Agronomic and physiological response of giant reed (Arundo donax L.) to soil salinity

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Abstract

The soil salinity increase in the Mediterranean basin is one of the consequences of the climate change. The aim of this study was to evaluate the adaptability of giant reed (Arundo donax L.) to salinity, in conditions of higher temperatures, in order to hypothesise the future use of giant reed under these conditions. The trial was carried out in pots under a permanent metal structure, open on the sides and with a clear PE on the top. Four levels of soil salinity in the range 3.3-15.5 dS m–1 were imposed. The stem number of the most stressed treatment was about 45% lower than the control and the stem height was lower than in all other treatments. The green and yellow leaf number decreased as the soil salinity increased, and their sum was significantly lower in the two most stressed treatments. Osmotic potential of the leaf sap was not affected by salinity. Leaf water potential and stomatal conductance in the saline treatments were lower than in the control. Assimilation rate showed similar pattern of stomatal conductance. Intrinsic WUE remained almost stable until July and increased during August under the most stressful conditions. PSII photochemistry was not affected by soil salinity. Biomass yield was not different from the control until to soil ECe 12.0 dS m–1; only the most stressed treatment (15.5 dS m–1) caused yield losses (50%). Tolerance threshold to salinity was 11.2 dS m–1 and the relative yield losses were 11.6% per dS m–1.

Introduction

In the perspective of climate changes, new cropping systems have to be developed by integrating energy, food, and animal feed production. In this framework, the low-input perennial species are considered the more suitable crops for supplying biomass to produce bio-fuels and biomaterials (Smith and Olesen, 2010). Nevertheless, energy crops must not compete for land with food crops: this means that they must be cultivated in soils not useful for food crops, such as saline, polluted, and other marginal soils (Impagliazzo et al., 2017).

Climate change scenarios in the northern Mediterranean basin show a temperature increase, particularly in their summer peak values (Brunetti et al., 2000). Furthermore, cropland salinity is predicted to increase due to climate change (Yeo, 1998). This means that crops grown in the Mediterranean area, in the forthcoming years could be subjected to multiple stresses that are directly or indirectly related to greenhouse effects and climate changes.

A rise of 5-29 cm in the sea level is expected in the next 30-40 years due to temperature increase, and its effects will be: waterlogging of coastal humid areas; fastest coastal erosion; intrusion of saline wedge into the estuary and into the river delta, and an increase of saline water infiltration into the groundwater of the coastal areas (Duce, 2005).

Salinity is one of the most important limiting factors for crop yield in many parts of the world; high saline concentration in soil solution (>4 dS m–1) reduce germination rate, growth and yield (Maas and Grattan, 1999). The first symptoms of saline stress are a growth decrease (Maas, 1986) and a reduction of leaf area index, followed by an increase in the specific leaf weight (leaf weight to leaf area ratio) (Longstreth and Nobel, 1979), a reduction in the number and dimension of the stomata (De Pascale et al., 1999), premature leaf senescence, and an overall yield decrease.

Greenway and Munns (1980) reported three different types of physiological stresses due to salinity: osmotic, nutritional, and toxic. The osmotic stress is due to the reduction of the soil water potential with a consequent reduction of root uptake, leaf water potential and stomatal conductance and, thus, of photosynthetic rate. When salt stress becomes more intense, photosynthesis is...
limited also by the mesophytic diffusive resistance and by metabolic damages (Corinic and Massacchi, 1996; Flexas et al., 2004; Chaves et al., 2009). The nutritional stress depends mainly on the modification in the nutrient uptake due to Na⁺ absorption (Munns and Termaat, 1986). Finally, the toxic stress depends on the accumulation of salts in the different organs of the plant, which, in turn, accelerates leaf senescence (Munns and Termaat, 1986). When the mortality rate of leaves overcomes the emission of new leaves, the whole-plant photosynthetic activity decreases.

All these adaptation mechanisms result in the yield decrease of glycophyte crops.

Among the different biomass crops for producing renewable energy aimed to reduce greenhouse gas emissions due to fossil fuels, giant reed (Arundo donax L.) is considered one of the most interesting species (Fagnano et al., 2015), thanks to its adaptability. It can grow in all soil textures, from heavy clay to sandy soils and it can withstand extended periods of severe drought (Lewandosky et al., 2003). Giant reed is native of Asia or of the Mediterranean basin, but it is now widely distributed in many temperate and sub-tropical areas of both hemispheres (Lewandosky et al., 2003).

Since it is a sterile plant, the natural variability may be due to spontaneous mutations and to natural selection as a response to different environmental conditions, and particularly to climatic stresses (Cosentino et al., 2006). It normally produces high lignocellulosic biomass yields in marginal lands, such as polluted soils (Fiorentino et al., 2010), hilly soil (Mantineo et al., 2009) also subjected to accelerated soil erosion (Fagnano et al., 2015), and under saline conditions (Williams et al., 2008; Williams et al., 2009). A. donax thanks to positive energy balance, due to high biomass production and low agro-management requirements (Lewandosky et al., 2003; Pompeiano et al., 2013), can become a promising crop for lignocellulosic feedstock (Pompeiano et al., 2017).

A. donax is a C₃ species with particularly high assimilation and transpiration rates (Rossi et al., 1998; Webster et al., 2016) that are comparable to several C₄ species. Presently, A. donax is not considered a halophyte because it primarily invades freshwater habitats (Nakley and Kim, 2015), but its salt tolerance has been proved worldwide: California, South Africa (Bell, 1997; Rossa et al., 1998), and Australia (Williams et al., 2008).

Recent studies report giant reed as moderately tolerant to salt stress (Nakley and Kim, 2015; Pompeiano et al., 2017), with an assimilation rate of 7-12 µmol CO₂ m⁻² s⁻¹ with 40 dS m⁻¹ of water salinity.

This study aims to evaluate the adaptability of A. donax to increasing salinity levels in higher temperature conditions, from an agronomic and physiological point of view, in order to suppose a future use of this energy crop in marginal soils for salinisation in a perspective of climate changes.

### Materials and methods

#### Plant material, salt treatments and experimental design

Rhizomes of giant reed were collected in March 2011 from a population growing in Bellizzi (SA), a coastal plain area of Southern Italy. The rhizomes were planted in 0.39 m² pots, filled with 0.10 m of gravel for drainage, and 0.45 m of loamy sandy soil (ISSS classification; Table 1). The 70 cm diameter and 60 cm height pots (200 L volume) were allocated in the experimental field of the Department of Agriculture (40° 48.870’ N; 14° 20.821’ E; 70 m a.s.l.) in Portici (Naples, Italy) under a 116 m² (14.5×8 m) permanent metal structure, open on the sides and with a clear PE on the top, to avoid the interference of rainfall. The plants were irrigated during the growth cycle about once a week, with 370 L per pot corresponding to the decennial average rainfall (950 mm per year).

Four levels of soil salinity [electrical conductivity (EC) 1:5 method] were initially imposed: control: EC soil between 0.2 and 0.6 dS m⁻¹ (Control); low saline stress: EC soil between 0.6 and 1.0 dS m⁻¹ (S1); medium-low saline stress: EC soil between 1.0 and 1.5 dS m⁻¹ (S2); moderate saline stress: EC soil between 1.5 and 2.0 dS m⁻¹ (S3). A completely randomised block design was adopted with 5 replicates per treatment.

The three levels of soil EC resulted from a precedent experiment on the response of food crops to salinity and they were obtained by saline irrigations at different salt concentrations (Mori et al., 2011). In order to maintain the initial soil salinity of the three treatments, every 15 days the soil EC was monitored by soil samplings in the layer 0-30 cm. If the soil salt level was lower than that initially imposed, irrigation of saline treatments was carried out, adding tap water with common salt. The soil electrical conductivity (1:5 method) was measured with a conductimeter (Basic 30 CRISON), and the pH with a digital pH meter. Besides, the percentage of soil water content was determined with the gravimetric method.

ECe (saturated past) was calculated by using the following robust (r²=0.98) equation, as proposed by Sommez et al. (2008) for sandy soils:

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ECe = 8.22 \times EC \times (1:5) - 0.33
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#### Biometric and physiological measurements

The measurements were carried out from April 2012 to January 2013 on re-sprouted plants after the first cut.

Plant growth was recorded nine times (every month), from 16 May 2012 to the harvest (early January 2013), by measuring the following parameters: number of stems, basal diameter, height of stems and number of green and yellow leaves.

At harvest, the fresh and dry weight of stems and leaves were measured. Dry matter values were determined after oven drying at 70°C until constant weight.

Measurements of all physiological parameters were taken on the uppermost sunlit well-expanded leaf in 5 replicate plants per treatment, with the exception of chlorophyll content index for which 12 leaves in each replicate plant were monitored (Table 1). Measurements were carried out between 10:00 am and 01:00 pm in the same days of the biometric samplings. Midday leaf water potential (Ψ, MPa) was measured by a Scholander pressure chamber (Model 3000; Soil Moisture Equipment Corp., Santa Barbara, CA, USA). Osmotic potential (Ψₛ, MPa) measurements were carried out on the same leaves used for gas-exchanges determination. The entire leaf lamina was frozen in liquid nitrogen and squeezed at fixed pressure to extract the cellular sap. The sap osmolality was measured by using a micro-osmometer (13/13 DR-Autocal-Hermann; Roelbing Messtechnik, Berlin, Germany) and it was then converted into osmotic potential by using Morse’s equation (Morse, 1914).

Net CO₂ assimilation rate (A, µmol m⁻² s⁻¹) at saturating light and stomatal conductance to water vapour (gₛ, mol m⁻² s⁻¹) were measured, by using a portable open-system gas-exchange analyser Li-6400 (Li-Cor Biosciences, Lincoln, NE, USA). Intrinsic water use efficiency (WUE) was calculated as the ratio A/gₛ (µmol mol⁻¹).

The CO₂ inside leaf chamber was supplied by an external cartridge.
to obtain a flow rate of 400 µmol mol⁻¹ air. A LED light source, with emission peaks centred at 670 nm in the red and at 465 nm in the blue, provided a photosynthetic photon flux density (PPFD) equal to 2000 µmol m⁻² s⁻¹ (90% red, 10% blue). The software of the instrument (Li-Cor, 2011) calculated the gas-exchange parameters based on the von Caemmerer and Farquhar (1981) model. Chlorophyll a fluorescence was measured, by using a continuous excitation Handy PEA fluorimeter (Hansatech Instruments Ltd., King’s Lynn, Norfolk, UK). Fluorescence was induced by a red (650 nm) light diode source for 1 s at the maximal available PPFD of 3500 µmol m⁻² s⁻¹ (Strasser et al., 2000; Giorio et al., 2011). The leaves were dark adapted for 30 min by means of the equipped white leaf-clips, prior to the determination of maximum quantum yield of PSII photochemistry calculated by the instrument software as Fv/Fm = (Fm – Fo)/Fm, according to Kitajima and Butler (1975).

The chlorophyll content was assessed as an optical index by using a handheld meter CCM-200 plus (Apogee Instruments, Inc., Logan, UT, USA). The meter measures the ratio between the leaf transmittance at 653 and 931 nm and calculates the chlorophyll content index, which is proportional to the amount of chlorophyll in the leaf.

**Statistical analysis**

The experimental design was one factor randomised complete block design combined over years (dates) with soil salinity as main factor and dates as repeated measures over the time.

The biometric data were analysed with MSTAT software (Crop and Soil Science Department, Michigan State University, Version 2.0). Physiological variables were processed by one-way ANOVA, followed by Tukey’s multiple comparisons test, using GraphPad Prism version 6.00 (GraphPad Software, La Jolla, California, USA).

**Results**

**Air temperature**

The trend of minimum and maximum air temperature during the crop cycle is shown in Figure 1. The temperature was higher under the shelter than outside, especially in the summer months, when the difference was 14.5°C for Tmax and 3.2°C for Tmin.

**Soil moisture and salinity**

The soil EC varied during the cycle; however, across the average, they were within the range imposed (Table 2). The soil water content (average of all samplings) increased as soil salinity increased (Table 2).

**Plant growth**

The main effects of soil salinity and sampling date were significant for all the growth parameters (Table 3). Interactions were not significant.

The effects of the two factors are reported in Table 4. The stem number per pot increased until the begin of August, although since July there were no significant differences (Table 4). The salt stress caused a significant decrease of stems emission (Table 4); the stem number of the most stressed treatment was about 45% lower than the control, but it was not different from the S2 treatment.

The increase in the stem height lasted until the beginning of July there were no significant differences (Table 4). The salt stress was significant for all the growth parameters (Table 3). Interactions were not significant for all the growth parameters (Table 3). Interactions were not significant. The main effects of soil salinity and sampling date were significant. The stem number per pot increased until the begin of August, although since July there were no significant differences (Table 4). The salt stress caused a significant decrease of stems emission (Table 4); the stem number of the most stressed treatment was about 45% lower than the control, but it was not different from the S2 treatment.
October (Table 4); in the two successive samplings, there were no further increases.

The effect of saline stress on the stem height was not very strong; only the S3 plants were statistically shorter than the all other treatments. The rate of stem elongation had a parabolic trend for all treatments (Figure 2); however, each reached the maximum peak in a different moment of season and with different values. The elongation of stems of the control plants began earlier than the other treatments with the highest rate (cm per day) until June, while the three saline treatments showed the higher stem elongation rates in the summer months.

During the cycle, the number of green leaves had a parabolic trend; the control reached the maximum in September, and the saline treatments in October; starting from the end of October, the green leaves number began to decrease significantly (Figure 3). The effect of saline stress on the green leaf number was very strong: the most stressed treatment showed the half of the green leaves of control (Table 4).

The first yellow leaves were recorded in July, and their number significantly increased until the harvest (Table 4). The yellow leaves decreased as the soil salinity increased, but only in S2 and S3 treatments they were different from the control (Table 4). The sum of the green and yellow leaves was significantly lower in the two most stressed treatments, about 28% and 43% less than the control.

The percentage of the number of green and yellow leaves on the total number of leaves (Figure 4), in the four most important moments of cycle (beginning, first emission of yellow leaves, full activity of plants, and end cycle/senescence), confirms that the control plants had a faster cycle; in fact, the senescence began earlier in control plants, and only at the end of October the percentage of yellow leaves of S3 plants was comparable with the control.

**Physiological responses**

Leaf water potential ($\Psi_l$) in May did not show significant differences among all the treatments, with an overall mean of −1.71 MPa (Figure 5A). Afterwards, $\Psi_l$ was higher in the control than in the saline treatments, with a 0.70 MPa difference between control and S3 in July and August.

The osmotic potential of the leaf sap (Figure 5B) was not affected by salinity from June to August, showing an average of −1.63 MPa in all treatments. In October, it was 0.48 MPa lower in S3 than in the control plants (−1.93 vs −1.45 MPa).

During the whole experiment, the stomatal conductance in S1, S2, and S3, was significantly lower than the control plants, with the exception of S1 in June (Figure 6A). However, no significant differences were found among the saline treatments. As an average of the three saline treatments, g, progressively decreased from 0.20 in June, to 0.045 in July, and 0.021 mol m$^{-2}$ s$^{-1}$ in August; the latter two corresponding to 10% and 13% of the control, respectively.

The assimilation rate (A) broadly followed the $g_s$ pattern (Figure 6B). The three salt treatments showed average values of 5.3 µmol m$^{-2}$ s$^{-1}$ in July and 6.7 in August, corresponding to 21% and 32% of the control, respectively. The two gas exchange parameters, partially recovered in October and November, when the data were comparable with those recorded in May.

Intrinsic water use efficiency remained almost stable until July, ranging from 86 to 123 µmol mol$^{-1}$ in all treatments (Figure 6C). As expected, a noticeable increase was observed in August under the most stressful conditions, when A/g, was 2.5-fold higher in S1 and S2, and 3.9-fold in S3 as compared to the control. Afterward, A/g, values progressively decreased in November as compared with those recorded in May.
The Fv/Fm ratio remained stable around an overall mean of 0.77 (r.u.) throughout the whole experiment, with no significant differences between the control plants and those subjected to salt treatment (S1, S2, S3) (Figure 6D).

The chlorophyll content index, measured over a few dates, resulted as an average 22% lower in the control than in the saline treatments (Figure 7).

**Biomass yield**

The yield and related parameters were all significantly affected by the soil salinity (Table 5).

The biomass yield (fresh weight) of the two less stressed treatments (S1 and S2) were not significantly different from the control; the S3 yield was about the half of the average value of the other three treatments (Table 6).

S1 and S2 plants showed a higher average weight of stems and a lower emission capacity of stems than the control but the differences were not significant.

The S3 plants always showed the lowest values for all parameters of yield (Table 6). In addition, the percentage of dry matter was higher for the control and the two less-stressed treatments (51.5 vs 49.1%, average of values of control, S1 and S2 and value of S3 respectively).

Yield was strongly correlated with soil salinity ($r^2=0.99$) (Figure 8); the model estimates 100% yield loss at EC of 2.3 dS m$^{-1}$ (1:5 method), corresponding to the ECe of 18.6 dS m$^{-1}$.

Similarly, the tolerance of the giant reed to salinity was also confirmed by applying Maas and Hoffman equation (1977). In fact, we found that the tolerance threshold to salinity (value beyond which the yield loss begins) for *A. donax* is 11.2 dS m$^{-1}$ and the relative growth decrease was 11.6% per 1 dS m$^{-1}$ increase. The percentage incidence of leaves dry matter increased with

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**Table 5.** Analysis of variance on yield parameters at harvest: significance of main factors.

| Parameter          | Soil salinity |   Significance | LSD |
|--------------------|---------------|---------------|-----|
| Stem number per pot| 0.05          | 3.06          |     |
| Stem basal diameter| 0.01          | 0.14          |     |
| Stem height        | 0.05          | 43.47         |     |
| Stem average weight| 0.05          | 7.71          |     |
| Yield (kg m$^{-2}$ fresh weight) | 0.05 | 0.45 |

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**Table 6.** Effect of soil salinity on yield (fresh weight) and its parameters at harvest.

| Treatment | Number (No.) | Height (cm) | Diameter (cm) | Average weight (g stem$^{-1}$) | Yield Fresh weight (kg m$^{-2}$) | Dry matter (%) |
|-----------|--------------|-------------|---------------|-------------------------------|---------------------------------|----------------|
| Control   | 9.5$^a$      | 158.2$^b$   | 1.11$^a$      | 39.0$^b$                      | 1.14$^a$                       | 52.0           |
| S1        | 8.5$^a$      | 192.1$^a$   | 1.08$^a$      | 51.3$^a$                      | 1.43$^a$                       | 52.5           |
| S2        | 6.8$^b$      | 188.8$^b$   | 0.98$^b$      | 54.1$^b$                      | 1.22$^b$                       | 50.1           |
| S3        | 5.3$^b$      | 137.5$^b$   | 0.77$^b$      | 35.2$^b$                      | 0.64$^b$                       | 49.1           |

$^a$-$^b$ Values with different letters indicate significant differences at P=0.05 and only for stem basal diameter at P=0.01.
saline stress, in fact the two more stressed treatments was slightly higher than the other two ones, 30.1% vs 23.0% respectively (Figure 9).

Discussion and conclusions

The salt tolerance of a crop depends not only on the intensity of saline stress but also on genotype, other biotic or abiotic stress (temperature, water deficit, soil physical conditions) and crop practices (Läuchli and Epstein, 1990). Many studies were carried out about the effect of salt on growth, physiology and yield of several food crops: tomato (Mori et al., 2008); potato (Patel et al., 2001); bean (Bayuelo-Jimenez et al., 2003); snap bean (Mori et al., 2011); cauliflower and broccoli (De Pascale et al., 2005), while,

![Figure 6. Stomatal conductance to water vapour (A), net CO₂ assimilation (B), intrinsic water use efficiency (C) and maximum quantum yield of PSII (D). Values represent average measurements, n=5 for A), B), C) and n=20 for D). Bars with the same letter are not significantly different according to Tukey test (P<0.05).](image)

![Figure 7. Content of chlorophyll a. Values represent average measurements, n=20. Bars with the same letter are not significantly different according to Tukey test (P<0.05).](image)

![Figure 8. Yield (t ha⁻¹ F.W.) vs average electrical conductivity of soil. The line represents the best fit regression model.](image)

![Figure 9. Percentage incidence of stems and leaves dry matter on total dry matter of plants of four treatments.](image)
few studies were carried out on biomass crops. To get complete information about the response of a crop to salinity, it is necessary to evaluate the behavior of plants at different times of the cycle in which the salt stress occurs. Therefore, it is important to evaluate: i) the survival of plants (sowed or transplanted) on saline soil; ii) the plant growth (absolute or relative, as compared to no-saline condition); and iii) the yield reduction. About the first point, our results confirmed that A. donax is able to survive on saline soil; in fact, the rhizomes of all treatments sprouted, in agreement with results reported by Nackley and Kim (2015) for stem cuttings of A. donax.

However, as it is well-known about food crops (Shahbaz et al., 2012), likewise for giant reed, the early growth phases are the more sensitive; in fact, sprouting capacity of rhizomes reduces to increasing saline stress (the stem number of the most stressed treatments was almost the half of control), as also reported by Sanchez et al. (2015).

The stem height decreased with saline stress; probably this parameter was also negatively influenced by the number of stems, according to Angelini et al. (2009); we found a relation between the number of stems per pot and their height (y = 2.7982x² – 67.924x + 536.4x – 1245.5; R²=0.999). This equation individuates, in our experimental conditions, an optimal range of stems per square meter, that ranged from 15 to 20, and that allowed the maximum elongation of stems. If the number of stems per pot is higher than 8, like it was for control plants, the height of stems is reduced, probably due to the higher competition for space and/or nutrients, reaching height values similar to those of the plants of most stressed treatments. On the other hand, as also found by Sanchez et al. (2015), the stem height decreased in the saline and saline-water stressed treatments. These researchers also found a decrease in stem diameter that ranged from 0.7 cm in control plants (well-watered with no-saline solution) to 0.6 cm in the most stressed plants (low watered with saline solution). We also found a similar effect of saline stress, but our values of basal diameter of stems was higher and, at harvest, ranged from 1.1 cm of control plants to 0.77 cm of S3 plants.

Additionally, the growth rate of plants was affected by salt stress in particular, which is especially evident for stem elongation rate. In fact, the control plants already showed a maximum elongation of stems in the first months of cycle, according to Spencer et al. (2005), which found that growth (stem elongation, leaf production and RGR) was most rapid prior to the second week in June, and to Nasi o Di Nasso et al. (2011) which highlighted the maximum crop growth rate at the middle of June. On the other hand, Triana et al. (2014) identified four growth stage for Arundo donax: i) initial (crop sprouting-beginning of stem elongation); ii) crop development, corresponding to stem elongation; iii) mid-season (end of stem elongation-beginning of canopy senescence); iv) late season (canopy senescence-end of water uptake), and in both test years the phase of elongation stem started about at beginning-middle of May and ended at most at 12 June, according to our findings about the control plants.

Probably the elongation activity for the stressed plants started later because the plants grown under salt stress needed a period of adjustment to saline conditions before starting to develop, and this period seems to be longer with greater salt concentration, as already indicated by the rhizomes sprouting capacity.

This hypothesis (the necessity of a period of adjustment) seems to be confirmed by the observation that from July the rate of elongation stems of these plants was even higher than the elongation rate of control plants. It is possible that in the first months of the cycle, these plants mainly used their energy to activate the mechanisms of adaptation to salt stress and/or develop roots, as already reported as a response of these crops to other environmental stresses (Fiorentino et al., 2017). In addition, the early emission of the yellow leaves of the control plants, seems to indicate that the cycle of these plants was faster, as confirmed at the harvest also by a higher dry matter percentage than the two more stressed treatments. However, the plants grown in saline conditions recovered from the delay, except for the most stressed plants that were the only ones to reduce biomass yield.

Similarly to the results reported by Sanchez et al. (2015), for the total dry weight of plants watered with saline solution of 16 dS m⁻¹, we also observed a negative relationship between the above-ground biomass per plant and salinity, but only in the most stressed treatments.

The physiological performances of potted giant reed in the control treatment were comparable with those reported in other experiments made with plants grown either in pots or in open field conditions: the mean seasonal stomatal conductance (0.298 mol m⁻² s⁻¹) and assimilation rates (21.4 µmol m⁻² s⁻¹) in control plants were similar to those reported by Sanchez et al. (2015) in potted plants grown in greenhouse, by Cosentino et al. (2016) in two-year old fully irrigated plants in open field conditions, and by Webster et al. (2016) in natural stands. We are not aware of research reporting leaf water potential of giant reed under salt stress conditions; however, Cosentino et al. (2016), in plant irrigated with 50% ETM, and Mann et al. (2013), under mild drought (9% soil moisture, soil matric potential, Ψₘ = -0.5 MPa), reported leaf water potential values proximal to the values (–2.08 MPa) measured in our S3 treatment. Therefore, we can infer that the most salt-stressed plants of our experiment experienced mild stress conditions. Under open field conditions during the summer, a decrease of gᵣ in control plants as a mechanism of drought avoidance (Jones, 1992) is normally observed, because of the increased atmospheric evaporative demand. In our experiment, plants grown under the shelter, experienced higher VPD (data not shown) and temperature (Figure 1) than in open air, which induced an evident gᵣ reduction in early August. The interaction between such environmental conditions and salinity can explain the marked decrease of stomatal conductance. We observed a good relationship between A and gᵣ (data not shown) in all treatments as reported by Nackley and Kim (2015) on the same species. This would indicate that the reduction of assimilation rate could be ascribed to a lower availability of CO₂ due to stomatal closure. Moreover, same Authors reported that plants irrigated with saline water (40 dS m⁻¹) reached stomatal conductance of about 0.05 mol m⁻² s⁻¹ without impairment of PSII photochemistry, as showed by high ΦPSII values. We found high values of Fᵥ/Fₘ in all treatments during the entire experiment, indicating that, under our experimental conditions, PSII reaction-centres were not compromised (Baker, 2008). Indirect evidence that photosynthetic limitation was only due to stomata and not due to photosynthetic machinery can also be deduced by the strong increase in intrinsic WUE at low gᵣ, (Cifre et al., 2005; Flexas et al., 2005), as observed under the most stressful conditions in summer.

Salt stress causes a reduction in the chlorophyll content as found in rice (Ali et al., 2004) and pearl millet (Sneha et al., 2014). As regards Arundo donax, Nackley et al. (2015) found no variation in plants irrigated with saline water (ECw 0-40 dS m⁻¹), while Sanchez et al. (2015) observed either an increase or no variation in moderately salt stressed plants, depending on genotypes, as we found in our experiment. It can be speculated that shrinkage or reduced expansion of leaves in stress treatments may have contributed to enhance the leaf specific chlorophyll content (Tang and Boyer, 2007). Both gas-exchange and fluorescence parameters
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