

15 **1. INTRODUCTION**

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17 Bedding plant industry has been exponentially expanded around the world according to 18 significant costs decrease. The last has been related to higher bedding pot plant vield per unit 18 significant costs decrease. The last has been related to higher bedding pot plant yield per unit
19 areenhouse area through two grower currently decision-making: plug cell volume during nursery 19 greenhouse area through two grower currently decision-making: plug cell volume during nursery
20 and growing media quality for both nursery and pot cycle. Commercial profits have been related 20 and growing media quality for both nursery and pot cycle. Commercial profits have been related
21 to a decrease in plug cell volume [1] and the use of lower expensive growing media [2,3,4]. 21 to a decrease in plug cell volume [1] and the use of lower expensive growing media [2,3,4].
22 However, these business choices imply that plants will suffer different root restriction stresses 22 However, these business choices imply that plants will suffer different root restriction stresses
23 during most growing cycle. during most growing cycle.

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25 25 Usually, the 'root restriction syndrome' has been defined as a physical stress imposed on a root
26 system when plants are grown in small containers, which leads to a pronounced decrease in 26 system when plants are grown in small containers, which leads to a pronounced decrease in 27 both root and shoot growth at the transplant stage. The pre- and post-transplant effects of the 27 both root and shoot growth at the transplant stage. The pre- and post-transplant effects of the
28 container volume during nursery [5,6] have been extensively studied by our laboratory and we 28 container volume during nursery [5,6] have been extensively studied by our laboratory and we
29 found that growth restrictions would be closely related to endogenous cytokinin synthesis by 29 found that growth restrictions would be closely related to endogenous cytokinin synthesis by
30 roots. A limited plug cell volume gives a vertical root restriction when root apical meristem 30 roots. A limited plug cell volume gives a vertical root restriction when root apical meristem
31 **comes** to the bottom of the cell or the pot. At this moment, both primary root growth and root 31 comes to the bottom of the cell or the pot. At this moment, both primary root growth and root 32 branching decrease [7] and cytokinins, the main endogenous hormone synthesized by root 32 branching decrease [7] and cytokinins, the main endogenous hormone synthesized by root and approximation and ap apical meristems would decrease as well [8,9]. 34

35 On the other hand, when a growing media quality decreases, it changes their physical
36 properties [3,10], which generally results in decreased pore sizes, and would be taken into 36 properties [3,10], which generally results in decreased pore sizes, and would be taken into
37 account as an abiotic stress [4]. As pore size decreases, total porosity and air-filled porosity account as an abiotic stress [4]. As pore size decreases, total porosity and air-filled porosity 38 decrease as well. Bailey-Serres and Colmer [11] and Voesenek and Sasidharan [12] have 39 indicated that a lack of oxygen inhibits respiration, decrease metabolic plant adaptations to cope
40 with the hypoxic and anoxic conditions and resulting energy deficits, as well as change 40 with the hypoxic and anoxic conditions and resulting energy deficits, as well as change 41 anatomical and morphological adaptations to improve internal $O₂$ supply. These metabolic 41 anatomical and morphological adaptations to improve internal $O₂$ supply. These metabolic 42 changes would give the same effects on endogenous cytokinin synthesis as plug cell volume 42 changes would give the same effects on endogenous cytokinin synthesis as plug cell volume
43 [13.14]. $[13, 14]$.

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45 45 With the goal of maximizing bedding plant yield to identify the main limiting factor it is
46 imperative. Previous reports on ornamentals [15.16.17] and vegetables [18.19.20.21] have imperative. Previous reports on ornamentals $[15,16,17]$ and vegetables $[18,19,20,21]$ have 47 shown that nursery growth has a significant effect on post-transplant biomass accumulation.
48 The precise effects of combined plug cell volume and growing media quality on nursery have The precise effects of combined plug cell volume and growing media quality on nursery have 49 been recently indicated as well [22,23,24]. However, the simultaneously post-transplant
50 interactions between these two **stresses source** during both nursery and pot growth is lacking. interactions between these two **stresses source** during both nursery and pot growth is lacking.

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52 *Impatiens walleriana* (Hook.f.) is a commercially important year-round garden crop for landscape, and the first best-selling bedding plants in both developed and undeveloped 54 countries. Most *Impatiens* genotype produce a compact green foliage and covers itself with 55 extremely uniform growth habit and bright blooms. *Impatiens* F₁ genotypes prefer partial
56 sun/shade (8-25 mol photons m⁻² day⁻¹) [25]. Dry mass and flowering increase from 14 to 28°C sun/shade (8-25 mol photons m⁻² day⁻¹) [25]. Dry mass and flowering increase from 14 to 28°C
57 [26]. Plants only grow well with 100% evapotranspiration [27]. *I wallweriana* has been included 57 [26]. Plants only grow well with 100% evapotranspiration [27]. *I wallweriana* has been included in most research from our laboratory for the last decade.

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The aim of this work was to evaluate *I. walleriana* yield to the end of the pot growth stage when 61 four different pre-transplant cell volume and four pre or post-transplant growing media with
62 different physical properties were used. The hypothesis tested was that only one of the 62 different physical properties were used. The hypothesis tested was that only one of the 63 potentially negative stress source (pre-transplant cell volume or growing medium quality) is the 63 potentially negative stress source (pre-transplant cell volume or growing medium quality) is the main responsible for decreasing biomass accumulation at the post-transplant pot growing cycle. 65

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67 **2. MATERIALS AND METHODOLOGY**

68 **2.1 Plant Material**

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Experiment were carried out under a greenhouse at the Faculty of Agronomy, University of 71 Buenos Aires, Argentina (34° 35' 59"S, 58° 22' 23"W) from October 10th 2012 to December 9th 72013 and repeated from October 16th 2013 to December 15th 2014. 2013 and repeated from October 16th 2013 to December 15th 2014.

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74 *I. walleriana* 'Xtreme White' seeds (Goldsmith Inc., NY, USA) were germinated and grown in 75 $50-$, 128-, 288- and 512-cell plug tray⁻¹ (55.70, 17.37, 6.18 and 2.50 cm³ cell⁻¹ respectively) in 76 four different pre-transplant growing media as follows:

- 77
- 78 1) Klasmann 411® medium (Klasmann-Deilmann, GmbH, Germany): Canadian
79 Sohagnum peat moss-perlite-vermiculite (70/20/10 v/v/v) (K) 79 *Sphagnum* peat moss-perlite-vermiculite (70/20/10 v/v/v) (**K**)
80 2) *Sphagnum maguellanicum*-perlite (80/20 v/v) (**S**)
	- 80 2) *Sphagnum maguellanicum*-perlite (80/20 v/v) (**S**)
	-
- 81 3) River waste-perlite (80-20 v/v) (**R**)
82 4) Sphagnum maguellanicum-river was 82 4) *Sphagnum maguellanicum*-river waste-perlite (40-40-20, v/v/v) (**SR**).

83
84 When seedlings reached the transplant stage, they were transplanted into 1,200 cm³ pots filled
85 buth a post-transplant Klasmann 411® medium (Klasmann-Deilmann, GmbH, Germany), At the 85 with a post-transplant Klasmann 411® medium (Klasmann-Deilmann, GmbH, Germany). At the 86 same time, plants grown at the pre-transplant stage in a Klasmann 411[®] medium (Klasmann-
87 Deilmann, GmbH, Germany) were transplanted to 1,200 cm³ pots filled with the four different pre-87 Deilmann, GmbH, Germany) were transplanted to 1,200 cm³ pots filled with the four different pre-
88 transplant growing media tested, given 32 combinations of plug cell volume-growing media. transplant growing media tested, given 32 combinations of plug cell volume-growing media.

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91 91 **2.2 Cultivation and Meteorological Data**
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93 Plants were irrigated daily at saturation with high quality tap water using intermittent overhead
94 mist. Growing media **was** weekly fertilized with 1.0: 0.5: 1.0: 0.5 ($v/v/v/v$) N: P: K: Ca through the 94 mist. Growing media was weekly fertilized with 1.0: 0.5: 1.0: 0.5 (v/v/v/v) N: P: K: Ca through the 95 overhead irrigation water (Stage 2: 50 mg L^{-1} N; Stage 3-4: 100 mg L^{-1} N; pot: 150 mg L^{-1} N). overhead irrigation water (Stage 2: 50 mg L⁻¹ N; Stage 3-4: 100 mg L⁻¹ N; pot: 150 mg L⁻¹ N).

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97 Daily mean temperatures (22.26 to 25.06°C) and daily photosynthetic active radiation (4.24 to 98 5.03 mol photons m^{-2} day⁻¹) for the experiments were recorded with a HOBO sensor (H08-004-5.03 mol photons m^{-2} day⁻¹) for the experiments were recorded with a HOBO sensor (H08-004-
99 02) (Onset Computer Corporation, MA, USA) connected to a HOBO H8 data logger. The plants 99 02) (Onset Computer Corporation, MA, USA) connected to a HOBO H8 data logger. The plants 100 were arranged at a density of 25 plants m^2 , which avoided mutual shading. were arranged at a density of 25 plants m^2 , which avoided mutual shading.

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102 103 **2.3 Sampling and Growth Evaluations** 104

105 Samples of each substrate were collected at the beginning of the pot experiments (before 106 transplant to 1,200-cm³ pots) and total porosity, air-filled porosity, bulk density and container transplant to 1,200-cm³ pots) and total porosity, air-filled porosity, bulk density and container 107 capacity were determined according to Fonteno [28]. Data are indicated in Table 1 and show 108 significant physical properties differences in of the growing media tested.

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 Table 1: Physical properties for the growing media tested. K: [Canadian *Sphagnum* **peat (70%) + Perlite (20%) + Vermiculite (10%)], S: [***Sphagnum maguellanicum* **(80%) + Perlite (20%)], R: [River waste (80%) + Perlite (20%)], SR: [***Sphagnum maguellanicum* **(40%) + River waste (40%) + perlite (20%)]. Standard errors are indicated.**

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116 Plants were harvested at the transplant stage and at 15, 30, 45, and 60 days after transplanting. 117 Roots were washed and root, stem and leaf fresh weights (FW) were recorded. Dry weights 118 (DW) were obtained after drying roots, stems and leaves to constant weight at 80°C for 96 118 (DW) were obtained after drying roots, stems and leaves to constant weight at 80°C for 96
119 **hours**. The number of leaves was recorded, and each leaf area was determined using the hours. The number of leaves was recorded, and each leaf area was determined using the 120 ImageJ® (Image Processing and Analysis in Java) software. The number of stems and nodes 121 has been recorded as well. has been recorded as well.

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123 The relative rate of leaf area expansion (RLAE) was calculated as the slope of the regression of 124 the natural logarithm (In) of total leaf area versus time (in days). The rate of leaf appearance 124 the natural logarithm (ln) of total leaf area versus time (in days). The rate of leaf appearance
125 (RLA) was calculated as the slope of the number of fully expanded leaves versus time (in 125 (RLA) was calculated as the slope of the number of fully expanded leaves versus time (in 126 weeks). The specific leaf area on a FW basis (SLA) and leaf weight rate (LWR) were calculated 126 weeks). The specific leaf area on a FW basis (SLA) and leaf weight rate (LWR) were calculated
127 as the ratio between the area of the new individual leaf and leaf FW and the ratio between the 127 as the ratio between the area of the new individual leaf and leaf FW and the ratio between the 128 leaf DW and the total plant DW respectively at the end of the experiments. The relative growth 128 leaf DW and the total plant DW respectively at the end of the experiments. The relative growth 129 rate (RGR) was calculated as the slope of the regression of the In of whole plant DW versus 129 rate (RGR) was calculated as the slope of the regression of the In of whole plant DW versus 130 time (in days). $time (in days)$. 131

132 The mean net assimilation rate (NAR) and leaf area ratio (LAR) were calculated according to 133 Potter and Jones [29] as follows: Potter and Jones [29] as follows:

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NAR = \frac{k_w W_0 e^{k_w t}}{A_0 e^{k_a t}}
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LAR = \frac{k_w}{NAR}
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139 139 where W₀: extrapolated value of total DW (g) at time zero; k_w : RGR (day⁻¹); A₀: extrapolated 140 value of leaf area (cm²) at time zero: k_a: RLAE (day⁻¹): t: time (days) at the midpoint of the 140 value of leaf area (cm²) at time zero; k_a : RLAE (day⁻¹); t: time (days) at the midpoint of the 141 experimental period and e: base of the ln.

143 The allometric coefficients between root and shoot were calculated as the slope (**β**) of the 144 straight-line regression of the ln of the root DW versus the ln of the shoot DW. The Root: Shoot 145 ratio (at the end of the experiment) was performed as well. ratio (at the end of the experiment) was performed as well.

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148 148 **2.4 Statistical Analysis**

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150 150 The experimental design was a randomised factorial with three blocks of five single-pot 151 replications of each treatment combination (plug cell volume x growing medium x pre- and post-151 replications of each treatment combination (plug cell volume × growing medium × pre- and post-
152 transplant). Since there were no significant differences between the two experiments, they were 152 transplant). Since there were no significant differences between the two experiments, they were 153 considered together (n = 30). Data were subjected to three-way analysis of variance (ANOVA). 153 considered together (n = 30). Data were subjected to three-way analysis of variance (ANOVA).
154 STATISTICA 8 (StatSoft) software was used and the assumptions of the ANOVA were checked. 154 STATISTICA 8 (StatSoft) software was used and the assumptions of the ANOVA were checked.
155 Means were separated by Tukey's tests ($P \le 0.05$). Slopes from straight-line regressions of 155 Means were separated by Tukey's tests ($P \le 0.05$). Slopes from straight-line regressions of 156 MATR package [30]. RLA, RLAE, RGR and allometric values were tested using the SMATR package [30].

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159 **3. RESULTS** 160 **3.1. Biomass accumulation**

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162 162 Total fresh weight at the end of the pot growth cycle (60 days from transplant) was higher in 163 plants from 50-plug tray⁻¹ and decreased according cell **numbers** increased in all growing media plants from 50-plug tray⁻¹ and decreased according cell **numbers** increased in all growing media
164 tested. Anyway, growing media significantly changed post- transplant biomass accumulation on 164 tested. Anyway, growing media significantly **changed** post- transplant biomass accumulation on
165 a FW base as during the pre- as the post-transplant stage, but especially during nursery. The a FW base as during the pre- as the post-transplant stage, but especially during nursery. The 166 higher FW was found in plants grown in **R-** and **S-**growing media (Fig. 1A). When the mean 167 acris and the mean to and the mean the mea 167 aerial FW was plotted against the mean root FW (Fig. 1B), a positive correlation was found (r^2 = 168 = 0.661; P < 0.001). $0.661; P < 0.001$).

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171 **Figure 1. Total fresh weight at the end of the experiment (60 days from transplanting) for** *Impatiens walleriana* **plants grown in four plug cell volumes (50-, 128-, 288-, and 512-cell tray-1) at the pre-transplant stage and four growing media at the pre- or post-transplant stage. Bars indicate standard errors and vertical line indicate least significant differences** (LSD). Panel B. Relationships between **shoots and roots** FW according to four plug cell **volumes (50-, 128-, 288-, and 512-cell tray-1) at the pre-transplant stage and four growing media at the pre- (full symbols) or post-transplant stage (empty symbols). For substrate abbreviations see Table 1. F: -◊; R: ■-□; S: ●-○; SR: ▲-∆. The straight-line regression** 179 was: Shoot FW = 6.65 Root FW + 6.73 (r^2 = 0.661). The probability of the slope being zero 180 was P < 0.001. was P < 0.001.

3.2. Leaf area expansion

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185 Total leaf area at the end of the experiment was once again higher in plants from 50-plug cells
186 and in those grown at different growing media during the pot grow stages (Fig. 2A). Although 186 and in those grown at different growing media during the pot grow stages (Fig. 2A). Although
187 there were significant differences in individual leaf area according to different pre-transplant cell there were significant differences in individual leaf area according to different pre-transplant cell 188 volume and pre- or post-transplant growing media, changes were smaller than in total leaf area
189 (Fig. 2B). (Fig. 2B).

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192 **Figure 2. Total (A) and individual (B) leaf area at the end of the experiments (60 days from** 193 **transplanting) for** *Impatiens walleriana* **plants from four plug cell volumes (50-, 128-, 288-,** and 512-cell tray⁻¹) at the pre-transplant stage and four growing media at the pre- or post-

195 **transplant stage. Bars indicate standard errors and vertical line indicate least significant** 195 **transplant stage. Bars indicate standard errors and vertical line indicate least significant** differences (LSD). For substrate abbreviations see Table 1.

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198 **Table 2: Changes in RLA, RLAE and SLA for** *Impatiens walleriana* **plants from four plug cell volumes (50-, 128-, 288-, and 512-cell tray-1** 199 **) at the pre-transplant stage and four growing media at the pre- or post-transplant stage. Different lower-case letters indicate significant differences (P< 0.05) between pre-transplant plug cell volumes, while different capital letters indicate significant differences (P < 0.05) between pre- and post- growing media. For substrate abbreviations see Table 1. The probability of the RLA and RLAE** slopes being zero was P < 0.001.

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207 207 The higher plug cell volume the higher RLA and RLAE but the lower SLA (leaves has a higher 208 thickness). The response to a change in growing media quality at the pre- or post-transplant 208 thickness). The response to a change in growing media quality at the pre- or post-transplant 209 changed according pre-transplant plug cell volume and growing media tested in the same way 209 changed according pre-transplant plug cell volume and growing media tested in the same way
210 that total leaf area (Table 2). that total leaf area (Table 2).

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212 212 Stem numbers per plant showed significant differences in plants grown at nursery in different 213 both plug cell volume and growing media; they were positively stimulated by a change in post-213 both plug cell volume and growing media; they were positively stimulated by a change in post-
214 transplant growing media in plants from K-growing media. An inverse or no significant result 214 transplant growing media in plants from K-growing media. An inverse or no significant result
215 was found when S-, R- or SR-growing media were used at the post-transplant stage (Fig. 3A). A 215 was found when S-, R- or SR-growing media were used at the post-transplant stage (Fig. 3A). A
216 similar response from the number node per plant was found (Fig. 3B). similar response from the number node per plant was found (Fig. 3B).

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230 **Fig. 3. Stem (A) and node (B) number at the end of the experiments (60 days from** 231 **transplanting) for** *Impatiens walleriana* **plants from four plug cell volumes (50-, 128-, 288-,** and 512-cell tray⁻¹) at the pre-transplant stage and four growing media at the pre- or post-
233 **but a transplant stage. Bars indicate standard errors and vertical line indicate least significant** 233 **transplant stage. Bars indicate standard errors and vertical line indicate least significant** differences (LSD). For substrate abbreviations see Table 1.

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237 **3.3. Dry weight accumulation**

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239 Due there is no DW significant differences between treatments (data not shown), the traditional 240 arowth analysis approach would be performed. During the experiments. RGR values were growth analysis approach would be performed. During the experiments, RGR values were 241 significantly different from plants grown in different plug cell volume although data from post-
242 transplant growing media were higher than those from plants grown at the same growing media 242 transplant growing media were higher than those from plants grown at the same growing media
243 at the pre-transplant stage. When RGR was separating from their 'physiological' (NAR) and at the pre-transplant stage. When RGR was separating from their 'physiological' (NAR) and 244 'morphological' (LAR) components, NAR decreased according plug cell volume decrease with 245 significant differences between growing media tested and time (lesser pre-transplanted plants 246 than post-transplanted ones). Quite opposite responses were found for LAR (Table 3). than post-transplanted ones). Quite opposite responses were found for LAR (Table 3).

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 Table 3: Changes in RGR, NAR and LAR for *Impatiens walleriana* **plants from four plug cell volumes (50-, 128-, 288-, and 512-cell tray-1** 250 **) at the pre-transplant stage and four growing media at the pre- or post-transplant stage. Different lower-case letters indicate significant differences (P< 0.05) between pre-transplant plug cell volumes, while different capital letters indicate significant differences (P < 0.05) between pre- and post- growing media. For substrate abbreviations see Table 1. The probability of the RGR slope being zero was P < 0.001.**

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261 inverse relationship between RGR and LAR (r^2 = 0.191 and 0.328) (Fig. 4B).

²⁶³ ²⁶⁴ **Figure 4: The net assimilation rate (NAR) (A) and the leaf area ratio (LAR) (B) related to** 265 **the relative growth rate (RGR) for** *Impatiens walleriana* **plants from four plug cell volumes (50-, 128-, 288-, and 512-cell tray-1** 266 **) at the pre-transplant stage and four growing media at** 267 the pre- or post-transplant stage. The probability of the NAR and LAR <mark>slopes</mark> being zero
268 was P < 0.001. For substrate abbreviations see Table 1. F: ●-○; R: ■-□; S: ♦-◊; SR: ▲-Δ. 268 **was P < 0.001. For substrate abbreviations see Table 1. F: ●-○; R: ■-□; S: -◊; SR: ▲-∆.** 269 The straight-line regressions were: NAR_{Pre} = 189.25 RGR - 1.67 (r^2 = 0.656); NAR_{Post} = 270 **739.27 RGR - 29.99 (** \tilde{r}^2 **= 0.635); LAR_{Pre} = - 1,596.20 RGR + 132.07 (** \tilde{r}^2 **= 0.191) and LAR_{Pre} = -**271 **5,572.40 RGR + 368.89 (r²= 0.328).**

272 273 Positive relationships between RLAE (r^2 = 0.645 P < 0.001) (Fig. 5A), RLA (r^2 = 0.627 P < 274 0.001) (Fig. 5B), RGR (r^2 = 0.665 P < 0.001) (Fig. 5C), NAR (r^2 = 0.602 P < 0.001) (Fig. 5D), 275 and root DW were found. The higher values were those from plants grown in different growing
276 media at the post-transplant stage. media at the post-transplant stage. 277

 Fig. 5. Relationship between RLAE (A), RLA (B), RGR (C), NAR (D) and root dry weight (RDW) for *Impatiens walleriana* **plants from four plug cell volumes (50-, 128-, 288-, and 512-cell tray-1** 282 **) at the pre-transplant stage and four growing media at the pre- (full symbols) or post-transplant stage (empty symbols). The straight-line regressions were: RLAE = 0.067**
284 **RDW + 0.023 (r² = 0.645 P < 0.001). RLA = 0.735 root DW + 0.44 (r² = 0.627 P < 0.001). RGR = RDW + 0.023 (r² = 0.645 P < 0.001), RLA = 0.735 root DW + 0.44 (r²** 284 **= 0.627 P < 0.001), RGR = 0.036 RDW + 0.03 (r² = 0.665 P < 0.001), NAR = 11.86 RDW + 2.69 (r² = 0.602 P < 0.001). The probability of the slopes being zero was P < 0.001. F: -◊; R: ■-□; S: ●-○; SR: ▲-∆.**

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291 **3.4. Photo assimilates partitioning**

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293 The higher plug cell volume the lower β coefficient for root: shoot allometries for all growing 194 media at the pre-transplant, which showed a higher photo assimilates partitioning to roots. The 294 media at the pre-transplant, which showed a higher photo assimilates partitioning to roots. The 295 same B response pattern was found during the post-transplant but absolute values were even 295 same β response pattern was found during the post-transplant but absolute values were even
296 lower than for the pre-transplant growing media. An increase in root: shoot ratio according plug lower than for the pre-transplant growing media. An increase in root: shoot ratio according plug 297 cell volume decrease were found as well with significant differences between growing media
298 (Table 4). $(Table 4)$.

299
300 **Table 4: Changes in allometric relationships between roots and shoots for** *Impatiens walleriana* plants from four plug cell volumes (50-, 128-, 288-, and 512-cell tray⁻¹) at the 302 pre-transplant stage. **pre-transplant stage and four growing media at the pre- or post-transplant stage. Different lowercase letters indicate significant differences (P< 0.05) between pre- transplant plug cell volumes, while different capital letters indicate significant differences (P < 0.05) between pre- and post- growing media. For substrate abbreviations see Table 1. The probability of the slopes being zero was P < 0.001.** 307

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310 **4. DISCUSSION**

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312 312 In a recent previous report [24], we have found that, in *I. walleriana* seedlings, the abiotic stress imposed by the growing medium quality during nursery had a higher effect on biomass 314 accumulation (on both fresh and dry base), leaf area expansion and photo assimilates partitioning than plug cell volume and constitute an interactive process associated with cytokinin 316 synthesis. However, this novelty approach did not exclude the plug cell volume involvement as 317 a limiting abiotic stress source during other parts of the *I. walleriana* growth cycle. In this context 317 a limiting abiotic stress source during other parts of the *I. walleriana* growth cycle. In this context we have evaluated pot biomass accumulation of this bedding plant to four growing media at 319 both nursery and pot stage in plants propagated in four different plug cell volumes.

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321 A 'root restriction' syndrome related to either plug cell volume or growing media quality is an 322 exogenous signal, which let plants to sense the volume space and decrease or increase root exogenous signal, which let plants to sense the volume space and decrease or increase root 323 system accordingly [31,32]. Having in mind that noot is a major source of cytokinins [33], which
324 boontrol the source of biomass accumulation such as the shoot apical meristem [34]. the 324 control the source of biomass accumulation such as the shoot apical meristem [34], the 325 similarity between *l. walleriana* plants grown in the best growing conditions and exogenous 325 similarity between *I. walleriana* plants grown in the best growing conditions and exogenous 326 cytokinin-sprayed plants [23,24,35], it is not unexpected. Our results from Fig. 5, which shown a positive relationships between the most growing parameters performed (RLAE, RLA, RGE, 328 NAR) according to a root dry weight increases are in agreement with this previous reports. On
329 the other hand, a positive relationship between **shoots and roots** fresh weight (there were not 329 the other hand, a positive relationship between **shoots and roots** fresh weight (there were not 330 significant differences between fresh and dry weight) was found (Fig. 1B) in agreement with 330 significant differences between fresh and dry weight) was found (Fig. 1B) in agreement with 331 previous reports [4,15,17,21,22,23,24]. previous reports [4,15,17,21,22,23,24].

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333 Although growing media performed through a high quality *Sphagnum sp*. peat base (Klasman® 334 commercial growing media) has been indicated as the best for pot plants [1], *I. walleriana* 335 previous results indicated that better growing media for this bedding plant can be found [36].

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337 Results showed significant fresh weight changes at the end of the experiments (Fig. 1A)
338 according a decrease in plug cell volume and a change in growing media quality, which are in 338 according a decrease in plug cell volume and a change in growing media quality, which are in
339 agreement to RGR changes (Table 3). Methodology usually used to describe changes in 339 agreement to RGR changes (Table 3). Methodology usually used to describe changes in 340 biomass accumulation on both fresh and dry weight included: (i) stems appearance: (ii) leaf 340 biomass accumulation on both fresh and dry weight included: (i) stems appearance; (ii) leaf
341 area expansion: (iii) photosynthetic capacity and: (iy) photo assimilate partitioning. In 341 area expansion; (iii) photosynthetic capacity and; (iv) photo assimilate partitioning. In 342 ornamental bedding plants, additional traits such as tolerance to biotic and abiotic stresses and 342 ornamental bedding plants, additional traits such as tolerance to biotic and abiotic stresses and 343 aesthetics must be included [37]. aesthetics must be included [37].

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345 Stem branching is an important aspect to consider in ornamental bedding plants because it takes
346 Dart as the biomass accumulation as the aesthetic appearance. Results from Fig. 3 indicated that 346 part as the biomass accumulation as the aesthetic appearance. Results from Fig. 3 indicated that 347 both shoot number and node number decreased according to plug cell volume decrease, which 347 both shoot number and node number decreased according to plug cell volume decrease, which
348 indicates a desirable commercial ideotype with a lower branching and compact growth habit. 348 indicates a desirable commercial ideotype with a lower branching and compact growth habit.
349 However, growing media quality changes the impact on the response to different pre-transplant plug 349 However, growing media quality <mark>changes</mark> the impact <mark>on</mark> the response to different pre-transplant plug
350 cell volume. cell volume.

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352 Aesthetically, total leaf area is the main trait related to plant quality for commercial acceptation
353 of ornamentals and it determines the time of plant sale and at the same time. leaves are the 353 of ornamentals and it determines the time of plant sale and at the same time, leaves are the 354 plant organs responsible to light interception. In physiological terms, it implies to expand leaf plant organs responsible to light interception. In physiological terms, it implies to expand leaf 355 area at the higher growth rate which included both individual leaf sizes and leaf numbers. Data
356 from Fig. 2A shown that the total leaf area, with minor effects on individual leaf area (Fig. 2B) 356 from Fig. 2A shown that the total leaf area, with minor effects on individual leaf area (Fig. 2B)
357 was significantly affect by both plug cell volume and growing media guality. Three growth was significantly affect by both plug cell volume and growing media quality. Three growth 358 parameters can be used to characterize leaf area development: (i) RLA, which is an estimator of 359 leaf initiation and plastochron length, (ii) RLAE, which let to quantify leaf expansion and, (iii) leaf initiation and plastochron length, (ii) RLAE, which let to quantify leaf expansion and, (iii) 360 SLA, which characterize leaf thickness. Data from Table 2 showed that a decrease in plug cell
361 volume and a change in growing media quality decreased RLA and RLAE while increased SLA. 361 volume and a change in growing media quality decreased RLA and RLAE while increased SLA.
362 These results implies that the changes in total leaf area are mainly related to the meristematic 362 These results implies that the changes in total leaf area are mainly related to the meristematic 363 shoot apex capacity to initiate and to expand leaf primordia [38]. Both processes are mediated 364 by the down requiation of *KNOTTED* and *WUSCHEL* genes [39] associated to a high cytokinin: 364 by the down regulation of *KNOTTED* and *WUSCHEL* genes [39] associated to a high cytokinin: 365 low gibberellin ratio [40]. On the other hand, the lower SLA the higher leaf thickness, which it is a pre-requisite for a high photosynthetic rate [41]. In this way, Gandolfo et al. [9] found positive 367 relationships between leaf thickness, intercellular spaces and NAR in *I. walleriana* rootrestricted plants. When the mesophyll thickness of the leaf is increased, the maximum 369 photosynthetic rate increased as well. This probably explains the strong relationship between 370 NAR and mesophyll thickness.

371

372 Variation in RGR has the result of two key traits: the 'physiological component' NAR and the 373 ('morphological component' LAR. RGR, which ultimate quantify biomass accumulation. is 373 'morphological component' LAR. RGR, which ultimate quantify biomass accumulation, is 374 greatly influenced by photosynthetic efficiency. Although the higher the plug cell volume the 375 higher the RGR and NAR, growing media quality at the post-transplant stage increased both higher the RGR and NAR, growing media quality at the post-transplant stage increased both 376 growth parameters (Table 3). Shipley [42] indicated that, in general, NAR was the best general 377 predictor of variation in RGR, in agreement with our results from Fig. 4. Root restrictions often
378 depresses photosynthetic capacity [44] and decreased energy synthesis [45]. The positive 378 depresses photosynthetic capacity [44] and decreased energy synthesis [45]. The positive 379 relationships between NAR and RGR (Figure 4A) are in agreement with Shi et al. [44.45]. 379 relationships between NAR and RGR (Figure 4A) are in agreement with Shi et al. [44.45].

380

381 Root-restricted plants change photo assimilates partition as a response to abiotic stresses
382 (Table 4). At the end of the experiments, the higher root restriction the higher root: shoot ratio. 382 (Table 4). At the end of the experiments, the higher root restriction the higher root: shoot ratio.
383 Root: shoot allometries let to explain these results because showed a higher photo assimilates 383 Root: shoot allometries let to explain these results because showed a higher photo assimilates
384 Dartitioning to roots (lower ß coefficients) during the greater part of the experiments in root-384 partitioning to roots (lower β coefficients) during the greater part of the experiments in rootlimited treatments.

386
387

387 As opposed to a previous report [23], which indicate that that growing media quality would be a
388 more limited factor than plug cell volume for *I. walleriana* seedlings during nursery, our results 388 more limited factor than plug cell volume for *I. walleriana* seedlings during nursery, our results 389 showed that both abiotic stresses would be interactive restricting technological factors during the 390 post-transplant pot stage. post-transplant pot stage.

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392

393 **5. CONCLUSIONS** 394

395 The effect of an abiotic stress and the relationships between multiples stress sources is not the 396 same according to the plant growth stage. In this context, different *I. walleriana* growth 396 same according to the plant growth stage. In this context, different *I. walleriana* growth 397 besponses to both plug cell volume and growing media found in our experiments would not be
398 bunexpected results but to extend to other ornamental bedding plants and to perform a 398 unexpected results but to extend to other ornamental bedding plants and to perform a
399 commercial suggestion much more research must be reguired. commercial suggestion much more research must be required.

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