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Chlorophyll a fluorescence analysis reveals divergent photosystem II responses to saline, alkaline and saline-alkaline stresses in the two *Lotus japonicus* model ecotypes MG20 and Gifu-129.

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Abstract:	<p>Saline and alkaline stresses affect more than 10% of the World's arable land, limiting agricultural production. Salt-induced stress may affect the Photosystem II (PSII) function, altering fluorescence emission. Therefore, changes in fluorescence are used to quantify and analyze abiotic stress responses in plants. So far, no study has been focused on the response of PSII to saline, alkaline and saline-alkaline stresses in the model legume <i>Lotus japonicus</i>. For the saline, alkaline and saline-alkaline treatments, plants of the <i>L. japonicus</i> ecotypes MG20 and Gifu-129 were cultivated in sand with nutrient solution added with NaCl and NaHCO₃ in different proportions. Growth, gas exchange, and chlorophyll a fluorescence transient kinetic and OJIP parameters were measured, and chlorophyll a and b determined. The analysis of the kinetic of chlorophyll a fluorescence showed that NaCl-derived stress sources affect the photochemical events in PSII in both ecotypes, being this effect more evident under higher pH condition, whereas alkalinity per se has a mild or no effect on these events. The saline-alkaline stress induced a more severe effect on Gifu B-129, compared with Miyakojima MG20, whereas NaCl improved primary photochemistry in MG20. Our results allow us to accept the hypothesis that both ecotypes deploy differential responses under the three stressful treatments and that the saline-alkaline stress</p>	

causes higher damage levels than saline and alkaline stresses alone in relation with structures and sub-processes of the PSII.

Cover letter

Ms. Ref. No.:ACPP-D-18-00793

Dear Przemyslaw Wojtaszek,

Editor-in-chief

Please find attached a revised version of our manuscript “Chlorophyll *a* fluorescence analysis reveals divergent Photosystem II responses to saline, alkaline and saline-alkaline stresses in the two *Lotus japonicus* model ecotypes MG20 and Gifu-129.”

As you requested in your email, we have re-written specific passages in the Introduction and Discussion sections. We hope that changes performed will make the Similarity index acceptable now.

We have noticed that the software to screen for plagiarism was applied to the whole document, which included the response to the reviewers, affiliations and cites. We wonder if matches from those parts, and those from Materials and methods, where there are paragraphs taken from our previous works (which were not modified, following your recommendation) are being taken into consideration. Please, let us know if they need to be changed also.

The present work is not being considered for publication by another journal.

All authors have agreed to publication

We shall look forward to hearing from you at your earliest convenience.

Yours sincerely,

Ana Menéndez

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Title page

-Title. Chlorophyll *a* fluorescence analysis reveals divergent Photosystem II responses to saline, alkaline and saline-alkaline stresses in the two *Lotus japonicus* model ecotypes MG20 and Gifu-129.

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2 Abstract
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7 Saline and alkaline stresses affect more than 10% of the World's arable land, limiting agricultural
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9 production. Salt-induced stress may affect the Photosystem II (PSII) function, altering fluorescence
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11 emission. Therefore, changes in fluorescence are used to quantify and analyze abiotic stress
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13 responses in plants. So far, no study has been focused on the response of PSII to saline, alkaline
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19 with nutrient solution added with NaCl and NaHCO₃ in different proportions. Growth, gas exchange,
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21 and chlorophyll *a* fluorescence transient kinetic and OJIP parameters were measured, and chlorophyll
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23 *a* and *b* determined. The analysis of the kinetic of chlorophyll *a* fluorescence showed that NaCl-
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25 derived stress sources affect the photochemical events in PSII in both ecotypes, being this effect
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29 events. The saline-alkaline stress induced a more severe effect on Gifu B-129, compared with
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31 Miyakojima MG20, whereas NaCl improved primary photochemistry in MG20. Our results allow us
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33 to accept the hypothesis that both ecotypes deploy differential responses under the three stressful
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35 treatments and that the saline-alkaline stress causes higher damage levels than saline and alkaline
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37 stresses alone in relation with structures and sub-processes of the PSII.
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42 Keywords
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45 Alkalinity; Chlorophyll *a* fluorescence; *Lotus japonicus*; OJIP transient; Photosynthesis; Salinity
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56 Introduction
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1 Saline stress represents a soil condition where neutral salts (NaCl or Na₂SO₄) predominates,
2 whereas alkaline stress is mainly related to the occurrence of alkaline salts (Na₂CO₃ or NaHCO₃; Yang
3 et al. 2007). Both salt stresses disturb more than 10% of the world's cultivable area, hindering
4 agricultural outputs (Läuchli and Lüttge 2002). Negative effects of saline stress include reduction of
5 water uptake from soil (Munns 2002) and Na⁺ or Cl⁻ build-up within cells, leading to enzymatic activity
6 inhibition, all altering photosynthetic and energetic processes (Yeo 1998; Tester and Davenport 2003).
7 Alkaline salts may cause deficiency of nutrients such as phosphorus, iron and zinc (Clark 1982;
8 Marschner 1995), inhibiting photosynthesis and plant growth, besides altering balances of reactive
9 oxygen species and causing cell dying (Kukavica et al. 2013). Fluctuating neutral to alkaline salt
10 proportions often co-exist according to the soil (Shi and Wang 2005; Li et al. 2010). Several authors
11 have reported that when mixed, saline and alkaline stresses present synergistic detrimental actions on
12 plant development (e.g.: Shi and Sheng 2005; Paz et al. 2012; Paz et al. 2014; Vu et al., 2015; Li et al.,
13 2017; Kumar et al., 2018; Jia et al., 2019)

14 Salt stress often limits the photosynthesis (Ashraf and Harris 2013). In the long term, salinity-
15 induced reductions in the photosynthetic activity may result from decreases in the chlorophyll (Chl) and
16 carotenoid contents (Parida and Das 2005; Munns and Tester 2008; Duarte et al. 2013), increased
17 stomatal closure (Kyle et al. 1987) and/or non-stomatal restrictions, such as electron transport chain
18 disruption (Parida and Das 2005; Chaves et al. 2009; Xiang et al. 2016), among other factors. Indeed,
19 several stress sources may affect the Photosystem II (PSII) function, which alters fluorescence
20 emission. Therefore, changes in fluorescence are used to assess and analyze abiotic stress responses
21 in plants (Demetriou et al. 2007; Kalaji et al. 2011; Mathur et al. 2013; Zushi and Matsuzoe 2017).

22 In plants, fluorescence is primarily produced by Chl *a* integrating the PSII antenna complexes.
23 Its detection and analysis constitutes one of the most informative approaches to monitor the
24 photosystem II (PSII) functioning. Changes in fluorescence are related to processes occurring within
25 and around the PSII reaction centers, which reveal alterations in the acceptors redox balance and the
26 energy quantum yield (Goltsev et al. 2016). After a dark-adapted plant has been illuminated, the
27 intensity of Chl *a* fluorescence varies in a typical way. This variation in fluorescence intensity over time
28 is acknowledged as fluorescence transient and may be represented by a polyphasic curve (or Kautsky
29 curve, Papageorgiou 1975). This curve shows a phase of rapid increase presenting four steps
30 symbolized as O-J-I-P (Strasser et al. 1995). The O-J-I-P test gives clues on the condition of quinone
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1 A (Q_A), quinone B (Q_B) and plastoquinone (PQ) pools (Strasser and Govindjee 1992). The rise of Chl
2 a fluorescence from its minimal level "O" (the minimum value of fluorescence, F_0) to a "J" level (or F_J)
3 occurs at about 2 ms as result of Q_A reduction by PSII; this is followed by a fluorescence rise to the "I"
4 level (or F_I) at about 30 ms, by cause of the filling up of the quinone pool (Q_A and Q_B); finally a rise from
5 the "I" level to the "P" level occurs (the maximum value of the fluorescence, F_M), due to congestion of
6 electron traffic at the acceptor side of PSI.
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11 *Lotus japonicus* (Leguminosae) has been broadly used as model plant to study many important
12 physiological aspects during adaptation to salt stress, whose genome has been sequenced, providing
13 numerous tools for genomic/genetic research (Handberg and Stougaard 1992; Sato et al. 2006).
14 Several authors have compared the most widely employed genotypes of this species, Gifu B-129 and
15 Miyakojima MG20 (MG20) by using classic physiology and/or omic tools, regarding their tolerance and
16 response to saline (Melchiorre et al. 2009; Sánchez et al. 2008; 2011; 2012) and alkaline stresses
17 (Babuín et al. 2014; Campestre et al. 2016; Bordenave et al. 2017), showing that the first ecotype
18 deploys a higher sensitivity level than the second ecotype to both stresses. In 2016, Campestre and
19 collaborators registered a decline in PSII performance in plants of several *L. japonicus* ecotypes grown
20 under alkaline condition. However, no study has been undertaken to compare Gifu B-129 and MG20
21 responses to combined saline-alkaline (S-A) estres, nor has any study been directed on those
22 constituents of the PSII, which are substantially altered by saline, alkaline and S-A stresses in these
23 ecotypes.
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38 Our hypothesis are: 1- PSII is vulnerable to salt stress in both ecotypes, 2- Gifu B-129 and
39 MG20 deploy different responses to S-A stress, 3- the S-A stress causes higher damage levels than
40 separated saline and alkaline stresses. To test this, we compared the impacts of three distinct classes
41 of salt-induced stresses, alkaline, saline and S-A, on plant growth, net photosynthesis rate, stomatal
42 conductance, Chl contents and specific Chl fluorescence parameters in the two model *L. japonicus*
43 ecotypes.
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54 Materials and methods

57 *Plant material and growth conditions.*

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L. japonicus seeds from MG20 and Gifu B-129 ecotypes were scarified with concentrated sulfuric acid (98%) 3 min, washed ten times with sterile distilled water and sown in Petri dishes containing water-agar (0.8%). Plates were incubated in a growth chamber at 30 °C in darkness until germination, 7 days after sown. Each resultant seedling was transferred to a 8 × 20 cm (diam x length) cylindrical pot containing washed sand mix (50% fine/50% coarse sand; pH 7.0; E.C. = 0.05 mS cm⁻¹) and irrigated with 0.5 x Hoagland's nutrient solution (Hoagland and Arnon 1950). Transferred plants were cultured in a growth chamber with a 16/8 h photoperiod at 24 °C/21 °C ± 2 °C (day/night) and 55/65 ± 5% relative humidity and 250 μmol photons m⁻² s⁻¹ light intensity provided by Gro-lux fluorescent lamps (F 40W), until the end of all experiments. A drip irrigation system (9001 Digital Watering Timer Weekly Program, ELGO®, www.elgo.co.il; flow rate = 6.25 ml/h) was used according to Paz et al. (2012).

Experimental design and stress treatment

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Experiments were performed according to a completely randomized design of one factor (stress) and four levels: control, saline, alkaline and S-A stresses.

The 0.5x Hoagland solution was used as the nutrients source for all stressful and control conditions. The separate alkaline and saline conditions in a pot were created by adding respectively, 10 mM NaHCO₃ and 100 mM NaCl to the nutrient solution. For the S-A treatment, 90 mM NaCl + 10 mM NaHCO₃ were added, thus obtaining a stress solution with the same Na⁺-derived EC but a higher alkalinity than that of the saline stress treatment. Control treatment consisted of plants irrigated with 0.5x Hoagland solution without NaCl or NaHCO₃. The pH and EC of irrigation solutions were monitored every 3 days with a combined pH meter/conductimeter (HI 255; Hanna Instruments, Padova, Italy) and maintained at 5.8/1.2, 5.8/11.0, 8.0/1.9 and 8.0/11.0 pH units/dS.m⁻¹, for control, saline, alkaline and S-A conditions, respectively. In the case of alkaline treatment, 8-day-old seedlings received the final salt concentration: 10 mM NaHCO₃. In order to avoid any osmotic shock in the saline and S-A treatments, plants were first subjected to acclimation. For this, 8-day-old seedlings initially received 30 mM NaCl (saline), and 20 mM NaCl + 10 mM NaHCO₃ (S-A). Then, NaCl concentration was stepwise increased during 1 week until reaching the final treatments concentrations. After acclimation, fifteen-day old plants were grown under their respective treatments for further 20 days. Gas exchange parameters, Chl *a* fluorescence transients and chlorophyll contents were measured in one month-old plants, on intact, fully

1 expanded leaves, which were basal to the internode immediately below the apical bud. Gas exchange
2 and Chl a fluorescence transients were measured at midday. Plants were harvested at the age of 35
3 days, splitted into shoots and roots and kept for further determinations.
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6 There were six plants per treatment for each measured parameter (n = 6), and the experiment
7 was performed twice. Only most representative results are shown.
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10 11 *Evaluation of Gifu B-129 and MG20 tolerance.*

12 The tolerance of MG20 and Gifu B-129 genotypes to the different stress sources was evaluated
13 on the base of the plant growth, in terms of total dry matter production. For this, roots and shoots were
14 dried at 60 °C until constant weight.
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22 *Measurement of gas exchange parameters*

23 Net photosynthesis rate (Pn), stomatal conductance (Gs), transpiration rate (E) and internal
24 CO₂ (Ci) were measured under light saturation (1500 μmol photons m⁻² s⁻¹ illumination, LED light (peak
25 Wavelength: 625 nm), ambient carbon dioxide (400 ppm, average) at 24 °C, using a portable
26 photosynthesis system (TPS-2 Portable Photosynthesis System, MA, USA). Data was collected at
27 midday, 2 min to average steady-state.
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38 *Measurement of Chl a fluorescence transient kinetic and OJIP parameters*

39 Non-invasive Chl fluorescence fast-transient test (OJIP test) was conducted with a portable Chl
40 fluorometer (PocketPEA v.1.1, Hansatech Instruments, Ltd., UK) at room temperature according to
41 Babuin et al. (2014). Blade sections of intact leaves were covered with a leaf clip to adapt them to
42 darkness for 20 min and then exposed for 3 s to 3500 μmol photons m⁻² s⁻¹ (637 nm peak wavelength)
43 and Chl a fluorescence was recorded. The fluorescence data was processed by PEA plus software
44 (Hansatech Instruments, U.K.) to obtain the OJIP parameters. Measurements were performed on 6
45 plants per treatment for each ecotype. A summary of OJIP parameters used in this study is shown in
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58 *Chlorophyll a and b determination*

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1 Leaves were harvested and stored at -80 °C until use. For pigments extraction, 40 mg of plant
2 material grounded in liquid nitrogen was shaken in 100% acetone (4 °C, overnight). The extract was
3 cold centrifuged and the supernatant removed. Measurements were made at wavelength 663 nm (Chl
4 a) and 647 nm (Chl b) in a spectrophotometer (Perkin Elmer Lambda 25 UV/VIS spectrometer), and
5 pigments concentration calculated according to Lichtenthaler et al. (1987).
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10 11 *K⁺, Na⁺ and Cl⁻ determinations*

12 The K⁺ and Na⁺ contents were extracted from shoots with 100 mM HCl and estimated by
13 standard flame photometry according to Chen et al. (2001). Briefly, chloride was determined by a
14 thiocyanate-Hg-based colorimetric reaction. For this, 12.5 mg of powdered dry plant material was
15 extracted in 0.5 ml of a solution containing H₂O₂ (30%):concentrated HNO₃:isoamyl alcohol:H₂O at
16 1:1:0.08:7.9 (v/v), incubated at room temperature for 15 min, diluted to 5 ml with Milli-Q water, and
17 vigorously agitated in a Vortex. Then, 1.5 ml of the extraction mixture was centrifuged (10,000 rpm, 5
18 min) and the supernatant transferred to another Eppendorf tube. The colorimetric reaction solution
19 contained polyethylene glycol dodecyl ether–water (Brij 35®, 4%):mercuric thiocyanate (4.17 g/l
20 methanol):(NO₃)₃Fe (202 g/l Milli-Q water plus 21 ml concentrated HNO₃): Milli-Q water at 0.05:15:15:70
21 (v/v). One milliliter of this reaction was added to 320 µl of the supernatant (control treatment). In the
22 case of saline treatments, 50 µl of the supernatant was previously diluted to 320 µl with the extraction
23 solution. Sample absorbance was determined at 450 nm with a spectrophotometer (Hitachi U-1100)
24 and interpolated into a KCl calibration curve (0, 5, 10, 15, 20 ppm) to calculate Cl⁻ concentration.
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42 *Statistical analysis*

43 The data of all parametric measurements were tested within each genotype through to one-way
44 analysis of variance (ANOVA) and to multiple comparisons by Duncan's test. Statistical analyses were
45 performed with the InfoStat package (Di Rienzo et al. 2010).
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52 **Results**

53 *Impact of salinity, alkalinity and saline-alkaline treatments on plant growth and photosynthesis*

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1 Neutral, alkaline and mixed NaCl and NaHCO₃ salts significantly reduced total dry weight of
2 Gifu B-129 and MG20 plants respect to the control treatment (Fig. 1). The magnitude of the induced
3 reductions due to salinity or mixed S-A stress tended to be higher in Gifu B-129 than in MG20
4 (respectively 53% and 83% in the first, versus 34% and 61% in the second ecotype), and similar under
5 alkalinity (Gifu B-129= 61%; MG20= 64%). Pn was reduced in both ecotypes by the mixed S-A stress,
6 and by the alkaline stress in Gifu B-129 plants, whereas Gs and E were strongly reduced by the three
7 types of stress in both ecotypes and Ci did not significantly varied (Table 2).
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16 *Effect of alkaline, saline and saline-alkaline treatments on Chl a fluorescence kinetics and*
17 *behavior of PSII photosynthetic machine, determined by OJIP parameters*
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20 Chlorophyll fluorescence rise transients obtained from control leaves of both genotypes showed
21 a typical OJIP shape (Fig. 2). The “spider plot” diagrams (Fig. 3) showed that the behavior of the
22 different parameters characterizing PSII functioning varied according with the ecotype and the stress
23 type. Overall, most significant variations with respect to the control situation were found in the S-A
24 stress, being those variations more evident in Gifu B-129 than in MG20. Following paragraphs
25 describing variations in fluorescence parameters were statistically supported by Duncan’s Test at $P <$
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34 No significant salt-induced changes in the OJIP curves were observed, excepting in Gifu B-129
35 plants treated with S-A stress, which showed a much lower fluorescence rise. Despite the fact that no
36 salt-induced changes in F_0 and F_m were observed in Gifu B-129 (neither in MG20), some OJIP
37 parameters derived from their relationship were affected by the S-A stress in this ecotype (Figs. 3A, 3B;
38 Table 3): F_v/F_m (ϕP_0) and F_v/F_0 decreased (-5 and -10%, respectively), whereas F_0/F_m increased
39 (20%).
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46 Salt treatment also affected parameters normalized to F_v , although differently, according to the
47 ecotype (Figs. 3A, 3B, Table 3). dV/dt_0 , or M_0 , reflects the accumulation rate of the active reaction
48 centers (RC) fraction that are closed, thus providing information about the acceptor side of PSII
49 (Strasser and Srivastava 1995; Strauss et al. 2003; Strasser et al. 2004; Xia et al. 2004; Mehta et al.
50 2010a, b). M_0 was increased 68% by S-A stress in Gifu B-129 and reduced (21%) in MG20 treated with
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A rise in V_J suggests an increase of the proportions of closed RC and reduced Q_A at J step, whereas a rise in V_I indicates accumulations of reduced Q_A and plastoquinone, which cannot transfer electrons to the dark reactions (Pan et al. 2010). V_J was increased (40%) in Gifu B-129, and showed no change in MG20, whereas V_I showed mild increases in MG20 and Gifu B-129 under S-A stress, and in MG20 treated with NaHCO_3 alone. V_J and M_0 also defined the OJIP parameters $S_s=V_J/M_0$ and $S_m=\text{Area}/(F_M-F_0)$ that represent the normalized total complementary areas corresponding to the O-J phase and OP phase of the kinetic Chl fluorescence, respectively. S_s and S_m reflect the single-turnover Q_A reduction events and the multiple-turnover Q_A reduction events, respectively. S_s and S_m were reduced by S-A stress in Gifu B-129 (-25% and -39%, respectively). Conversely, MG20 increased the S_s values under salinity and S-A stress (22 and 24%, respectively) and showed a slightly fall of S_m (-8%) under S-A stress.

In the S-A treatment, the specific energy fluxes ABS/RC (or also, the Absorption flux (of antenna chlorophylls) per RC, DI_0/RC and TR_0/RC showed contrasting behaviors between genotypes, since they decreased (15% average) in NaCl-treated MG20 plants, but increased (27% average) in Gifu B-129 ones (Fig. 3C, D; Table 3). On other hand, ET_0/RC was 15% average reduced by NaCl in both ecotypes, regardless alkalinity.

Under NaCl treatment, the density of RC or amount of RC per excited cross section (RC/CS_0) and the active Chl associated to RC (γRC) of MG20 leaves were increased in a 20% and a notorious 130%, respectively, while these parameters showed no salt-induced change in Gifu B-129 (Figs. 3C, 3D; Table 3).

In Gifu B-129 plants, S-A stress induced 4% , 30% and 30% decreases of ϕP_0 , and ψE_0 and ϕE_0 , respectively (Figs. 3E, 3F; Table 3), whereas no changes in these parameters were registered in the remaining treatments.

PI_{ABS} is a multiparametric expression elaborated through independent components, all contributing to photosynthesis. PI_{ABS} was increased (29%) by NaCl as a sole stress source in MG20, whereas it was dramatically decreased by the S-A stress in Gifu B-129 (-63%). In line, PI_{ABS} components behaved markedly different according to the ecotype and the type of salt (Figs. 3E, 3F; Table 3).

Effect of alkaline, saline and saline-alkaline treatments on the chlorophyll and ions contents

1 In Gifu B-129 plants, the S-A treatment led to decreased Chl *a* and total Chl contents, and
2 reduced Chl *a/b* ratio, whereas no effect was observed on these parameters with the other two stresses
3 (Fig. 4). In MG20, Chl *b* was reduced by the three types of salt stress, whereas Chl *a* was not affected.
4 As result, total Chl content was reduced by the S-A stress, and no salt-induced change of the Chl *a/b*
5 ratio was observed in this ecotype.
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10 Higher shoot Na⁺ and Cl⁻ levels were found in plants of both ecotypes when confronted with
11 saline or S-A treatments, compared with respective controls (Supplementary Table 1). In contrast, a
12 non-significant change was registered in these levels for plants treated with the alkaline salt alone.
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14 Regardless alkalinity, NaCl reduced K⁺ contents and K⁺/Na⁺ ratios in plants of both ecotypes, whereas
15 the alkaline salt added as a sole stress source affected these parameters only in Gifu B-129 plants.
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22 Discussion

23 24 25 26 *Salt stress impact on growth, gas exchange and ions contents.*

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28 Although the main objective of this work was to gain insight into the salt-induced behavior of
29 the *L. japonicus* photosynthetic apparatus, plant growth, gas exchange and ion accumulation were also
30 analyzed in order to improve our knowledge on *L. japonicus* responses to the three stresses.
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34 Our results from total biomass (Fig. 1), showed that the two ecotypes diverged in their sensitivity
35 level to salinity, in agreement with earlier works (Sánchez et al. 2008; 2010), and that such divergence
36 also applies to the S-A stress. In addition, we showed that both ecotypes were less tolerant to the last
37 stress than to separate alkaline and saline stresses (in that order). A comparable result was previously
38 reported by Paz et al. (2012) on plants of a different *Lotus* species, *L. tenuis*, subjected to the same
39 three stress sources. Previous studies on other plant species have revealed that, when compared at
40 the same concentration level, alkaline salts exert a more detrimental action on plant growth than neutral
41 ones (Shi and Yin 1993; Tang and Turner 1999; Yang et al., 2008; Yang et al., 2009; Gong et al., 2013).
42 It is worthy to note that the alkaline treatment here used had a much lower strength (10 mM of salt) than
43 the saline one (100 mM of salt), what prevents us of confirming former statement. On other hand, the
44 S-A stress solution here employed, had equal electrical conductivity to that of the saline treatment
45 (neutral salt) and equal pH to that of the alkaline one. Therefore, the worse growth performance
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1 registered in the S-A treatment could be a result of a possible additive effect of high saline strength and
2 high pH.
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4 In this work, some expected cause-effect relationships involving biomass and Gs parameters
5 were not observed. Specifically, the negative effect of salt stress on plant biomass was not always
6 accompanied by lower Pn values, suggesting that factors such as a limited nutrient condition (Munns
7 2005; Tsai et al. 2004; Hong et al. 2009) or toxicity (Marschner 1995) may have intervened in such
8 decreases. Indeed, this could be the case in our work: the reduced shoot K⁺ content and K⁺/Na⁺ ratio,
9 or the increased shoot Na⁺ and Cl⁻ accumulation registered in some saline and alkaline treatments
10 (Supplementary Table 1) to levels possibly toxic for diverse metabolic pathways, could have contributed
11 to the reduction of the total biomass in those treatments. In fact, the higher ability of MG20, compared
12 to Gifu B-129 to prevent Na⁺ reaching the shoot was the physiological parameter better explaining the
13 more satisfactory response to salt stress of the first ecotype, in terms of plant biomass. Likewise,
14 significant stress-induced stomatal closures were registered in both genotypes, in the absence of
15 photosynthetic performance variation. The last phenomenon has been previously reported by other
16 authors (Munns and Tester 2008; James et al. 2002) and may be explained through salt-induced
17 changes in leaf anatomy leading to the improvement in CO₂ diffusion (Acosta-Motos et al. 2015a;b;
18 Gómez-Bellot et al. 2015). In this regard, it is worth to mention that alkaline and S-A salts caused a
19 replacement of palisade by spongy parenchyma in leaves of *L. tenuis* (Paz et al. 2014). Further
20 anatomical studies performed on *L. japonicus* ecotypes MG20 and Gifu B-129, testing the hypothesis
21 of a salt-induced reorganization of leaf mesophyll would be valuable for a better understanding of
22 present gas exchange results.
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44 *Salt stress impact on PSII photosynthetic performance*

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46 Data pertaining to the Chl fluorescence analysis showed that the most significant effects
47 induced by salt addition concerned Gifu B-129 plants treated with combined neutral-alkaline salts, and
48 MG20 plants treated with NaCl. The two photosynthesis-related indexes, ϕP_0 and PI_{ABS} were found to
49 be notably altered by diverse environmental constraints in several plant species (Pietrini et al. 2005;
50 Gazquez et al. 2015; 2018). As ϕP_0 may provide clues of photoinhibition occurrence (Yamane et al.,
51 2008; Goh et al., 2012), our data showing no significant salt-induced variation in this parameter (or a -
52 4% decrease in Gifu under S-A) suggest the absence of photoinhibition events due to salt treatment.
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1 The ϕP_0 also reflects the PSII capacity to reduce the primary acceptor Q_A (Calatayud and Barreno
2 2001). The PI_{ABS} is regulated by the three main functional steps of photosynthetic activity by the PSII,
3 which are absorption of light energy (ABS), trapping of excitation energy (TR) and conversion of
4 excitation energy (ET; Strasser et al. 2000). The fact that both parameters were reduced by combined
5 NaCl and $NaHCO_3$ salts in Gifu B-129 plants (Fig. 3), but not in MG20, supports the notion extracted
6 from biomass data of a higher tolerance to the S-A stress in the last ecotype, compared with Gifu B-
7 129. Interestingly, NaCl alone increased PI_{ABS} in MG20 plants, indicating an improvement of the PSII
8 functioning in these plants. However, last result was not reflected by P_n values. This lack of parallelism
9 between net photosynthesis and the mentioned indices may be explained by the electron flux being not
10 used in carbon metabolism, but re-routed to other biochemical pathways such as Mehler reaction, or
11 photorespiration (which may sum 36% of photosynthetic electrons dissipated; e.g.: in tomato under
12 water stress; Haupt-Herting and Fock et al. 2002).

23 *Salt stress-induced modifications of the photosynthetic apparatus structures*

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25 RC/ABS and RC/CS are measures of the density of RC per ABS and cross section or CS,
26 respectively. The most commonly observed RC response to salinity in glycophytes is their reduction,
27 (e.g. *Scenedesmus obliquus*, Demetriou et al. 2007; *Cucumis melo*, Xiang et al. 2016; *Triticum sp*,
28 Mehta et al. 2010); *Wolffia arrhiza*, Wang et al. 2011 and *Solanum lycopersicum*, Zushi and Matsuzoe
29 2017). In this study, the RC pool size was another parameter differently altered by salt stress according
30 to the ecotype. RC reduction was registered in Gifu B-129 plants exposed to combined NaCl and
31 $NaHCO_3$ salts, confirming the glycophytic *L. japonicus* status (Sanchez et al. 2008; Sanchez et al.
32 2011). In contrast, both RC densities (per both cross section and absorption energy flux) increased in
33 NaCl-treated MG20 plants, in line with higher values of γRC , registered in this ecotype. Having in mind
34 that salt-treated plants of the halophyte *Artemisia anethifolia* presented higher RC/CS than
35 corresponding controls (Wen et al. 2005), former result could be interpreted as an adaptation, through
36 improvement of RC stability, of the photosystem machinery to NaCl in this ecotype. Interestingly, the
37 MG20 ecotype was considered as a moderate halophyte by Melchiorre and collaborators (2009) when
38 confronted with 50 mM NaCl. Thus, the opposite S-A-induced responses observed in the RC/ABS
39 between genotypes, an increase in MG20 and a decrease in Gifu B-129, could have also contributed
40 with the contrasting behaviour of PI_{ABS} in these ecotypes.

1 Variations in the effective antenna size of PSII may be inferred from changes in ABS/RC,
2 chlorophyll content and Chl *a/b* ratio. These parameters varied concomitantly in Gifu B-129 plants
3 under the mixed S-A treatment, supporting the notion that mixed NaCl and NaHCO₃ salts induced a
4 higher effective antenna size in this last ecotype, with preferential loss of reactions centre complexes
5 (DI₀/RC, Table 3; Figs. 3 and 4). In parallel, the chlorophyll contents and chl *a/b* data support specific
6 loss of LHC in MG-20.
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14 *Salt effects on electron transport chain and carriers*

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16 Changes in several obtained and calculated parameters support the notion that, in Gifu B-129
17 plants confronting mixed (neutral and alkaline) salts, photoinhibition took place on both sides of PSII
18 (acceptor and donor). On one hand, the reduction in Fv/F₀ reflects a decrease in the water-splitting
19 complex activity, at the donor PSII site (Schreiber et al. 1994; Sayed 1998; Kalaji et al. 2011). Also, the
20 decrease of S_s values in Gifu B-129 under S-A stress indicated a detriment of the processes associated
21 to the primary photochemistry. Interestingly, higher S_s values in MG20 under saline and S-A stresses
22 would indicate an improvement of the primary photochemistry. On the other hand, the decrease in ψE_0
23 and the raise in V_i, V_J and M₀ in Gifu B-129 plants treated with combined NaCl and NaHCO₃ salts
24 conform the idea of a slowdown of electron transfer from Q_A to the secondary acceptor Q_B, on the
25 acceptor PSII side. Also, the decrease in S_m registered in Gifu B-129 plants under S-A stress suggests
26 a decline in the pool of electron carriers between PSII and PSI, as S_m reflects the total electron carriers
27 per RC (Jiang et al., 2008). Last results point to a reduction of the total electron acceptor capacity of
28 S-A treated Gifu B-129 leaves. Besides (unlike MG20), Gifu B-129 plants under S-A treatment
29 presented increased TR₀/RC despite there was no significant change in the RC level. According to
30 Weng et al. (2005), last situation indicates a lower Q_A re-oxidation. In fact, our data showing a reduction
31 in the electron transport per reaction center (ET₀/RC) in these plants, is in line with the last phenomenon,
32 and supports the decreased ψE_0 , and the increased DI₀/RC (non-photochemical quenching of energy)
33 registered. Indeed, this scenery of electrons transport blockage, in addition to a higher effective
34 antenna size as it was pictured above for Gifu B-129, explains the higher energy dissipation of the
35 excess excitation energy trapped in the non-RC (DI₀/RC, Fig. 3D, Table 3). A rise in non-photochemical
36 quenching may be the outcome of changes in either photoinhibition or protective high-energy-state
37 quenching. As we stated above, our results from ϕP_0 suggested the absence of photoinhibition events
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1 due to salt treatment. In tomato, increasing non-photochemical quenching, associated to lower ϕP_0
2 was suggested to intervene in excess energy dissipation to keep photosynthetic apparatus from being
3 dismantled (Gong et al., 2013). Energy dissipation prevents the reduction of Q_A to Q_A^- and therefore, it
4 does not contribute to the variable fluorescence (see Strasser and Strasser; 1995; Strasser et al. 2000;
5 Krüger et al.; 1997; Appenroth et al.; 2001; Jafarina and Shariati 2012). That is why the OJIP curves
6 obtained from Gifu B-129 plants treated with combined NaCl and NaHCO₃ salts presented lower
7 fluorescence levels than those from control plants (Fig. 2). Taken as a whole, these results explain the
8 observed PI_{ABS} and ϕP_0 decreases in the Gifu B-129 ecotype.
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16 Conclusions

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19 Figure 5 summarizes the effects of different stresses on the photosynthetic apparatus of both
20 genotypes, explained on the base of their energy fluxes, quantum yields and efficiencies, and pools of
21 each photosynthetic machine component. Our results allow us to accept the proposed hypothesis.
22 Here we conclude that NaCl-derived stress sources affect the photochemical events in PSII in both
23 ecotypes, being this effect more evident under higher pH condition, whereas alkalinity *per se* has a mild
24 or no effect on these events. Our study also allowed us to determine quantitative and qualitative
25 differences between both ecotypes regarding Chl *a* fluorescence response to salt stresses. The S-A
26 treatment induced a more severe effect on Gifu B-129, compared with MG20, particularly on those
27 parameters related with the functioning of a section spanning the donor (water splitting complex) and
28 the first acceptor (Q_A) sides of the electron transport chain. In contrast, NaCl improved primary
29 photochemistry in MG20, although the step between Q_B and PSI was functionally compromised in this
30 ecotype. The fact that some components, or sequential steps of the PSII photosynthetic activity in Gifu
31 B-129 and MG20 had diverged during their salts-induced responses, indicates that both model ecotypes
32 would not be equivalents for performing experiments addressing more specific hypothesis on
33 physicochemical or structural aspects of the *L. japonicus* photosynthetic machinery. Therefore, we
34 consider the information here obtained is useful and should be taken into account in future research on
35 structural and biochemical aspects of the photosynthetic apparatus, aimed at elucidating the
36 mechanisms that make possible the tolerance to these stressful conditions.
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8 Author's contributions

9 ABM: Conceptualization; RR, CDB, MPC: Methodology; ABM, AAR: Formal analysis; SJM,
10 OAR: Funding acquisition; AMB: Writing - original draft; AMB, AAR, CDB: Writing - review &
11 editing.
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Legends

Figure 1. Total dry weight of *L. japonicus* plants. Same letter within each ecotype means no statistically different (Duncan's test, $P < 0.05$). Empty bars = MG20. Grey bars = Gifu B-129.

Figure 2. Chlorophyll a fluorescence OJIP transient curves obtained from leaves MG20 and Gifu B-129.

Figure 3. Spider plots of the parameters measured and deduced from Chl a fluorescence OJIP transient curves in MG20 (A, C, E) and Gifu B-129 (B, D, F).

Figure 4. Contents of Chl *a*, *b*, Chl *a/b* ratio and total Chl content in apical leaves of Gifu B-129 and MG20 plants. Asterisks mean statistically different from control at $P < 0.05$, according with Duncan's Test.

Figure 5. Schematic representation of OJIP test results. Central block represents a control condition for both ecotypes. Right and left blocks depict the sets of salt-induced events revealed by Chl fluorescence data in MG20 and Gifu B-129. Black boxes with A, S or S-A mean alkaline, saline or S-A treatments, respectively. RC, active reaction center; RC with dashed line means "non-RC"; Chl_{RC}, active Chl in RC; Q_A and Q_A⁻, quinone A oxidized and reduced, respectively. Smaller and bigger arrows and

letters with respect to the central block mean negative and positive changes regarding quantum efficiencies and the activity of the water-splitting complex, compared with the untreated control.

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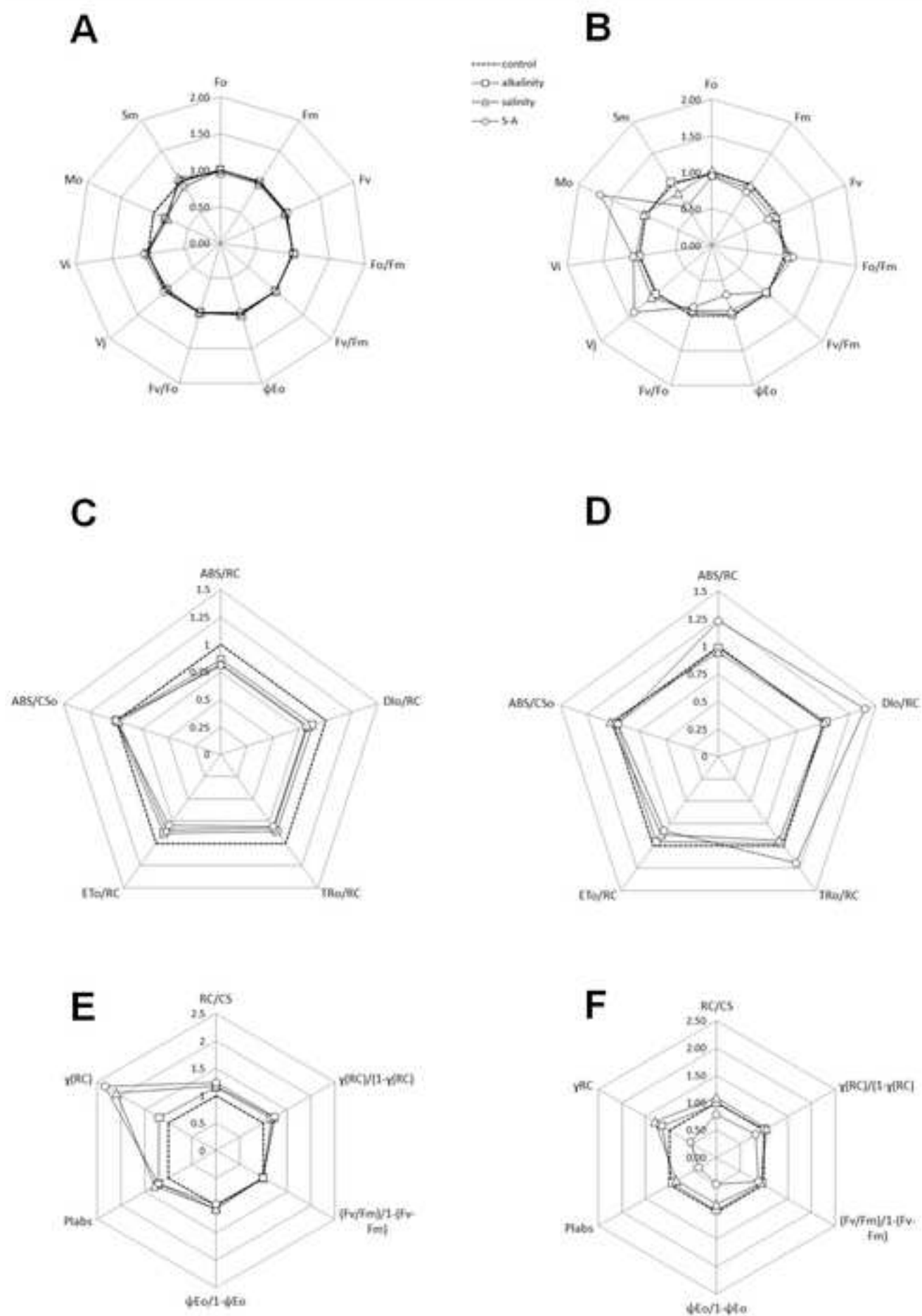
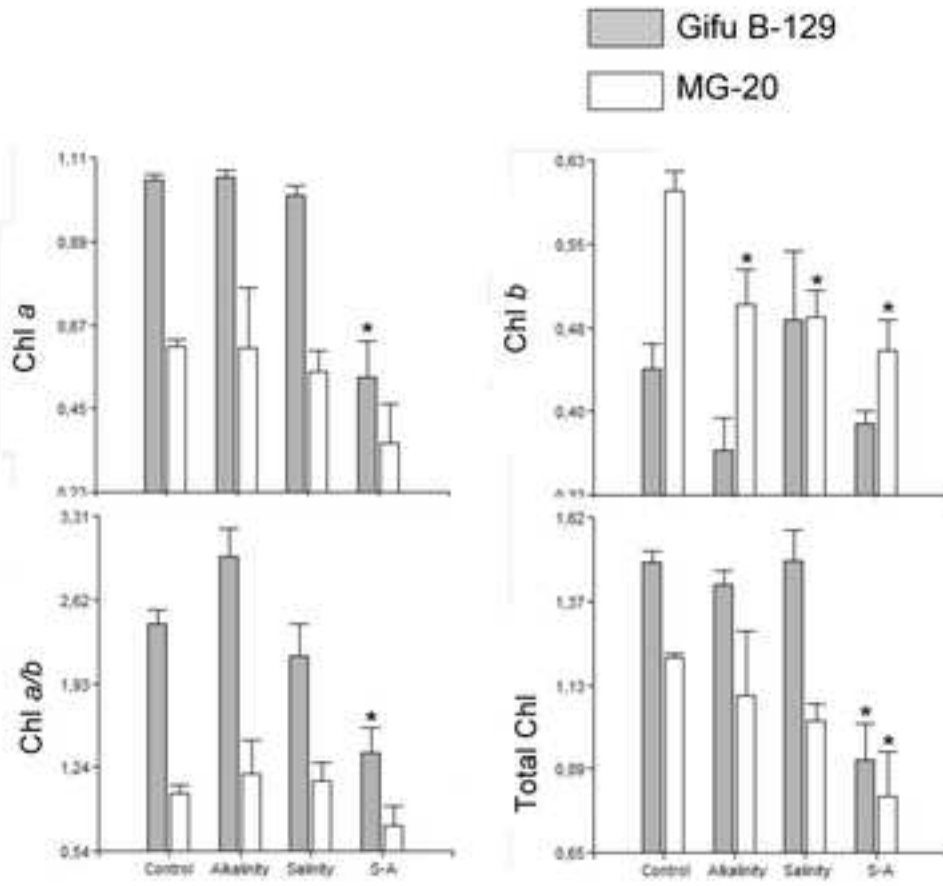


Figure 4



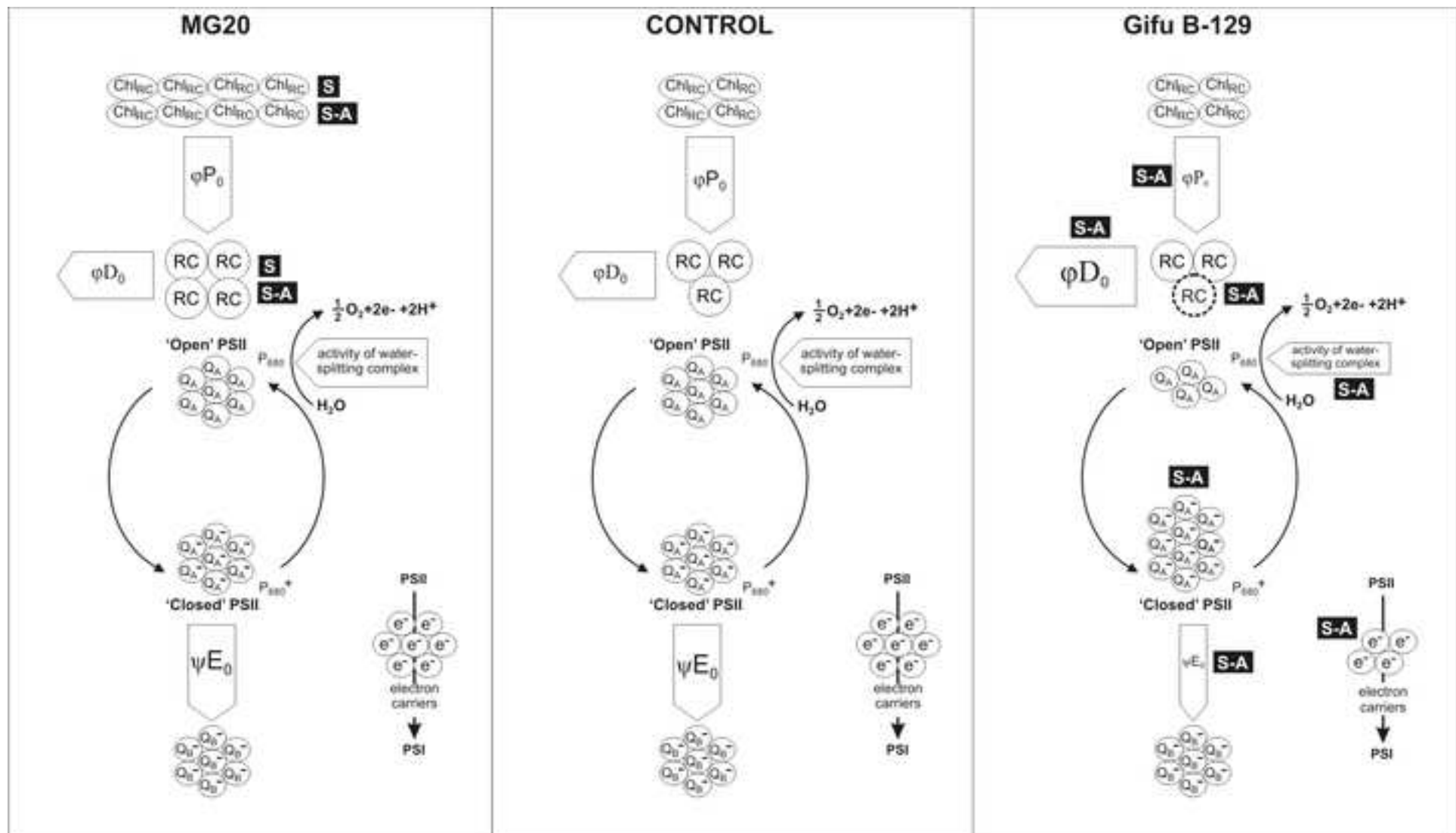


Figure 1

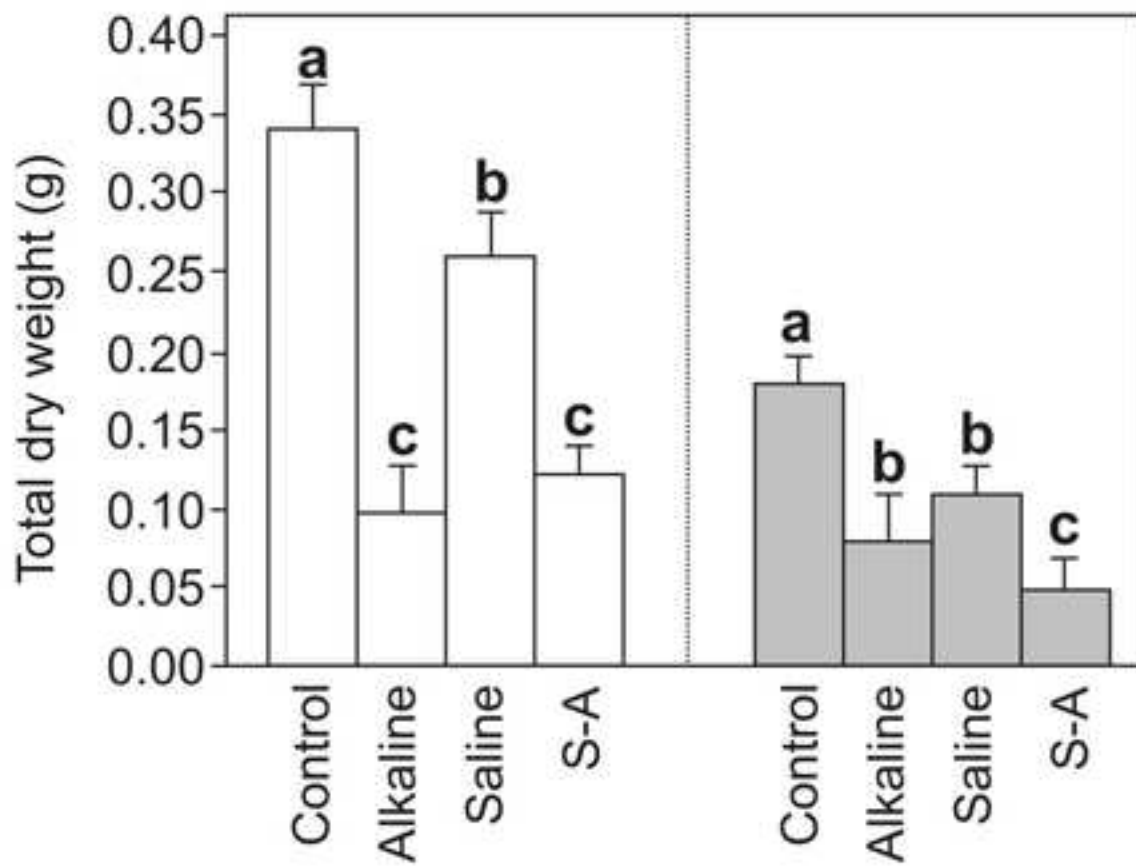


Figure 2

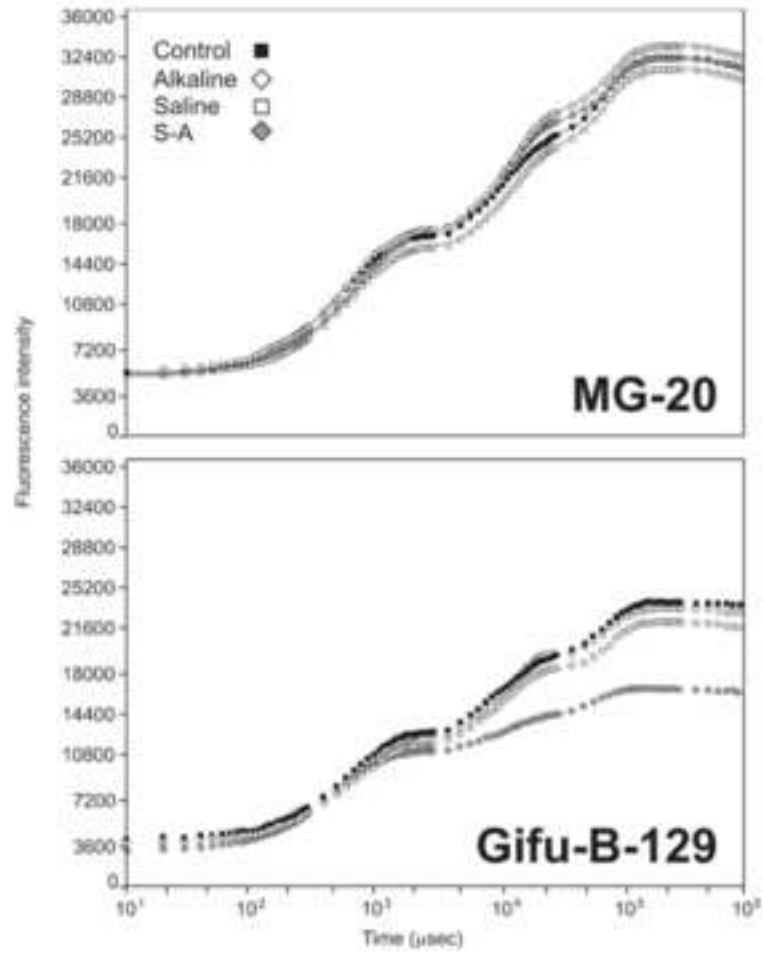


Table 1. Summary of parameters, formulae and their description using data extracted from chlorophyll *a* fluorescence (OJIP) transient. Strasser et al. 1999, Tsimilli-Michael and Strasser 2008).

<i>Extracted parameters</i>	<i>Description</i>
F_0	Minimum fluorescence at $t = 0$, when all RC of PSII are open
F_M	Maximum fluorescence, when all RC of PSII are closed
Area	total complementary area between the fluorescence induction curve and $F = F_M$
<i>Derived parameters</i>	
$F_V = F_M - F_0$	Maximal variable fluorescence
F_V/F_0	Proportional ratio to the activity of the water-splitting complex
$V_J = (F_{2\text{ ms}} - F_0)/F_V$	Relative variable fluorescence at the J-step (2 ms)
$V_I = (F_{30\text{ ms}} - F_0)/F_V$	Relative variable fluorescence at the I-step (30 ms)
$M_0 = 4 (F_{300\ \mu\text{s}} - F_0)/F_V$	Approximated initial slope (in ms^{-1}) of the fluorescence transient
$S_s = V_J/M_0$	Normalized total complementary area corresponding only to the O-J phase
$S_m = \text{Area}/F_V$	Normalized total complementary area corresponding to the O-P phase or total electron carriers per RC
<i>Quantum efficiencies</i>	
F_V/F_M (or ϕP_0)	Maximum quantum efficiency of the PSII (primary photochemistry) at $t = 0$
F_0/F_M (or ϕD_0)	Maximum quantum efficiency at $t = 0$ for energy dissipation
$\psi E_0 = 1 - V_J$	Quantum efficiency that an electron moves further than Q_A^-
<i>Specific and phenomenological energy fluxes</i>	
$\text{ABS}/\text{RC} = M_0 \cdot (1/V_J) \cdot (1 - \phi P_0)$	Absorption flux (of antenna Chl) per RC
$\text{TR}_0/\text{RC} = M_0/V_J$	Trapped energy flux per RC at $t = 0$
$\text{ET}_0/\text{RC} = M_0 \cdot (1/V_J) \cdot (1 - V_J)$	Electron transport flux per RC at $t = 0$
$\text{DI}_0/\text{RC} = (\text{ABS}/\text{RC}) - (\text{TR}_0/\text{RC})$	The flux of energy dissipated in processes other than trapping per active PSII
$\text{ABS}/\text{CS}_0 = \text{Chl}/\text{CS}$	Absorption flux per cross section (CS) at $t = 0$
<i>RC densities and active Chl</i>	
$\text{RC}/\text{CS}_0 = \phi P_0 \cdot \text{ABS}/\text{CS}_0 \cdot S_s$	Amount of RC per CS at $t = 0$
$\text{RC}/\text{ABS} = \gamma \text{RC}/(1 - \gamma \text{RC})$	Amount of RC per Chl
$\gamma \text{RC} = (1/(\text{ABS}/\text{RC}))/((1 - (1/(\text{ABS}/\text{RC})))$	Probability that a PSII Chl molecule functions as RC
<i>Performance index and its components</i>	
$\gamma \text{RC}/(1 - \gamma \text{RC})$	Structural component of PI_{ABS} (or EC)
$\phi P_0/(1 - \phi P_0)$	Photochemical component of PI_{ABS} (or PC)
$\psi E_0/(1 - \psi E_0)$	Biochemical component of PI_{ABS} (BC)
$\text{PI}_{\text{ABS}} = \text{EC} \cdot \text{PC} \cdot \text{BC}$	Performance index based on the equal absorption

Table 2. Gas exchange parameters net photosynthesis rate (Pn; $\mu\text{mol CO}_2 \text{ m}^{-2} \text{ s}^{-1}$), stomatal conductance (Gs; $\text{mmol H}_2\text{O m}^{-2} \text{ s}^{-1}$), transpiration rate (E; $\text{mmol H}_2\text{O m}^{-2} \text{ s}^{-1}$) and internal CO_2 (Ci; $\mu\text{mol mol}^{-1}$) in leaves of *L. japonicus* Gifu B-129 and MG-20 plants. Same letter within each ecotype means no statistically different (Duncan's test, $P < 0.05$).

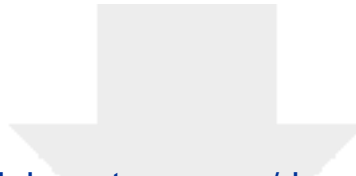
Ecotype	Treatment	Pn	Gs	E	Ci
Gifu B-129	Control	$1.8 \pm 0.3a$	$39 \pm 2.9a$	$0.58 \pm 0.03a$	$412 \pm 94ab$
	Alkaline	$0.6 \pm 0.3b$	$25 \pm 1.2b$	$0.33 \pm 0.02b$	$359 \pm 64ab$
	Saline	$1.5 \pm 0.3a$	$17 \pm 2.1c$	$0.26 \pm 0.03c$	$331 \pm 70b$
	Mixed S-A	$0.1 \pm 0.2b$	$20 \pm 1.2c$	$0.21 \pm 0.02c$	$581 \pm 85a$
MG-20	Control	$5.4 \pm 0.4a$	$103 \pm 6.4a$	$1.21 \pm 0.05a$	$403 \pm 35ab$
	Alkaline	$5.4 \pm 0.6a$	$43 \pm 4.3b$	$0.50 \pm 0.04b$	$363 \pm 32ab$
	Saline	$3.4 \pm 0.7a$	$37 \pm 2.7bc$	$0.43 \pm 0.04bc$	$341 \pm 33b$
	Mixed S-A	$0.1 \pm 0.05b$	$30 \pm 2.9c$	$0.35 \pm 0.05c$	$455 \pm 35a$

Table 3. Significant variations (%) observed in the different OJIP parameters in MG20 and Gifu B-129 plants under the three stressful conditions in comparison with the control (Duncan's test, $P < 0.05$). Dash line means no significant differences, compared with the control treatment.

Parameter	Description	MG20			Gifu B-129		
		Alkaline	Saline	S-A	Alkaline	Saline	S-A
<i>Extracted and derived parameters</i>							
F_0	Minimum fluorescence at $t = 0$, when all RC of PSII are open	-	-	-	-	-	-
F_M	Maximum fluorescence, when all RC of PSII are closed	-	-	-	-	-	-
F_V	Maximal variable fluorescence ($F_M - F_0$)	-	-	-	-	-	-
F_V/F_0	Proportional ratio to the activity of the water-splitting complex	-	-	-	-	-	-
V_i	Accumulations of Q_A^-	-	-	-	-	-	-10
V_i	Accumulations of Q_A^- and reduced PQ	5	-	5	-	-	40
M_0	Accumulation rate of closed RC	-	-21	-21	-	-	9
Ss	Reflects the single-turnover Q_A^- reduction events	-	22	24	-	-	68
Sm	Reflects the multiple-turnover Q_A^- reduction events or total electron carriers per RC	-	-	-	-	-	-21
<i>Quantum efficiencies</i>							
ϕP_0	Efficiency of PSII to reduce Q_A^-	-	-	-	-	-	-39
ϕD_0	Efficiency for energy dissipation	-	-	-	-	-	20
ψE_0	Efficiency to move an electron further than Q_A^-	-	-	-	-	-	-30
<i>Specific energy and phenomenological fluxes</i>							
ABS/RC	Absorption flux (of antenna Chl) per RC	-	-15	-19	-	-	23
TR ₀ /RC	Trapped energy flux per RC at $t = 0$	-	-16	-21	-	-	19
ET ₀ /RC	Electron transport flux per RC at $t = 0$	-	-10	-19	-	-13	-17
DI ₀ /RC	The flux of energy dissipated in processes other than trapping per active PSII	-	-13	-13	-	-	36
ABS/CS ₀	Absorption flux per cross section (CS) at $t = 0$	-	-	-	-	-	-
<i>RC densities, active Chl in RC, performance indices and components</i>							
RC/CS ₀	Amount of RC per CS at $t = 0$	-	18	24	-	-	-
γ RC	Probability that a PSII Chl molecule functions as RC	-	121	135	-	-	-
γ RC/(1 - γ RC)	Structural component of PI_{ABS}	-	17	23	-	-	-17
$\phi P_0/(1 - \phi P_0)$	Photochemical component of PI_{ABS}	-	-	-	-	-	-12
$\psi E_0/(1 - \psi E_0)$	Biochemical component of PI_{ABS}	-	-	-	-	-	-50
PI_{ABS}	Performance index based on the equal absorption	-	29	-	-	-	-63

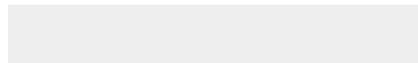
Author's contributions

ABMenéndez: Conceptualization; RRocco, CDBordenave, MPCampestre: Methodology;
ABMenéndez, AARodríguez: Formal analysis; SJMaiale, OARuiz: Funding acquisition;
ABMenéndez: Writing - original draft; ABMenéndez, AARodríguez, CDBordenave: Writing -
review & editing.



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