**Effect of the foliar application of microalgae hydrolysate (*Arthrospira platensis*) and silicon on the** **growth of *Pelargonium hortorum* L.H. Bailey under salinity conditions**

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Keywords: NaCl, biostimulants, abiotic stress, bedding plant

**Abstract**

Plant growth is limited by salinity stress yet there are few strategies for alleviating it. *Arthrospira platensis* and silicon can stimulate plants to grow under stress conditions. The aim of this work was to study the effects of both a single and a joint application of *Arthrospira platensis* and siliconon the growth of *Pelargonium hortorum* L.H. Bailey under salt stress conditions. Plants were exposed to 2.0, 3.0 and 3.5 dS m-1 EC, with and without the application of microalgae and silicon. At the end of the trial, biometric parameters and plant analysis were determined. The microalgae hydrolysate concentration was 5 g L-1 and the silicon concentration was 150 mg L-1. Foliar spraying was applied weekly.Pelargonium can be grown in moderately saline irrigation water (3.0 dS m-1). This bedding plant mitigates salt stress by avoiding the uptake of Cl- ions and by tolerating a high Na+ concentration in the tissue. The joint foliar application of *Arthrospira* microalgae and silicon stimulates root, shoot, leaf and flower formation in the *Pelargonium hortorum* L.H. Bailey crop under salinity conditions (3.5 dS m-1).

Keywords: NaCl, biostimulants, abiotic stress, ornamental plant

**1. Introduction**

There is a growing need to develop strategies aimed to reduce or replace the use of chemicals inputs, with natural or biological substances; plant biostimulants [1]. Plant biostimulants have attracted interest in modern agriculture as a tool for enhancing crop performance, resilience to environmental stress, and efficient nutrient use [2]. The development of innovative technologies based on bioresources, including plant biostimulants, have proven to be effective methods for improved crop performance [3]. Plant biostimulants contain substances and/or micro-organisms whose function, when applied to the plants or the rhizosphere, is to stimulate natural processes that enhance/benefit nutrient uptake, nutrient efficiency, tolerance to abiotic stress and crop quality [4].

Microalgae, which comprise eukaryotic and prokaryotic cyanobacteria (blue-green algae), are attracting increased interest from scientists, extension specialists, plant growers and private industry because of their versatile nature: this includes their simple unicellular structure, high photosynthetic efficiency, potential for heterotrophic growth and valuable co-products yield, adaptability to domestic and industrial wastewater, and amenability to metabolic engineering [2]. Microalgae are classified mainly by their pigmentation, life cycle and cell structure [5]. The dominating microalgae species available commercially are: *Isochrysis* spp., *Chaetoceros* spp., *Chlorella* spp., *Arthrospira* spp. and *Dunaliella* spp. Of these, *Arthrospira* spp. and *Chlorella* spp. are the main species cultivated and used commercially [5]. Microalgae are microscopic plants that contain potential bioactive materials in the form of proteins, lipids, glycerols, carotenes and vitamins [6]. The potential of these microorganisms as resources for bioactive compounds is attributed to their remarkable capacity to synthesize useful products such as polysaccharides, proteins, polyunsaturated fatty acids, lipids and other bioactive metabolites from atmospheric CO2 [3]. The average nutritional composition of the microalgae (expressed as g per 100 g dry weight) is protein 63, fat 4.3, carbohydrates 17.8, chlorophyll 1.15, magnesium 0.316, beta-carotene 0.12, vitamin B1 (thiamin) 0.001, vitamin B2 (riboflavin) 0.0045, vitamin B3 (niacin) 0.0149, vitamin B5 (pantothenic acid) 0.0013, vitamin B6 (pyridoxine), vitamin B9 (folic acid) and vitamin B12 (cobalamin) [6]. The presence of certain amino acids in microalgal extracts is expected to significantly increase the growth and yield of cultivated crops because they are the metabolic precursors of various phytohormones [7]. Arginine serves as the precursor to polyamines, which are involved in floral initiation and development, and in protecting against osmotic stress [8,2]. Several studies have reported a positive effect of microalgae on plant growth. It has been proved that foliar application of hydrolysates with microalgae accelerates plant development in terms of higher rates of root growth, leaf and shoot development, and earliness of flowering [9].

Silicon, on the other hand, can stimulate growth and yield both directly and indirectly [10, 11]. Plants take up Si in the form of silicic acid, which is transported to the shoot; following water loss, it is polymerized as silica gel on the surface of leaves and stems [11]. Silicon has been shown to mitigate the impact of many biotic and abiotic stresses [12,10]. The beneficial effect of Si may be due to the Si-induced decrease in transpiration, and to the partial blockage of the transpirational bypass flow [10]. There has been limited research addressing the effects of silicon on container-grown ornamentals [11]. Since most ornamental crops are grown in soilless media, where the silicon concentration is minimal [11], evaluating the effect of this element has attracted great interest. The beneficial effect of applying silicon to ornamental plants is evidenced by their improved health, firmness and proper flower formation [13]. Application of the Si+P fertilized counteracted the negative effects of water stress on vegetative growth and certain biochemical parameters of potted geranium plants [14]. However, the effect of Si supplements depending on Si source and concentration [15]. In the case of KSiO3, the most beneficial Si concentration was 140 g m-3 [15].

Previous studies revealed that using microalgae and silicon singly can stimulating growth, flowering and fruiting of different crops. However, limited research has been done to study the effect on plant growth of using of silicon in combined with microalgae. In this regarding, supplying banana plants with a mixture containing potassium silicate and seaweed extract gave the best results to growth, earlies shooting, yield and fruit quality [16].

The aim of this work was to study the effect of single and joint applications of *Arthrospira platensis* and siliconon *Pelargonium hortorum* L.H. Bailey growth under salt stress conditions.

**2. Material and methods**

**2.1. Greenhouse trial**

The trials were carried out in a 170 m2 greenhouse with zenith ventilation along with relative humidity and temperature control. The temperature and relative humidity were recorded every 15 min with a HOBO U12-013 data logger (Onset Computer Corporation, Bourne, MA, USA) placed at canopy height in the central part of the cultivation table where the plants are grown. External radiation was measured every 15 min with a Q20-B sensor. To estimate the internal radiation, the cover transmission coefficient was estimated as the ratio between the internal and external radiation using a manual quantum photoradiometer (Delta OHM, model RAD/ PAR). The average temperature, humidity and global radiation in the greenhouse were 17.5 ºC, 68.6 % and 1.40 MJ m2day-1, respectively; the maximum and minimum averages of the temperature and humidity were 24.6 and 12.4 ºC, and 79.6 and 52.9 %, respectively.

The species studied was *Pelargonium hortorum* L.H. Bailey**.** *Pelargonium hortorum* was chosen as a model plant because it has a fast crop cycle and is easily grown in a greenhouse. These results can be applied to others horticultural crops. Plant cuttings were transplanted into 2.5 L pots (one plant per pot). The substrate used was a mixture of peat and perlite 80:20 (v/v). The white peat substrate (Brill WPS medium) is a potting substrate with a high air capacity (41 %), a pH of 5.7 and a 1.0 g L-1 salt content. Projar expanded perlite has a total porosity of 92-98% and a 1-5 mm fraction size.

Fertigation was applied manually until the leachate fraction reached 20 %. The average dose for the *Pelargonium hortorum* was 85 ml per plant once every two days. For the C2, C3 and C3.5 treatments, leachates from 3 plants were collected in a tray with plastic saucers and covered with black film. The amounts of leached water, the EC and the pH were determined every 3 days using a test tube, a pH-meter and an EC-meter respectively. Water uptake was estimated as the difference between the water applied and the water leached.

**2.2. Treatments**

Twelve treatments were performed, single and joint foliar application with *Arthrospira platensis* and silicon and fertigation at EC 2.0, 3.0 and 3.5 dS m-1 (Figure 1).





Figure 1. Schematic diagram of the treatment performed. C- foliar application with water, A-foliar application with *Arthrospira platensis*, Si-foliar application with silicon, ASi- foliar application with *Arthrospira platensis* and silicon,  2- fertigation with an EC 2.0 dS m-1, 3- fertigation with an EC 3.0 dS m-1 and 3.5- fertigation with an EC 3.5 dS m-1 and *Pelargonium hortorum* plants cultivated in the greenhouse, under the different treatments tested.

The C2 concentrations (the standard nutrient solution) were: 0.52, 7.20, 0.71, 1.71, 3.00, 2.49 and 1.0 mmol L-1 of H2PO4-, NO3-, NH4+, SO42-, K+, Ca2+ and Mg2+, respectively. Phosphoric acid, nitric acid, calcium nitrate, potassium nitrate, ammonium nitrate and potassium sulphate were used to prepare the nutrient solutions. The pH and EC of the standard nutrient solution was 6 and 2 dS m-1 respectively. After the nutrient solution was prepared, NaCl (491 and 982 mg L-1) was added to obtain EC 3.0 and 3.5 dS m-1 respectively.

The potassium silicate was sprayed at a concentration of 150 mg L-1. The microalgal hydrolysates concentration was 5 g L-1. The density of algae cells used was 100 g L-1. Foliar spraying was applied weekly. *Arthrospira platensis* was supplied by the Biorizon company (Almeria, Spain). The microalgae hydrolysates were produced using Arnon medium, with addition of 16 g L-1 of bicarbonate, in a 100 m2 raceway reactor. The cultures were carried out under pH control (pH=10) by injection CO2. The oxygen concentration was controlled and remained below the 250 % saturation. The biomass was harvested continuously at a dilution rate of 0.2 1/day.

The nutrient content of the microalgae hydrolysate was N 1.198±0.005, P 0.187±0.004, K 0.157±0.005, Ca 0.157±0.004 and Mg 0.048±0.001 (%) [9]. The hormone content of the microalgae hydrolysate was cytokinins 2752, gibberellins 56.24, indoleacetic acid 10.3, ABA 1.03, salicylic acid 0.61 and jasmonic acid 0.84 ng g-1 [9].

**2.3. Nutrient solution analysis**

The parameters determined in the nutrient solutions tested were pH, EC, NO3-, SO42-, H2PO4-, Ca2+, Mg2+ and K+. The pH was measured with a Crison Micro pH 2001 pH-meter and the EC with a Crison Micro CM 2200 conductivity meter. Nitrate, SO42-, H2PO4-, Ca2+, Mg2+ and K+ were determined by HPLC (High Performance Liquid Chromatography; Metrohm 883 Basic IC Plus). NO3-, SO42- and H2PO4- were quantified using a Metrosep A SUPP 4 column (with an IC conductivity detector range of 0–15000 μS cm−1). The mobile phase was prepared by mixing 190.6 mg of CO32− and 142.8 mg of HCO3−, and then diluting this in 1 L of deionized water, acidified with H2SO4 (50 mM). The Ca2+, Mg2+ and K+ were quantified using a Metrosep C4 column (with an IC conductivity detector range of 0–15000 μS cm−1) and the mobile phase was prepared by mixing 117 mg of 2,6- Pyridinedicarboxylic acid and 1.7 ml of nitric acid (1 M) diluted in 1 L of deionized water.

**2.4. Biometric parameters**

 Growth parameters and ornamental quality are particularly important for the ornamental species, since can determine the price of the plant. Salinity can affect biomass production, leaf area, plant heights and water status. Therefore, fresh and dry weights, internode length, leaf length and width, and number of leaves and shoots were recorded at the end of the cultivation. The number of leaves and shoots per plant were counted. The internode length and the leaf length and width were measured using a ruler. The number of flowers per plant was counted each week during the crop cycle. After removing the substrate, the plant material (comprising 12 plants per treatment) was separated into different vegetal fractions: absorption organs (roots), conductive organs (stems and petioles), photosynthetic organs (leaves) and flowers, which were weighed separately on a COBOS G M5-1000 scale (with a precision of 0.005 g) to determine the fresh weight (FW). All the samples were washed and dried in a Nüve EFN500 oven (with a 30 to 300 ºC range) at 60ºC for 48 h to determine the dry weight (DW). The total DW was calculated as the sum of the roots, stems and petioles, leaves and flowers. The FW and DW totals were used to calculate the water content (WC) as (FW-DW)x100/FW. The leaf area was estimated by a non-destructive method using the formula S = *a* + *b*LW, as proposed by Giuffrida et al. (2011), where S is the foliar surface area, L is the leaf length (cm), W is the leaf width, and the a and b coefficients are specific to each species. For Pelargonium, a is 0.07 cm2 and b is 0.68 cm2. Leaf color was identified according to the Munsell chart for leaves. The Munsell color is based on a three-dimensional model in which each color is comprised of three attributes: hue (the color itself), value (lightness/darkness) and chroma (color saturation or brilliance). The color of the leaves was visually matched [17].

**2.5. Mineral analysis**

A subsample of the dry matter (0.2 g of dry weight) of the most significant treatments (C2, C3, C3.5, ASi2, ASi3 and ASi3.5) was ground up in a Wiley mill and digested (in 96% H2SO4) in the presence of hydrogen peroxide (H2O2, 30 %(w/v)) at 300 ºC in a sand bath to analyze the organic N, P, K, Ca and Mg. Each treatment was replicated four times. The total Na and K were directly measured by flame spectrophotometry using an Evans Electro Selenium LTB Flame Photometer (Halstead, Essex, England). The total Ca and Mgwere analyzed by atomic-absorption spectrophotometry using a Perkin Elmer Atomic Absorption Spectrometer 3300. P was analyzed using the method proposed by [18], and nitrogen was analyzed using the method proposed by [19].

**2.6. Experimental design and statistical analysis**

The experimental design comprised a completely randomized block with 12 treatments. There were 4 replicates per treatment and 3 plants (pots) per replicate (12 plants per treatments). The treatments’ effect significance (salinity and the microalgae and silicon application) was examined using the standard analysis of variance (one-way ANOVA) and Fisher’s Least Significant Difference (LSD) test, performed using Statgraphics Centurion XVI.II software (Statpoint Technologies, Inc. Warrenton, Virginia, USA). Differences were considered significant at *P*<0.05.

**3. Results**

**3.1. Salinity effect**

No differences in stems, leaves, flowers, or total dry weight were found between the salinity treatments (Table 1). However, Pelargonium underwent a negative response in root growth when the salinity level rose from 2 to 3.5 dS m-1. The treatments with an EC of 3.0 and 3.5 dS m-1 resulted in a reduction of more than 20%, compared with an EC of 2.0 dS m-1. The number of shoots, the width and length of the leaves and the internode length showed no significant differences (Table 2). However, the leaf area and leaf number improved in the treatment with an EC of 3.0 dS m-1 and number of flowers in the treatment with an EC of 3.5 dS m-1. Na concentration in leaf was high in all treatments [10], even in C2 treatment where sodium has not been added. This is due to the relatively high concentration of sodium in irrigation water. The Na levels were significantly higher in the leaves of plants that were given the salinity treatments (Table 3) However, no difference between treatments C2.0 and C3.0 were found. This similar concentration in the leaves was probably attributed to the effects on the reduction in the root dry weight. However, N, P, S Cl, Ca, and Mg were not affected by the salinity treatments. The K concentration in the leaves decreased as the salinity increased, reducing by 20 % in treatment C3.5 compared to C2. The evaluation of leached solution over the cultivation period is shown in Figure 2. The leachate pH did not present significant differences between the treatments (6.6-7.4). Nonetheless, the salinity treatment did affect the leachate EC, indicating that salt accumulation varied with treatment [21]; 59 days after transplanting, the leachate EC was 3.47, 5.10 and 6.06 dS m-1 for C2, C3 and C3.5, respectively. In all the treatments, the leachate EC increased during the assay, but this increase was greater in treatments C3 (50%) and C3.5 (60%) than in the C2 treatment (30%). However, no difference in the plant water content was found in treatments C3 and C3.5 compared to C2; the water content was 90.9, 91.1 and 91.0%, respectively. Differences in water uptake were not found until 55 days after transplanting. After 55 days after transplanting, plant water uptake was reduced in the NaCl treatments. The water uptake was 17 and 12 % lower in C3 and C3.5, respectively, compared to C2 (58 days after transplanting) (Figure 3).

**3.2. Silicon and microalgae application**

Combined applications of microalgae and Si improved the leaves, roots, inflorescence and total dry weight in the treatment with an EC of 3.5 dS m-1 while it improved the inflorescence dry weight in the treatment with an EC of 3.0 dS m-1 (Figures 4, 5 and 6). However, no effect was observed on the roots, ~~or~~ the inflorescence and stem dry weight between the treatments with an EC of 2.0 dS m-1. The Si treatments improved leaf dry weight compared with the control in the treatments with an EC of 2.0 and 3.5 dS m-1. A single application of microalgae improved the inflorescence dry weight in treatments with an EC of 3 dS m-1 although a reduction in total dry weight was observed in treatments A2 and A3.5. In treatments A2 and Si2, the number of shoots increased with an EC of 2 compared with the control, as it also did in treatments Si3.5 and ASi3.5 with an EC of 3.5 dS m-1 (Figure 7a). The leaf number increased compared to the control (by 22 and 28 %) as did the flower number (by 24 and 51%) in treatments Si3.5 and ASi3.5, respectively (Figures 7b and 8a). Furthermore, applying *Arthrospira platensis* and silicon increased the flower number in the treatments with an EC of 3.0 and 3.5 dS m-1. No differences were found in the internode length or the leaf area (Figure 8b and 9). The Munsell chart was used to define color for all the treatments at the end of the assay. No differences were found between treatments; the leaves were identified as 5GY 4/4 for color, value and chroma, respectively. The Cl and Na concentrations increased by 15 % in treatment ASi3.5 compared to C3.5, as did the P and Ca concentrations in treatment ASi3 compared to C3 (Figure 10). The Si levels were significantly higher in plant leaves that had been treated with the *Arthrospira* and silicon foliar application. In contrast, the nitrogen, magnesium and sulfur contents were not affected by the microalgae and silicon application.

**4. Discussion**

Salinity can reduce growth in plants; this growth reduction may result from the effects of salt on physiological processes, biochemical relationships, or a combination of such factors [22]. However, salinity affects the biomass and biochemical parameters differently depending on the species [23]. The result of this assay indicated that salinity caused a reduction in root growth. The root/shoot ratio for treatments C2, C3 and C3.5 were 0.065, 0.058 and 0.058 g g-1, respectively. Root growth appears to be limited by increasing Na concentrations in the growth medium [10]. However, the C3 treatment, which had an EC of 3 dS m-1, improved the leaf area and leaf number compared to C2 and C3.5 and the number of flowers increased in the most saline treatment. A number of herbaceous perennials, groundcover plants, floricultural crops, and landscape woody shrubs became more compact with little visual damage when irrigated with low to moderate salinity compared non-saline control [21]. Pelargonium may be considered a plant that is moderately tolerant to salinity [22].

Water uptake was affected 56 days after transplantation in treatments that had an EC of 3.0 and 3.5 dS m-1. Furthermore, no differences in water content were found. In spite of the high salt concentration in the soil causing decreases in water uptake, salt stress did not cause changes in the water status of the pelargonium leaves, which shows that the plants did not suffer from water deficit stress [24]. There are three salt tolerance mechanisms in plants: osmotic stress tolerance, tolerance of tissue to accumulated Na and/or Cl ions, and minimizing the entry of Na and/or Cl ions [25]. Both the proline and anthocyanins contents increased significantly in pelargonium leaves growing in peat substrate with the two highest NaCl doses [24]. These compounds may function as osmotic adjusters as well as scavengers of oxygen radicals protecting cytoplasmic structures and chloroplasts from the adverse effects of toxic ions [24]. According to our results, the ability to maintain turgor in the plant probably also resulted from Na+ accumulation in the vacuoles after Na+ transport from the root to the aerial parts; this is an important mechanism of salinity tolerance [24]. Soil salinity can affect nutrient acquisition by severely reducing root growth [26] and deficiencies or imbalances in the plants which depend on the ionic composition of the external solution [26]. In accordance with our results, the concentration of potassium ions in the leaves was found to change significantly with increased NaCl concentration in the growing medium [24]. Moreover, leaf P, N, Ca, Mg and S concentrations did not present any differences. Some species tolerate salt stress by avoiding the uptake of certain ions or by tolerating a high ion concentration in the tissue [21]~~.~~ The ability of a plant to tolerate excessive amounts of Na+ varies widely among species [26]. Pelargonium seems to tolerate high Na concentrations in the tissue. Despite observing an increased Cl concentration in the external solution, no statistical difference in the Cl concentration in the leaves was found. One of the most important traits associated with tolerance is the ability to restrict the entry of saline ions through the roots [27]; however, a good Cl- ion excluder is not necessarily a good Na+ excluder, and vice versa [23]. *Pelargonium hortorum* L.H. Bailey seems to be a good Cl- ion excluder.

The results obtained by [11] found that silicon could be used as a fertilizer additive to improve the growth and quality of silicon-responsive ornamental plants. In our assay, leaf dry weight and leaf area were positively affected by the single application of silicon in treatments with an EC of 2.0 dS m-1. A single application of *Arthrospira* increased the number of shoots in treatments with and EC of 2.0 dS m-1, and the flower dry weight and number of flowers in treatments with an EC of 3.0 dS m-1. Regarding ornamental plants [9] showed that a foliar application of *Arthrospira platensis* improved the root dry matter and the number of flowers per plant. In petunias, applying microalgae stimulates bud formation in the mother plant and induces flowering in commercial petunia plants [9,28]. In this assay, combined application of *Arthrospira* and silicon gave the best results comparing with using each compound alone. The increase on root growth and the stimulation of flower and shoot formation, may be due to the synergistic effect of microalgae and silicon and their effect on some hormones such abscisic acid and cytokines [9,29]. Cytokinin promote bud development and abscisic acid promote root growth. In addition, [16] suggest that the advancing effect of silicon and seaweed on flowering time might be attributed to their essential role in balancing the ratio between carbohydrates and nitrogen in favour of flowering. According our result, [13, 28] show that the positive effect of microalgae and silicon, has been shown to be particularly beneficial under stress conditions. A wide array of chemicals has been identified in seaweed extracts, including polysaccharides, phenolics, fatty acids, vitamins, osmolytes, phytohormones, and hormone-like compounds involved in the plant-signaling response to abiotic stress [2]. In *Spirulina spp*., carbohydrates may account for up to 46 % of the dry weight (DW) extract [2], and this molecule can play an important role in osmotic adjustment [27]. In this sense, the major role played by sugars in stress mitigation involves osmoprotection, carbon storage, and scavenging ROS [30]. Moreover, amino acids can contribute to mitigating the damage caused by abiotic stress [5]. Furthermore, the application of Si has been shown to alleviate the negative effects of numerous abiotic stresses, including salt. Several potential mechanisms associated with stress alleviation in higher plants have been identified: increased structural reinforcement, altered photosynthetic rate, changes in stomatal conductance and enhanced water use efficiency [29]. Other possible mechanisms may also include osmotic adjustments through increased water potential and water content, reductions in oxidative stress, alterations in mineral uptake and accumulation and alterations to phytohormone concentrations [29].

Notably, all the essential minerals (about 7%) are available in Spirulina, including potassium, calcium, chromium copper, iron, magnesium, manganese, phosphorus, selenium, sodium and zinc [6]. The nutrient analysis of *Arthrospira* spp. dry biomass revealed that it contained 6.70, 2.47 and 1.14% (on a dry basis) of N, P and K, respectively , while the microalgae’s calcium (Ca) content was lower, relative to the other minerals [5]. In contrast to the results of other authors, no clear effect on nutrient concentration in the leaves was found after applying only the microalgae. On the other hand, the joint application of the microalgae and Si increased the P and Ca concentrations in the leaves in the treatment with an EC of 3.0 dS m-1, whereas it increased the Cl and Na in the treatment with an EC of 3.5 dS m-1. Conversely, [31] observed that applying silicon significantly decreased the Na+ concentration in the leaves while increasing the K+ concentration, compared with no saline treatment. In accordance with our results, it was found that the nitrogen, potassium, magnesium and sulfur contents were not affected by applying *Arthropira* to *Petunia x hybrida* [16].

**5 Conclusions**

*Pelargonium hortorum* L.H. Bailey is a plant that can be considered moderately tolerant to salinity. The mechanisms that confer salinity tolerance to *Pelargonium* are its ability to avoid Cl- uptake and to exhibit moderate tolerance to high internal Na+. Our results show that the joint application of *Arthrospira platensis* and silicon improves the flower dry weight and the number of flowers in treatments with an EC of 3.0 dS m-1, while improving the leaf, root, flower and total dry weight, and the number of flowers, shoots and leaves in treatments with an EC of 3.5 dS m-1. Combined applications of microalgae and silicon can improve plant growth and yield production by stimulating vegetative growth and promoting plant flowering. This strategy can be an alternative to chemicals used in agriculture.

**Declaration of Competing Interests**

The authors declare that they have no competing financial or personal interests that could have influenced the work reported in this manuscript.

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Table 1. Dry weight (DW) (g) of S (stems and petioles, conductive organs), R (roots), L (leaves), I (Inflorescence) and T(total) at the end of the trial for the C2 (fertigation EC 2.0 dS m-1 ), C3.0 (fertigation EC 3.0 dS m-1) and C3.5 (fertigation EC 3.5 dS m-1 ) treatments. Different letters indicate significant differences between treatments at the P<0.05 level using the LSD test**.** Values are the means of 12 plants.

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|   | SDW | RDW | LDW | IDW  | TDW |
| C2 | 1.46±0.09 | 0.30±0.01a | 1.85±0.13 | 0.48±0.07 | 4.62±0.21 |
| C3 | 1.52±0,08 | 0.24±0.01b | 2.04±0.13 | 0.33±0.07 | 4.14±0.20 |
| C3.5 | 1.50±0.07 | 0.25±0.02b | 1.97±0.03 | 0.49±0.07 | 4.64±0.14 |

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|   | Leaf area  | Leaf number  | Internode lenght  | Soot number  | Flower number  | Width Leaf  | Length Leaf  |
| C2 | 680.4±34b | 18.41±0.2b | 21.3±1.26 | 0.83±0.76 | 18.2±2.3 | 12.52±1.38 | 11.14±1.49 |
| C3 | 855.4±33a | 19.72±0.2a | 18.97±1.26 | 1.16±0.79 | 12.22±2.4 | 12.48±0.52 | 11.33±0.46 |
| C3.5 | 761.2±37b | 16.00±0.2b | 18.65±1.26 | 0.91±0.83 | 15.45±2.2 | 12.33±1.15 | 11.11±1.03 |

 Table 2. Leaf area, internode length, leaf, shoot , and flower numbers and width, and leaf length for the C2 (fertigation EC 2.0 dS m-1), C3.0 (fertigation EC 3.0 dS m-1 ) and C3.5 (fertigation EC 3.5 dS m-1 ) treatments. Different letters indicate significant differences between treatments at the P<0.05 level using the LSD test**.** Values are the means of 12 plants.

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|   | N (%)  | P (%) | S (%) | Cl (%) | K (%) | Ca (%) | Mg (%) | Na (%) |
| C2 | 3,72±0,20 | 0,25±0,01 | 0,11±0,01 | 0,88±0,11 | 1,59±0,19a | 0,75±0,07 | 0,09±0,01 | 0,47±0,04b |
| C3 | 3,82±0,30 | 0,23±0,01 | 0,11±0,00 | 0,91±0,03 | 1,55±0,09a | 0,71±0,01 | 0,09±0,02 | 0,53±0,02ab |
| C3.5 | 3,65±0,25 | 0,23±0,01 | 0,11±0,00 | 1,02±0,03 | 1,26±0,10b | 0,78±0,01 | 0,07±0,01 | 0,59±0,03a |

Table 3. N, P, S, Cl, K, Ca, Mg and Na concentration in the leaves at the end of the trial for the C2 (fertigation EC 2.0 dS m-1), C3.0 (fertigation EC 3.0 dS m-1 ) and C3.5 (fertigation EC 3.5 dS m-1 ) treatments. Different letters indicate significant differences between treatments at the P<0.05 level using the LSD test**.** Values are the means of 12 plants.

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Figure 2. Ph (a) and EC (b) of the leachate collated during the day after transplanting for the C2 (fertigation EC 2.0 dS m-1) , C3.0 (fertigation EC 3.0 dS m-1 ) and C3.5 (fertigation EC 3.5 dS m-1 ) treatments.

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Figure 3. Water uptake during the day after transplanting for the C2 (fertigation EC 2.0 dS m-1), C3.0 (fertigation EC 3.0 dS m-1) and C3.5 (fertigation EC 3.5 dS m-1) treatments.

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Figure 4. Leave and stems dry weight (g) at the end of the trial for the C2 (foliar application with water and fertigation EC 2.0 dS m-1), A2 (foliar application with *Arthrospira platensis* and fertigation EC 2.0 dS m-1), Si2 (foliar application with Si and fertigation EC 2.0 dS m-1), ASi2 (foliar application with *Arthrospira platensis* and Si and fertigation EC 2.0 dS m-1), C3.0 (foliar application with water and fertigation EC 3.0 dS m-1), A3.0 (foliar application with *Arthrospira platensis* and fertigation EC 3.0 dS m-1), Si3 (foliar application with Si and fertigation EC 3.0 dS m-1) , ASi3 (foliar application with Si and *Arthrospira platensis* and fertigation EC 3.0 dS m-1) treatments, C3.5 (foliar application with water and fertigation EC 3.5 dS m-1), A3.5 (foliar application with *Arthrospira platensis* and fertigation EC 3.5 dS m-1), Si3.5 (foliar application with Si and fertigation EC 3.5 dS m-1) and ASi3.5 (foliar application with Si and *Arthrospira platensis* and fertigation EC 3.5 dS m-1) treatments.. Different letters indicate significant differences between treatments at the P<0.05 level using the LSD test. Values are the means of 12 plants. Error bars show standard deviations.

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Figure 5. Roots and inflorescence dry weight (g) at the end of the trial for the C2 (foliar application with water and fertigation EC 2.0 dS m-1), A2 (foliar application with *Arthrospira platensis* and fertigation EC 2.0 dS m-1), Si2 (foliar application with Si and fertigation EC 2.0 dS m-1), ASi2 (foliar application with *Arthrospira platensis* and Si and fertigation EC 2.0 dS m-1), C3.0 (foliar application with water and fertigation EC 3.0 dS m-1), A3.0 (foliar application with *Arthrospira platensis* and fertigation EC 3.0 dS m-1), Si3 (foliar application with Si and fertigation EC 3.0 dS m-1) , ASi3 (foliar application with Si and *Arthrospira platensis* and fertigation EC 3.0 dS m-1) treatments, C3.5 (foliar application with water and fertigation EC 3.5 dS m-1), A3.5 (foliar application with *Arthrospira platensis* and fertigation EC 3.5 dS m-1), Si3.5 (foliar application with Si and fertigation EC 3.5 dS m-1) and ASi3.5 (foliar application with Si and *Arthrospira platensis* and fertigation EC 3.5 dS m-1) treatments.. Different letters indicate significant differences between treatments at the P<0.05 level using the LSD test. Values are the means of 12 plants. Error bars show standard deviations.

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Figure 6. Total dry weight (g) at the end of the trial for the C2 (foliar application with water and fertigation EC 2.0 dS m-1), A2 (foliar application with *Arthrospira platensis* and fertigation EC 2.0 dS m-1), Si2 (foliar application with Si and fertigation EC 2.0 dS m-1), ASi2 (foliar application with *Arthrospira platensis* and Si and fertigation EC 2.0 dS m-1), C3.0 (foliar application with water and fertigation EC 3.0 dS m-1), A3.0 (foliar application with *Arthrospira platensis* and fertigation EC 3.0 dS m-1), Si3 (foliar application with Si and fertigation EC 3.0 dS m-1) , ASi3 (foliar application with Si and *Arthrospira platensis* and fertigation EC 3.0 dS m-1) treatments, C3.5 (foliar application with water and fertigation EC 3.5 dS m-1), A3.5 (foliar application with *Arthrospira platensis* and fertigation EC 3.5 dS m-1), Si3.5 (foliar application with Si and fertigation EC 3.5 dS m-1) and ASi3.5 (foliar application with Si and *Arthrospira platensis* and fertigation EC 3.5 dS m-1) treatments.. Different letters indicate significant differences between treatments at the P<0.05 level using the LSD test. Values are the means of 12 plants. Error bars show standard deviations.

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Figure 7. Shoot number (a) and leaves number (b) at the end of the trial for the C2 (foliar application with water and fertigation EC 2.0 dS m-1), A2 (foliar application with *Arthrospira platensis* and fertigation EC 2.0 dS m-1), Si2 (foliar application with Si and fertigation EC 2.0 dS m-1), ASi2 (foliar application with *Arthrospira platensis* and Si and fertigation EC 2.0 dS m-1), C3.0 (foliar application with water and fertigation EC 3.0 dS m-1), A3.0 (foliar application with *Arthrospira platensis* and fertigation EC 3.0 dS m-1), Si3 (foliar application with Si and fertigation EC 3.0 dS m-1) , ASi3 (foliar application with Si and *Arthrospira platensis* and fertigation EC 3.0 dS m-1) treatments, C3.5 (foliar application with water and fertigation EC 3.5 dS m-1), A3.5 (foliar application with *Arthrospira platensis* and fertigation EC 3.5 dS m-1), Si3.5 (foliar application with Si and fertigation EC 3.5 dS m-1) and ASi3.5 (foliar application with Si and *Arthrospira platensis* and fertigation EC 3.5 dS m-1) treatments.. Different letters indicate significant differences between treatments at the P<0.05 level using the LSD test. Values are the means of 12 plants. Error bars show standard deviations.

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Figure 8. Flowers number (a) and Internode length (b) at the end of the trial for the C2 (foliar application with water and fertigation EC 2.0 dS m-1), A2 (foliar application with *Arthrospira platensis* and fertigation EC 2.0 dS m-1), Si2 (foliar application with Si and fertigation EC 2.0 dS m-1), ASi2 (foliar application with *Arthrospira platensis* and Si and fertigation EC 2.0 dS m-1), C3.0 (foliar application with water and fertigation EC 3.0 dS m-1), A3.0 (foliar application with *Arthrospira platensis* and fertigation EC 3.0 dS m-1), Si3 (foliar application with Si and fertigation EC 3.0 dS m-1) , ASi3 (foliar application with Si and *Arthrospira platensis* and fertigation EC 3.0 dS m-1) treatments, C3.5 (foliar application with water and fertigation EC 3.5 dS m-1), A3.5 (foliar application with *Arthrospira platensis* and fertigation EC 3.5 dS m-1), Si3.5 (foliar application with Si and fertigation EC 3.5 dS m-1) and ASi3.5 (foliar application with Si and *Arthrospira platensis* and fertigation EC 3.5 dS m-1) treatments.. Different letters indicate significant differences between treatments at the P<0.05 level using the LSD test. Values are the means of 12 plants. Error bars show standard deviations.

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Figure 9. Leave area at the end of the trial for the C2 (foliar application with water and fertigation EC 2.0 dS m-1), A2 (foliar application with *Arthrospira platensis* and fertigation EC 2.0 dS m-1), Si2 (foliar application with Si and fertigation EC 2.0 dS m-1), ASi2 (foliar application with *Arthrospira platensis* and Si and fertigation EC 2.0 dS m-1), C3.0 (foliar application with water and fertigation EC 3.0 dS m-1), A3.0 (foliar application with *Arthrospira platensis* and fertigation EC 3.0 dS m-1), Si3 (foliar application with Si and fertigation EC 3.0 dS m-1) , ASi3 (foliar application with Si and *Arthrospira platensis* and fertigation EC 3.0 dS m-1) treatments, C3.5 (foliar application with water and fertigation EC 3.5 dS m-1), A3.5 (foliar application with *Arthrospira platensis* and fertigation EC 3.5 dS m-1), Si3.5 (foliar application with Si and fertigation EC 3.5 dS m-1) and ASi3.5 (foliar application with Si and *Arthrospira platensis* and fertigation EC 3.5 dS m-1) treatments.. Different letters indicate significant differences between treatments at the P<0.05 level using the LSD test. Values are the means of 12 plants. Error bars show standard deviations.

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Figure 10. N, P, S and Cl (a) and K, Ca, Mg and Na (b) concentration in the leaves (%) for the C2 (foliar application with water and fertigation EC 2.0 dS m-1), A2 (foliar application with *Arthrospira platensis* and fertigation EC 2.0 dS m-1), Si2 (foliar application with Si and fertigation EC 2.0 dS m-1), ASi2 (foliar application with *Arthrospira platensis* and Si and fertigation EC 2.0 dS m-1), C3.0 (foliar application with water and fertigation EC 3.0 dS m-1), A3.0 (foliar application with *Arthrospira platensis* and fertigation EC 3.0 dS m-1), Si3 (foliar application with Si and fertigation EC 3.0 dS m-1) , ASi3 (foliar application with Si and *Arthrospira platensis* and fertigation EC 3.0 dS m-1) treatments, C3.5 (foliar application with water and fertigation EC 3.5 dS m-1), A3.5 (foliar application with *Arthrospira platensis* and fertigation EC 3.5 dS m-1), Si3.5 (foliar application with Si and fertigation EC 3.5 dS m-1) and ASi3.5 (foliar application with Si and *Arthrospira platensis* and fertigation EC 3.5 dS m-1) treatments.. Different letters indicate significant differences between treatments at the P<0.05 level using the LSD test. Values are the means of 12 plants. Error bars show standard deviations.