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Effect of salt stress on different tiller positions in rice and the regulatory effect of Prohexadione calcium

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Soil salinization has resulted in a significant decrease in crop yields, particularly affecting the production of crops like rice (Oryza sativa L.). Prohexadione calcium (Pro-Ca) can enhance crop resilience against failure by managing plant height. However, its impact on various tiller positions during the tillering phase of rice under salt stress remains unknown. This study explores the distinct effects of salt stress on the physiological traits of the main stem and different tiller segments of rice plants, along with the role of Pro-Ca in mitigating salt stress. The findings revealed that under salt stress conditions, the number of tillers and leaves on the main stem decreased significantly in rice. Moreover, the levels of malondialdehyde (MDA) and H₂O₂ in the leaves and stems of each tiller position notably increased. The percentage of tillers experiencing reduction or elevation was higher than that of the main stem compared to the respective control. Application of Pro-Ca through foliar spraying under NaCl stress effectively alleviated the impact of salt stress on the tiller growth of rice during the tillering phase. It also boosted the activities of antioxidant enzymes like superoxide dismutase (SOD) and peroxidase (POD) in the leaves and stems of the tillers. Furthermore, it successfully mitigated the damage inflicted by salt stress on the cell membrane of rice tillers during the tillering phase. The regulatory effect of calcium on cyclic acid was particularly pronounced in alleviating the impact on the tillers under salt stress conditions.

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30	Abstract
31	Soil salinization has resulted in a significant decrease in crop yields, particularly affecting the
32	production of crops like rice (Oryza sativa L.). Prohexadione calcium (Pro-Ca) can enhance crop
33	resilience against failure by managing plant height. However, its impact on various tiller
34	positions during the tillering phase of rice under salt stress remains unknown. This study
35	explores the distinct effects of salt stress on the physiological traits of the main stem and
36	different tiller segments of rice plants, along with the role of Pro-Ca in mitigating salt stress. The
37	findings revealed that under salt stress conditions, the number of tillers and leaves on the main
38	stem decreased significantly in rice. Moreover, the levels of malondial dehyde (MDA) and $\rm H_2O_2$
39	in the leaves and stems of each tiller position notably increased. The percentage of tillers
40	experiencing reduction or elevation was higher than that of the main stem compared to the
41	respective control. Application of Pro-Ca through foliar spraying under NaCl stress effectively
42	alleviated the impact of salt stress on the tiller growth of rice during the tillering phase. It also
43	boosted the activities of antioxidant enzymes like superoxide dismutase (SOD) and peroxidase
44	(POD) in the leaves and stems of the tillers. Furthermore, it successfully mitigated the damage
45	inflicted by salt stress on the cell membrane of rice tillers during the tillering phase. The
46	regulatory effect of calcium on cyclic acid was particularly pronounced in alleviating the impact
47	on the tillers under salt stress conditions.
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Introduction

56	In recent years, soil salinization has become a global issue due to rising sea levels and expanding
57	salinized land areas caused by global warming (Ahmed et al., 2021), severely reducing global
58	crop yields and agricultural production (Munns & Tester, 2008; Dai et al., 2022). Enhancing crop
59	salt tolerance and effectively addressing the damage from salt stress on yield has emerged as a
60	critical research focus (And et al., 2011). Salt stress is a key abiotic factor that impacts the
61	growth of most plants. In general, salt concentrations lead to changes in physiological and
62	biochemical functions, limiting the growth and development of above-ground plant parts and
63	root systems (dos Santos et al., 2022), When soil salinity increases, the water potential of the soil
64	solution decreases below that of the plant root cells. This makes root uptake challenging and
65	results in osmotic stress. Osmotic stress leads to the closure of plant stomata, hindering CO ₂
66	uptake, weakening photosynthesis, causing nutrient deficiencies. The accumulation of Na ⁺ and
67	Cl in cells affects mineral uptake and transport, inhibits enzyme activity, and leads to
68	dehydration of plant cells. Salt stress can also result in the accumulation of reactive oxygen
69	species (ROS), damaging cellular structures and biomolecules, thus limiting the growth of major
70	crops like rice. (Hussain et al., 2018). Prolonged exposure to harsh environments has led to the
71	evolution of a range of salt tolerance mechanisms in crops, including: changes in morphology,
72	water relations, photosynthesis, hormones, ion distribution and biochemical adaptations (dos
73	Santos et al., 2022), within certain limits, allow them to competitively obtain water from the soil
74	and maintain nutrient balances in the body in response to ionic stresses thereby surviving adverse
75	soil conditions (Hoang et al., 2016).
76	Rice (Oryza sativa L.), a moderately salt-sensitive crop (Joseph et al., 2010), is severely
77	affected by salt stress in its growth and development (Zhang et al., 2012), and this effect varies
78	according to developmental stage, degree and duration of stress, and variety (Zheng et al., 2023).
79	It has shown that the effects of salt stress on rice germination and emergence are mainly
80	characterized by a reduction in germination rate, germination speed, and germination energy,



leading to a reduction in shoot length, root length, and dry weight of rice (Taratima et al., 2022). 81 Salt stress during the seedling period is mainly manifested in the damage to leaves and root 82 83 system (Chang et al., 2019). Salt stress affects rice tillering mainly by reducing tillering capacity and delaying the reproductive process, and the duration of delay is positively correlated with the 84 degree of salt stress, and primary and secondary tillers being more affected than the main stem. 85 Additionally, salt stress reduces soil fertility and causes nutrient imbalance, and salinity stress 86 87 inhibits nutrient uptake by the root system, ultimately leading to reduced tillering or tiller death due to nutrient deficits (Ruan et al., 2008). Primary stems and primary tillers contribute more to 88 crop yield than secondary tillers due to asymmetric competitive advantages under stress 89 conditions, and these advantages are associated with increased leaf number. Transportation of 90 91 water and nutrients between the primary stem and tiller through the vascular bundles at the tiller 92 nodes is essential for tiller development and survival (Yang et al., 2022). Salt stress in the formation of young spikes and spiking and flowering stage of rice is mainly manifested in the 93 following ways: yellowing of leaves, delayed spiking, prolonged spiking period, increase in the 94 number of degradation of glumes, shorter spike lengths, decrease in the number of solid grains, 95 less full grains, more black rotting of roots in the late stage, early senescence, and ultimately 96 affecting the yield of rice (Chang et al., 2019). 97 Plant growth regulators, as organic compounds with effects on growth and development 98 similar to natural plant hormones, control plant growth by initiating various physiological and 99 metabolic processes (Kaya et al., 2023; Zhao et al., 2023). The formation of the endogenous 100 plant hormone gibberellin requires hydroxylase enzymes to catalyze a series of hydroxylation 101 reactions, and these hydroxylases require 2-ketoglutarate as a coenzyme. Prohexadione calcium 102 (Pro-Ca) imitates the coenzymes' structure and competitively hinders their function, thus 103 impeding the synthesis of active gibberellins. Among these hydroxylation reactions, the reaction 104 pathway for the formation of GA1 is the most sensitive to Pro-Ca, whereas the pathway for the 105 formation of GA4 is not involved in the β-hydroxylation reaction, so that Pro-Ca selectively 106 inhibits the synthesis of gibberellin GA1. GA1 is mainly found in the nutrient organs, controlling 107



108	the elongation and growth of stems and leaves, while GA4 is mainly found in the reproductive
109	organs, controlling flower bud differentiation and hot grain development. Pro-Ca is an ideal
110	dwarfing agent because of its strong synthetic activity in inhibiting GA1. Pro-Ca inhibits active
111	gibberellin synthesis while protecting the activity of both surviving gibberellins, so Pro-Ca has
112	dual activity on gibberellin metabolism (Kim et al., 2010; Ilias & Rajapakse, 2005). Pro-Ca has
113	been shown by previous authors to have specific regulatory effects on rice, apple, strawberry, etc.
114	(Kim et al., 2010; Kim et al., 2007; Lee et al., 1998). In previous studies, researchers have
115	explored the risks of salt stress, the characteristics of tillering, and the mechanism of action of
116	Pro-Ca acid. Our own research has shown that Pro-Ca acid can mitigate the harm caused by
117	NaCl to the antioxidant capacity, photosynthetic properties, and cell membranes during the
118	tillering stage in rice (Zhang et al., 2023; Huang et al., 2023). However, further investigations are
119	necessary to understand the varying impacts of salt stress on the main stem and tiller, as well as
120	the regulatory function of Pro-Ca.
121	In this study, we aimed to investigate the differential effects of salt stress on rice main stems
122	and tillers and the regulatory role of calcium switched acid by comparing the relevant
123	morphology building indexes, antioxidant enzyme activities, membrane damage indexes, and
124	soluble protein contents in leaves and stems of rice main stems, first tillers and second tillers at

127

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Materials and Methods

the tillering stage under different treatments.

128 Materials and reagents

- Huanghuazhan (conventional rice) was provided by Longping Seed Co. Ltd (Hunan, China), and
- 130 Xiangliangyou900 (hybrid rice) was provided by Nianfeng Seed Science and Technology Co.
- 131 Ltd (Hunan, China).
- The original solution of the test regulator 5% Pro-Ca used in this experiment was provided
- by Sichuan Runer Technology Co. Ltd (Chengdu, Sichuan).

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Experimental designs

Full and uniform rice seeds were selected, sterilized with 3% H₂O₂ for 15 min and then washed 136 repeatedly with distilled water, distilled water was added until the seeds were submerged, and the 137 seeds were soaked for 24 h at 30°C, after which they were germinated under dark conditions for 138 24 h. The experiment was selected to be carried out in the daylight linkage greenhouse of the 139 College of Coastal Agriculture, Guangdong Ocean University, and the germinated seeds were 140 uniformly sown on the rice-planting trays (specifications of 28~30 cm × 58~60 cm), about 5-8 141 seeds in each hole, and the soil used for seedling was a 3:1 mixture of brick red soil and nutrient 142 soil. 143 After transplanting cultivation using caliber \times bottom diameter \times height of $19 \times 15 \times 18$ cm 144 plastic pots, each pot containing 3 kg of sun-dried soil, before transplanting a fixed amount of 145 146 each pot to add 1 L of water, to be stabilized when the water surface line marking, and regularly replenish water to maintain the water layer. When the seedlings in the seedling tray were three 147 leaves and one heart, the seedlings with consistent growth were selected and transplanted, and 148 the depth of transplanting was about 1.5 cm, with 3 holes in each bucket and 1 plant in each hole. 149 After the end of greening and before tillering, select the evening of sunny weather at about 16:00 150 to carry out regulator treatment through foliar spraying, about 10ml per pot, to ensure that the 151 front and back of the leaf spraying evenly, in order to ensure its normal absorption. The regulator 152 treatment was followed by a 0.3% salt treatment 48 h later. Tagging and tracking marking of 153 154 tiller occurrence. Tillers were labeled with secondary and leaf positions. Ensure spatial distance between seedlings at each sampling to prevent competition between individuals due to different 155 spatial size. 156 The experiment was set up with eight treatments, Xiangliangyou900 variety included four 157 treatments as follows: XCK (distilled water), XS (0.3% NaCl), XPro-Ca (100 mg·L⁻¹ Pro-Ca), 158 XPro-Ca+S (100 mg·L⁻¹ Pro-Ca + 0.3% NaCl), and Huanghuazhan variety included four 159 treatments as follows: ZCK (distilled water), ZS (0.3% NaCl), ZPro-Ca (100 mg·L⁻¹ Pro-Ca), and 160 ZPro-Ca+S (100 mg·L⁻¹ Pro-Ca + 0.3% NaCl), with three replications per treatment. The leaves 161



and stems of the main stem, the first tiller, and the second tiller were taken every 7 d (7 d, 14 d,
21 d, 28 d, and 35 d after salt treatment) for the determination of related indexes.
Determination of morphological indices
Morphological indexes such as plant height, root length, and number of tillers were measured
directly by using vernier calipers to measure the intersection of stem and root to determine the
stem base width, and by using a leaf area meter (YX-1241) to measure the inverted two leaves
and inverted three leaves of each tiller position, and by using the conventional drying method,
the samples of each part of each treatment were killed for several and a half times for 30 min in
an oven at 105°C, and then dried at 80°C to a constant weight and then determined the dry
weight.
Determination of antioxidative enzyme activities
At 7 d, 14 d, 21 d, 28 d, and 35 d after NaCl stress, the leaves and stems of 15 rice plants with
different tiller positions were rapidly frozen in liquid nitrogen, and then stored in -80°C,
respectively. Three replicates of 0.5 g samples (leaves and stems weighed separately) were
ground in liquid nitrogen, and then 10 ml of pre-cooled phosphate buffer (0.05 mM PBS, pH 7.8)
was added, ground to homogenate, and then centrifuged at 6000 × g for 20 min at 4°C, and the
supernatant was aspirated and set aside in a 4°C refrigerator.
$\frac{3 \text{ ml}}{2}$ of reaction solution (50 ml PBS pH 6.0 + 28 μ L guaiacol + 19 μ L H ₂ O ₂ at 30%
concentration) was mixed with 40 μL of supernatant. The absorbance was recorded every 30 s for
4 times and the dynamic absorbance was measured at 470 nm using a spectrophotometer
(GENESYS 180 UV-Vis, Thermo Sci) to determine the peroxidase (POD) activity, 1 unit of
enzyme activity for each minute of OD increase of 0.01.
0.1 ml of supernatant was taken and mixed with 2.9 ml of reaction solution (100 ml of PBS
pH $7.0 + 0.05$ ml of H_2O_2 at 30% concentration) and the absorbance at 240 nm was measured
using a spectrophotometer ((GENESYS 180 UV-Vis, Thermo Sci) and recorded every 30 s for



four times. CAT activity was calculated using 0.01 decrease in OD per minute as a unit of 189 enzyme activity (Aebi, 1984). 190 Superoxide dismutase (SOD) activity was determined using the nitro blue tetrazolium (NBT) 191 method (Giannopolitis & Ries, 1977). 0.1 ml of supernatant was added to 2.9 ml of reaction 192 mixture (2.61 ml MET + 0.097 ml EDTA-Na₂ + 0.097 ml NBT + 0.097 ml riboflavin) and 193 irradiated for 20 min at 4000 lux light at 25°C. At the end of the reaction the absorbance at 560 194 195 nm was measured using the solvent in the unilluminated cuvette as a control tube, and the total activity of SOD was also calculated using 50% inhibition of NBT photochemical reduction as 196 one enzyme activity unit (U)... 197 $\frac{0.1 \text{ ml}}{0.1 \text{ ml}}$ of supernatant was mixed with the reaction solution (2.6 ml EDTA-Na₂ + 0.15 ml 198 AsA + 0.15 ml H_2O_2) and the absorbance at 290 nm was determined by spectrophotometer 199 recording every 30 s for 4 times. A change in absorbance of 0.01 in 1 min is defined as one unit 200 of enzyme activity u, and enzyme activity is expressed as u/g (FW) (Nakano & Asada, 1981). 201 202 **Determination of membrane damage index** 203 Malondialdehyde (MDA) content was determined by TBA method (Guo et al., 2018). 10 ml of 204 phosphate buffer (0.05 mM PBS, pH 7.8) was added to 0.5 g of sample and ground, then 205 centrifuged at 10,000 × g for 10 min at 4°C. 1 ml of the supernatant was taken and mixed with 2 206 ml of 0.6% TBA (thiobarbituric acid) in a centrifuge tube. The mixture was boiled in a boiling 207 water bath for 15 min and then centrifuged at 10000 × g and 25°C for 10 min. The absorbance of 208 the supernatant was measured spectrophotometrically at 450 nm, 532 nm, and 600 nm, 209 respectively. The MDA content was calculated according to Equation: MDA content (mmol·g⁻¹ 210 FW)= $[6.452 \times (OD_{532} - OD_{600}) - 0.559 \times OD_{450}] \times Vt / (Vs \times W)$, where Vt is the total volume of 211 extract, Vs is the volume of extract used for the assay, and W represents the fresh weight of the 212 sample. 213 0.5 g of the sample was taken, 5 ml of 0.1% TCA solution was added, ground in liquid 214 nitrogen and centrifuged at 10,000 × g for 10 min. Then, 0.5 ml of the supernatant was added to 215



216	0.5 ml of 10 mM PBS buffer and 1 ml of KI solution, and the reaction was carried out in the dark
217	at 28°C for 1 h. The H_2O_2 content of the sample was determined by spectrophotometric
218	(GENESYS 180 UV-Vis, Thermo Sci) at 390 nm to determine the H ₂ O ₂ content by measuring
219	the absorbance (Jessup et al., 2018).
220	.Subcellular localisation staining of plant leaf tissues for O_2^- and H_2O_2 was carried out with
221	a slight modification of the method of Romero Puertas et al. (2004).
222	For histochemical detection of analytical O ₂ ⁻ : fresh leaves were washed with distilled water,
223	evacuated, and placed in 10 mL centrifuge tubes containing 0.1% (w/v) nitrogen blue tetrazolium
224	(NBT, pH 7.8) for 24 h. Leaves were rinsed with distilled water, 95% ethanol was added, and
225	then boiled in a boiling water bath for 40 min, destained, and photographed.
226	Analysis Histochemical detection of H ₂ O ₂ : Fresh leaves were washed with distilled water,
227	evacuated, and soaked in 1 mg/L diaminobenzidine 13,3'-tetrahydrochloride (DAB, pH 3.8) in a
228	10 mL plastic centrifuge tube for 24 h. Leaves were rinsed with distilled water, then boiled in 95%
229	(v/v) ethanol for 40 min, decoloured, and photographed.
230	Histochemical staining of leaf cell death was referred to Martina Schraudner et al. (1998).
231	The treated leaves were washed with distilled water, and the leaves were immersed in 0.25%
232	(w/v) Evans blue solution for 24 h . The leaves were removed, rinsed with distilled water and
233	placed in boiling anhydrous ethanol:glycerol (9:1) for 30 min, and chlorophyll was removed until
234	the underside of the leaf appeared white and photographed.
235	
236	Determination of the soluble protein content
237	To determine the soluble protein content, the method of Bradford (1976) was used with the
238	Caumas Brilliant Blue G-250 staining method, 0.5 g of the sample was added to 10 ml of 0.05
239	mol/L pre-cooled phosphate buffer (pH 7.8) and ground in liquid nitrogen, and centrifuged at
240	$12,\!000 \times g$ at 4°C for 20 min, and the supernatant was the crude protein extract. The protein
241	content was determined by adding 1 ml of enzyme solution to 5 ml of koammas brilliant blue
242	solution and then shaking well, and the absorbance value at 595 nm was measured after 2 min of



243	reaction.
244	
245	Statistical analyses
246	Using Excel 2016 for statistical analysis and SPSS 25.0 for further examination, one-way
247	ANOVA and Duncan's method were employed for conducting ANOVA and multiple
248	comparisons. The outcomes were presented as mean $(X) \pm \text{standard error (SE)}$. Graphs were
249	created using Origin 2018 software, with distinct lowercase letters denoting significant
250	differences between treatments ($P < 0.05$).
251	
252	Results
253	Effect of salt stress on morphological indexes at the tillering stage of rice and regulation by
254	Pro-Ca
255	From the experimental results, it was found that salt stress negatively affected the tillering ability
256	of both rice varieties (Fig. 1a and b), and the number of tillers decreased by 30.0%~44.43% and
257	12.52%~33.35% in Xiangliangyou900 and Huanghuazhan, respectively, from the 7th to the 35th
258	days (Fig. 1c and d). The number of main stem leaves of Xiangliangyou900 decreased by 9.09%,
259	5.72%, 15.00%, and 2.04% on the 14th, 21st, 28th, and 35th days, respectively, and that of
260	Huanghuazhan decreased by 6.90%, 21.05%, 11.11%, 12.20%, and 8.16% on the 7th, 14th, 21st,
261	28th, and 35th days, respectively, after NaCl stress (Fig. 1e and f).
262	The plant heights of Xiangliangyou900 and Huanghuazhan were reduced by 4.07%~11.60%
263	and 10.31%~28.10%, respectively, compared with the control from the 7th to the 35th days after
264	salt stress (Table 1). The first tiller length of both varieties was reduced by 2.34% to 29.22% and
265	13.02% to 38.46%, respectively, compared with the CK from the 7th to the 35th days after salt
266	stress (Table 1). Compared with the main stem and the first tiller, the effect of NaCl stress on the
267	length of the second tiller was more significant, and the length of the second tiller of
268	Xiangliangyou900 and Huanghuazhan was reduced by 12.70%~31.86% and 19.60%~39.38%
269	from the 7th to the 35th d after salt stress, respectively (Table 1). The stem base width of the



main stem of Xiangliangyou900 was reduced by 16.45% to 25.68% from 7th to 35th days after 270 salt stress compared with the control, and that of Huanghuazhan was reduced by 17.41% to 38.99% 271 from 7th to the 35th days after NaCl stress (Table 1). Salt stress also significantly reduced the 272 stem base width of the first and second tillers of both varieties. Compared with the CK treatment, 273 the stem base width of the first tiller was reduced by 16.42%~44.02% and 7.74%~43.86%, and 274 the stem base width of the second tiller was reduced by 13.63%~36.32% and 26.62%~43.46%, 275 276 respectively, in Xiangliangyou900 and Huanghuazhan from the 7th to the 35th days after the salt stress (Table 1). 277 The main stem leaf area of Xiangliangyou900 was reduced by 19.83% to 41.70% from the 278 7th to the 35th days after salt stress compared with the control, and the main stem leaf area of 279 Huanghuazhan was reduced by 29.74% to 45.76% from the 7th to the 35th days (Table 2). 280 281 Compared with the control, the leaf area of the first tiller decreased by 36.83% to 60.23% and 30.66% to 73.22%, and the leaf area of the second tiller decreased by 30.63% to 43.52% and 282 43.26% to 65.35%, respectively, in Xiangliangyou900 and Huanghuazhan from the 7th to the 283 35th days after salt stress (Table 2). In addition, compared with the CK, the root length of 284 Xiangliangyou900 decreased by 25.73% to 48.08% from the 7th to the 35th days after salt stress. 285 and the root length of Huanghuazhan decreased by 15.75% to 34% from the 7th to the 35th days 286 after NaCl stress, which were significant differences (Table 2). 287 As shown in Table 3, the aboveground dry weight of the main stems of Xiangliangyou900 288 and Huanghuazhan decreased by 27.53% to 62.62% and 29.77% to 52.86%, respectively, 289 compared with the control from the 7th to the 35th d after NaCl stress. Compared with CK, the 290 dry weight of the first tiller decreased by 9.86% to 63.22% and 30.39% to 65.68%, and the dry 291 weight of the second tiller decreased by 8.44% to 37.46% and 20.99% to 65.03%, respectively, 292 in Xiangliangyou900 and Huanghuazhan from the 7th to the 35th days after salt stress (Table 3). 293 In addition, the root dry weight of Xiangliangyou900 decreased by 34.32% to 70.88% from the 294 7th to the 35th days after NaCl stress compared with the control, and which of Huanghuazhan 295 decreased by 36.33% to 70.21% (Table 3). 296



297	We can see from Fig. 1 that exogenous foliar application of Pro-Ca effectively alleviated
298	the inhibitory effect of NaCl stress on the growth parameters of the two varieties. Foliar
299	application of Pro-Ca under NaCl stress increased the number of tillers by 14.32% to 59.99%
300	and 42.86% to 100.04% in Xiangliangyou 900 and Huanghuazhan, respectively, from the 7th to
301	the 35th days. Compared with S treatment, the number of main stem leaves of both rice varieties
302	increased significantly in Pro-Ca+S treatment, where the number of main stem leaves of
303	Xiangliangyou900 increased by 4.16%, 23.33%, 3.03%, 23.53%, and 6.25%, respectively, and
304	the number of main stem leaves of Huanghuazhan increased by 11.11%, 10.00%, 15.62%, 19.44%
305	and 17.78%, respectively (Fig. 1e and f).
306	As shown in Table 1, compared with the control, foliar spraying Pro-Ca significantly
307	reduced the plant height of the two rice varieties, in which the plant height of Xiangliangyou900
308	was reduced by $5.00\%\sim15.00\%$ and Huanghuazhan's plant height was reduced by $3.51\%\sim20.42\%$
309	from the 7th to the 35th days. The spraying of Pro-Ca reduced the first tiller length of
310	Xiangliangyou900 and Huanghuazhan by $8.77\% \sim 30.02\%$ and $9.90\% \sim 26.17\%$ from the 7th to the
311	35th days, respectively, and the second tiller length of the two varieties was reduced by
312	$6.91\%\sim22.69\%$ and $6.52\%\sim38.96\%$ (Table 1). Compared with the S treatment, foliar spraying
313	Pro-Ca under NaCl stress significantly alleviated the stem base width of each tiller position in
314	both rice, in which the main stem basal width, first tiller basal width and second tiller basal width
315	of Xiangliangyou900 were increased by 15.24%~44.07%, 30.87%~45.30%, and
316	10.58%~48.32%, respectively, from the 7th to the 35th days, the stem base width of each tiller
317	position increased by 23.84%~47.56%, 11.19%~76.13%, and 36.28%~76.96%, respectively
318	(Table 1). Compared with S treatment, foliar spraying Pro-Ca under NaCl stress increased the
319	leaf area of main stem, first tiller leaf area and second tiller leaf area of Xianglaingyou900 by
320	$30.50\% \sim 65.47\%$, $30.11\% \sim 165.73\%$, and $38.19\% \sim 115.18\%$ from the 7th to the 35th days,
321	respectively, and the leaf area of each tiller position of Huanghuazhan increased by
322	$19.78\% \sim 81.97\%$, $74.45\% \sim 329.32\%$, and $69.65\% \sim 204.06\%$, respectively (Table 2). The root
323	lengths of Xiangliangyou900 and Huanghuazhan under Pro-Ca+S treatment increased by 13.43%



324	to 52.29% and 8.37% to 26.09%, respectively, from the 7th to the 35th days compared with that
325	of S treatment, and the differences were significant (Table 2). As shown in Table 3, compared
326	with NaCl, the main stem dry weight, first tiller dry weight and second tiller dry weight of Pro-
327	Ca+S treatment of Xiangliangyou900 increased by 35.75%~137.61%, 2.51%~152.69%, and
328	28.29%~54.31%, respectively, and the dry weights of each tiller position of Huanghuazhan
329	increased by 21.60%~88.90%, 61.33%~174.87%, and 61.22%~205.01%, respectively.
330	Compared with the S treatment, foliar spraying of Pro-Ca alleviated the suppression of below-
331	ground biomass by NaCl stress, and the root dry weight of Xiangliangyou900 increased by
332	23.64%~61.74% and that of Huanghuazhan increased by 39.22%~139.70% from the 7th to the
333	35th days (Table 3).
334	
335	Effect of salt stress on antioxidant enzymes in rice leaves at each tiller position at tillering
336	stage and regulation by Pro-Ca
337	Compared with the control, the SOD activity of the main stem leaves of Xiangliangyou900
338	increased by 6.88% to 31.25% from the 7th to the 35th days after salt treatment, and the SOD
339	activity of the main stem leaves of Huanghuazhan decreased by 10.86% and 9.81% on the 7th
340	and 14th days, respectively, and increased by 13.00% to 24.32% from the 21st to the 35th days
341	(Fig. 2a and b). The SOD activity of the first tiller leaves of Xiangliangyou900 decreased by
342	12.55% and 9.39% on the 7th and 14th days after salt treatment, respectively, and that of the first
343	tiller leaves of Huanghuazhan decreased by 14.48% on the 7th day, and did not show any
344	significant difference compared with the control in the following days (Fig. 2c and d). Compared
345	with the control, the SOD activity of the second tiller leaves of Xiangliangyou900 was
346	significantly increased by 22.51% on the 7th day but did not change significantly from the 14th
347	to the 35th days after NaCl treatment, however, the SOD activity of the second tiller leaves of
348	Huanghuazhan decreased by 3.17% to 32.22% from the 7th to the 35th days(Fig. 2e and f).
349	As can be seen from Fig. 3a and b, compared with the control, salt stress reduced the CAT
350	activity of main stem leaves of Xiangliangyou900 by 9.03% to 9.63% from the 7th to the 28th



351	days, and that of Huanghuazhan main stem leaves by 1.89% to 16.99% from the 14th to the 35th
352	days. Meanwhile, as shown in Fig. 3, NaCl stress reduced the CAT activity of the second tiller
353	leaves of Xiangliangyou900 by 0.50% to 32.15% from the 7th to the 35th days, respectively, and
354	that of Huanghuazhan's first tiller leaves by 3.90% to 8.54% from the 21st to 35th days after salt
355	stress, and that of the second tiller leaves by 0.78% to 5.62%.
356	NaCl stress reduced the POD activity of main stem leaves of Xiangliangyou900 by 2.83%
357	and 13.67% on the 7th and 14th days, respectively (Fig. 4a), and that of Huanghuazhan by 13.01%
358	and 6.55%, respectively, compared with the control (Fig. 4b). In addition, salt stress reduced the
359	POD activity of the first tiller leaves of Xiangliangyou900 by 10.81% and 8.82% on the 7th and
360	14th days, and increased it by 5.12% and 24.40% on the 21st and 28th days, respectively,
361	compared with the CK (Fig. 4c). The POD activity of the first tiller leaves of Huanghuazhan was
362	reduced by 5.71% to 10.71% from the 7th to the 21stdays and increased by 77.45% and 66.59%
363	at the 28th and 35th days, respectively (Fig. 4d). Under salt stress, the POD activities of the
364	second tiller leaves of Xiangliangyou900 and Huanghuazhan were reduced by 3.69%~44.75%
365	and 4.67%~26.11%, respectively, compared with the control from the 7th to the 35th days (Fig.
366	4e and f).
367	The results in Fig. 5a and b showed that NaCl stress increased increased the APX activity of
368	main stem leaves of Xiangliangyou900 by 9.02% to 89.09% from the 14th to the 35th days, and
369	decreased the APX activity of Huanghuazhan's by 56.24% and 7.31% on the 7th and 14th days,
370	respectively, and increased it by 8.55% to 16.30% from the 21st to 35th days. Compared with the
371	CK treatment, the APX activity of the first tiller leaves of Xiangliangyou900 and Huanghuazhan
372	decreased by $12.80\% \sim 54.26\%$ and $3.38\% \sim 50.04\%$, respectively, from the 7th to the 35th days
373	after NaCl stress (Fig. 5c and d), and that of the second tiller leaves of the two varieties
374	decreased by $5.62\% \sim 34.90\%$ and $2.74\% \sim 33.33\%$, respectively (Fig. 5e and f).
375	Under Pro-Ca+S treatment, the SOD activity of main stem leaves of Xiangliangyou900
376	significantly increased by 3.08% to 18.45% from the 7th to the 35th days, and that of
377	Huanghuazhan was only increased by 4.76% and 0.09% on the 7th and 14th days, respectively



(Fig. 2a and b). The SOD activity of the first tiller leaves of both varieties were increased by 1.93% 378 to 21.78% and 4.14% to 31.35% from the 7th to the 35th days after spraying with Pro-Ca 379 380 compared to S treatment, respectively (Fig. 2c and d). In addition, it can be seen from Fig. 2e and f that under salt stress, spraying Pro-Ca increased the SOD activity of the second tiller leaves of 381 Xiangliangyou900 by 11.64% and 10.13% on the 28th and 35th days, respectively, and that of 382 Huanghuazhan by 5.16%~23.86% from the 21st to the 35th days. 383 384 Compared with salt stress, spraying Pro-Ca before salt stress had no significant effect on the CAT activity of leaves of all tillers of Xiangliangyou900, in which, it increased the CAT activity 385 of leaves of the main stem of Huanghuazhan by 6.18%~17.17% from the 14th to the 28th days, 386 respectively. Compared with the S treatment, spraying Pro-Ca increased the CAT activity of the 387 first tiller leaves of Huanghuazhan by 2.37%~7.12% from the 14th to the 28th days, respectively 388 (Fig. 3). 389 As can be seen in Fig. 4, the POD activity of the main stem leaves of Xiangtwoyou 900 390 under Pro-Ca+S treatment increased by 2.38% to 51.79% from the 7th to the 28th days, and that 391 of the main stem leaves of Huanghuazhan increased by 8.27% to 26.63% from the 7th to the 21st 392 days, as compared with that of S treatment. The POD activities of the first tiller and second tiller 393 leaves of Xiangliangyou 900 under Pro-Ca+S treatment increased by 0.45% to 12.84% and 0.83% 394 to 23.74%, respectively, from the 7th to the 35th days (Fig. 4c and e). In addition, the spraying of 395 Pro-Ca increased the POD activity of the first tiller leaves of Huanghuazhan under salt stress by 396 4.84% to 26.07% from the 7th to the 28th days after salt stress, and that of the second tiller 397 leaves by 9.82% to 34.71% from the 7th to the 35th days (Fig. 4d and f). 398 Compared with NaCl stress alone, spraying Pro-Ca under NaCl stress increased the APX 399 activity of main stem leaves of Xiangliangyou900 by 8.48% to 38.46% from the 7th to the 35th 400 days (Fig. 5a). The APX activity of main stem leaves of Huanghuazhan increased by 1.10% to 401 57.35% from the 7th to the 35th days, but not significantly (Fig. 5b). Spraying Pro-Ca under 402 NaCl stress increased the APX activity of the first tiller leaves of Xiangliangyou900 and 403 Huanghuazhan by 6.15%~79.71% and 19.02%~103.47%, respectively, from the 7th to the 35th 404



405	days (Fig. 5c and d). The APX activity of the second tiller leaves of Xiangliangyou900 was
406	increased by 2.38%~40.31% from the 14th to the 35th days (Fig. 5e). The activity of the second
407	tiller leaves of Huanghuazhan was increased by 0.15%~51.94% from the 7th to the 28th days
408	(Fig. 5f).
409	
410	Effect of salt stress on antioxidant enzymes in rice stems at each tiller position at tillering
411	stage and regulation by Pro-Ca
412	Compared with the CK treatment, the SOD activity of the main stem of Xiangliangyou900
413	decreased by 7.64% to 30.56% from the 7th to the 35th days after NaCl treatment, and that of the
414	main stem of Huanghuazhan decreased by 1.43% to 13.72% from the 21st to the 35th days (Fig.
415	6a and b). The SOD activity in the stem of the first tiller of Xiangliangyou900 decreased by 3.81%
416	to 11.08% from the 7th to the 35th days after NaCl treatment, and that in the stem of the second
417	tiller decreased by 1.88% to 32.36% from the 7th to the 28th days (Fig. 6c and e). NaCl stress
418	reduced the SOD activity in the stem of the first tiller of Huanghuazhan by 4.98% to 52.21%
419	from the 21st to the 35th days, and reduced the SOD activity in the stem of the second tiller by
420	3.27% to 26.61% from the 7th to the 35th days, respectively (Fig. 6d and f).
421	Fig. 7 shows that NaCl stress reduced the main stem CAT activity of Xiangliangyou900 by
422	5.85% to 25.60% from the 7th to the 28th days. The CAT activity of Huanghuazhan decreased
423	by 6.67% and 26.94% on the 14th and 35th days, respectively, and increased but not
424	significantly at the 7th, 21st, and 28th days. The CAT activity of the first tiller stems of
425	Xiangliangyou900 decreased by 3.10% to 26.99% from the 21st to the 35th days, and that of
426	Huanghuazhan decreased by 3.14%, 3.27%, and 45.44% at the 7th, 21st, and 35th, respectively
427	(Fig. 7c and d). NaCl stress reduced the CAT activity in the stem of the second tiller of
428	Xiangliangyou900 by 8.69% to 38.70% from the 7th to the 28th days after NaCl stress, and that
429	of Huanghuazhan by 10.25% to 37.52% from the 7th to the 35th days (Fig. 7e and f).
430	Compared with their respective CKs, the POD activity of main stem of Xiangliangyou900
431	under NaCl stress increased by 5.02% to 44.88% from the 7th to the 35th days, and that of



432	Huanghuazhan increased by 7.39% to 11.37% from the 7th to the 21st days, and decreased by
433	18.49% and 35.13% at the 28th and 35th, respectively (Fig. 8a and b). The POD activity of the
434	first tiller stems of Xiangliangyou900 decreased by 5.51%% to 45.94% in 7~35 days under salt
435	stress (Fig. 8c). The POD activity in the stem of the first tiller of Huanghuazhan decreased by
436	2.64% to 10.67% from the 7th to the 28th days, and increased by 4.11% at the 28th day (Fig. 8d).
437	Compared with the control, salt stress reduced the POD activity in the stem of the second tiller
438	by 2.48% to 25.92% and 2.68% to 25.68% in the stem of the second tiller of Xiangliangyou900
439	and Huanghuazhan, respectively, from the 7th to the 35th days (Fig. 8e and f).
440	The main stem APX activity of Xiangliangyou900 decreased by 12.12% to 39.98% from
441	the 7th to the 35th days after NaCl stress, and that of Huanghuazhan decreased by 2.65% to
442	34.28%, respectively (Fig. 9a and b). The APX activity in the stem of the first and second tillers
443	of both varieties decreased by 2.57% to 49.67% and 5.29% to 64.09%, 21.47% to 63.03% and
444	21.35% to 34.96%, respectively, from the 7th to the 35th days after salt stress (Fig. 9).
445	Compared with the S treatment, spraying Pro-Ca under salt stress increased the SOD
446	activity of the main stem of Xiangliangyou900 by 13.66% to 65.88% from the 7th to the 28th
447	days, and that of the main stem of Huanghuazhan by 2.05% to 7.96% from the 21st to the 35th
448	days (Fig. 6a and b), the SOD activities in the stem of the first tiller increased by $6.34\% \sim 51.90\%$
449	and 6.91%~97.69% from the 7th to the 35th days in Xiangliangyou900 and Huanghuazhan,
450	respectively (Fig. 6c and d). Spraying Pro-Ca under salt stress increased the SOD activity of the
451	second tiller stems of Xiangliangyou900 by 2.68% to 123.51% from the 14th to the 35th days,
452	and that of the second tiller stems of Huanghuazhan by 3.27% to 26.61% from the 7th to the 35th
453	days (Fig. 6e and f).
454	Compared with the S treatment, the CAT activities of main stems sprayed with Pro-Ca
455	Xiangliangyou900 and Huanghuazhan under salt stress increased by 5.72% to 92.16% and 6.04%
456	to 40.09% from the 7th to the 35th days, respectively (Fig. 7a and b). Foliar spraying of Pro-Ca
457	also effectively increased the CAT activity in the first tiller stems of two rice varieties under salt
458	stress, which was increased by 0.83%~30.00% at the 7~28 days in Xiangliangyou900 and by



3.59%~95.57% at the 7~35 days in Huanghuazhan (Fig. 7c and d). CAT activity in the second 459 tiller stems of both varieties was increased by 0.93% to 66.28% and 7.57% to 65.13% under Pro-460 461 Ca+S treatment compared to S, respectively (Fig. 7e and f). Compared with the S treatment alone, spraying Pro-Ca under salt stress increased the main 462 stem POD activity by 5.89% to 51.39% and 4.76% to 33.32% from the 7th to the 35th days in 463 Xiangliangyou900 and Huanghuazhan, respectively. The POD activity in the first and second 464 tiller stems of both varieties increased by 0.18% to 24.07% and 13.49% to 98.03%, 14.51% to 465 31.29% and 4.15% to 33.22%, respectively, from the 7th to the 35th days (Fig. 8). 466 Compared with salt stress alone, spraying Pro-Ca under NaCl stress increased the main stem 467 APX activity of Xiangliangyou900 by 11.43% to 87.63% from the 7th to the 35th days, and that 468 of Huanghuazhan by 10.98% to 31.58%, respectively (Fig. 9a and b). It increased the APX 469 activity of the first and second tiller stems of both varieties by 0.85% to 103.18% and 10.74% to 470 99.20%, 4.11% to 34.69% and 18.49% to 83.24%, respectively, from the 7th to the 35th days 471 (Fig. 9). 472 473 Effect of salt stress on membrane damage index in rice leaves at each tiller position at 474 tillering stage and regulation by Pro-Ca 475 Salt stress significantly increased the MDA content of leaves of two rice varieties compared with 476 the control (Fig. 10). Among them, the MDA contents of main stem, first tiller and second tiller 477 leaves of Xiangliangyou900 increased by 13.04%~54.61%, 0.58%~79.40%, and 3.33%~39.59%, 478 respectively, from the 7th to the 35th days after salt stress (Fig. 10a, c, and e), the MDA content 479 of the leaves of each tiller position of Huanghuazhan increased by 10.62% to 127.93%, 13.25% 480 to 75.94%, and 6.01% to 64.67%, respectively, from the 7th to the 35th days after salt stress (Fig. 481 10b, d, and f). In addition, as seen in Fig. 11, salt stress increased the H₂O₂ content of the main 482 stem leaves of Xiangliangyou900 and Huanghuazhan by 4.57%~38.51% and 3.88%~21.84% at 483 7~35 days, and the H₂O₂ content of the first tiller leaves of two kinds of rice by 5.24%~39.80% 484 and 0.44%~41.06%. The H₂O₂ content of the second tiller leaves of the two varieties was 485



increased by 16.46%~28.88% and 8.37%~58.75% at 7~35 days, respectively (Fig. 11). 486 Compared with S treatment, foliar spraying of Pro-Ca before salt stress decreased the MDA 487 content of main stem leaves of Xiangliangyou900 and Huanghuazhan by 0.53%~27.59% and 488 8.65%~25.00%, respectively (Fig. 10a and b), and that of the first tiller leaves of the two 489 varieties under the same conditions by 17.24%~26.80% and 2.39%~29.12%, respectively (Fig. 490 10c and d), from the 7th to the 35th days. The MDA contents of second tiller leaves of the two 491 492 varieties under the same conditions were decreased by 0.09\%\~34.98\% and 1.60\%\~26.93\%, respectively (Fig. 10e and f). Spraying Pro-Ca before salt stress reduced the H₂O₂ content of 493 main stem leaves of Xiangliangyou900 and Huanghuazhan by 0.63%~22.87% and 494 5.91%~42.28%, respectively, from the 7th to the 35th days (Fig. 11a and b). Spraying Pro-Ca 495 reduced the H₂O₂ content of the first tiller leaves of Xiangliangyou900 by 4.03%~31.42% and 496 497 that of Huanghuazhan by 7.56%~26.04%, respectively at 7~35 days (Fig. 11c and d), and reduced the H₂O₂ content of the second tiller leaves of both rice varieties by 5.49%~29.10% and 498 8.23%~34.97% (Fig. 11e and f). The staining test revealed that there were more spots on the 499 leaves of both rice varieties under salt stress, and the spots on the tiller leaves were larger in area 500 and darker in color. In contrast, the area of spots on the leaves of the treatments sprayed with 501 Pro-Ca before salt stress decreased and became lighter in color (Fig. 12). 502 503 Effect of salt stress on membrane damage index in rice stems at each tiller position at 504 505 tillering stage and regulation by Pro-Ca Compared with the control, NaCl stress increased the MDA content of the main stem of 506 Xiangliangyou900 and Huanghuazhan by 14.85%~115.59% and 39.38%~95.94%, respectively 507 (Fig. 13a and b), and that of the first tiller stems of Xiangliangyou900 and Huanghuazhan by 508 4.98%~90.72% and 9.55%~72.13%, respectively, from the 7th to the 35th days after salt stress 509 (Fig. 13c and d), and the MDA content of the second tiller stems of the two varieties increased 510 by 22.21%~156.61% and 7.98%~61.68%, respectively (Fig. 13e and f). NaCl stress at 0.3% also 511 significantly increased the H₂O₂ content in the stems of both rice varieties (Fig. 14). Among 512



514	6.12% from the 7th to the 35th days, and increased the H_2O_2 content of the main stem of
515	Huanghuazhan by 7.61% to 24.09% from the 7th to the 28th days (Fig. 14a and b). Salt stress
516	increased the H ₂ O ₂ content of the first tiller stem of Xiangliangyou900 by 17.28%, 14.83%, and
517	8.49% at the 7th, 14th, and 35th days, respectively, and that of Huanghuazhan by 0.63%~9.16%
518	at 14~35 days (Fig. 14c and d). In addition, NaCl stress increased the H ₂ O ₂ content in the stem of
519	the second tiller of Xiangliangyou900 by 1.37%~18.04% from the 7th to the 35th days, and there
520	was no significant change on the 28th day, that of Huanghuazhan increased the content by 2.67%
521	and 28.36% on the at the 7th and 21st days (Fig. 14e and f).
522	Compared with S treatment, spraying Pro-Ca before salt stress caused a significant decrease
523	in MDA content, in which the MDA content of the main stems of Xiangliangyou900 and
524	Huanghuazhan decreased by 2.53% to 48.87% and 14.94% to 35.39%, respectively, from the 7th
525	to the 35th days (Fig. 13a and b). The MDA content of the first tiller stems of the two varieties
526	sprayed with Pro-Ca decreased by 3.79%~46.45% and 5.39%~27.28%, respectively, from the
527	7th to the 35th days after salt stress (Fig. 13c and d), compared with S treatment, spraying Pro-
528	Ca decreased the MDA content of the second tiller stems of Xiangliangyou900 and
529	Huanghuazhan by 1.70%~66.09% and 0.32%~37.09% from the 7th to the 35th days, respectively
530	(Fig. 13e and f). Foliar spraying of Pro-Ca before salt stress reduced H ₂ O ₂ in the main stem of
531	Xiangliangyou900 by 7.07%~12.62% from the 7th to the 28th days, and that of Huanghuazhan
532	by 2.79%~14.74% from the 7th to the 35th days (Fig. 14a and b), and reduced the H_2O_2 content
533	in the stem of the first tiller of Xiangliangyou900 and Huanghuazhan by 9.58%~15.60% and
534	3.20%~12.45% from the 7th to the 35th days respectively (Fig. 14c and d). Compared with salt
535	stress, spraying Pro-Ca reduced the H ₂ O ₂ content in the stem of the second tiller of the two rice
536	varieties by 1.35%~33.05% and 7.92%~24.15%, respectively, from the 7th to the 35th days (Fig.
537	14e and f).

them, salt stress increased the H₂O₂ content of the main stem of Xiangliangyou900 by 1.28% to

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Effect of salt stress on soluble protein content in rice leaves at each tiller position at



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541	As can be seen from Fig. 15, the soluble protein content of the main stem leaves of
542	Xiangliangyou900 increased by 0.88% to 2.59% from the 7th to the 28th days after salt stress,
543	and the soluble protein content of the main stem leaves of Huanghuazhan decreased by 1.59%
544	and 2.87% on the 7th and 14th days after salt stress, respectively, and increased by 0.44% to 2.41%
545	from the 21st to 35th d. NaCl stress increased the soluble protein content of the first tiller leaves
546	of Xiangliangyou900 by 0.61% and 5.33% on the 21st and 28th days, respectively, and
547	decreased the soluble protein content of the first tiller leaves of Huanghuazhan by 2.06% to 4.22%
548	from the 7th to the 28th days after NaCl stress (Fig. 15c and d). The soluble protein content of
549	the second tiller leaves of Xiangtwoyou 900 increased by 3.27% and 0.45% on the 7th and 21st
550	after NaCl stress, and decreased that of Huanghuazhan by 0.31% to 4.63% from the 7th to the
551	35th days (Fig. 15e and f).
552	Compared with the S treatment, the soluble protein content of the main stem leaves of
553	Xiangliangyou900 increased by 0.21% to 4.67% from the 7th to the 35th days in Pro-Ca+S
554	treatment, and that of the main stem leaves of Huanghuazhan increased by 1.28% to 1.90% from
555	the 7th to the 21st days (Fig. 15a and b). In addition, spraying Pro-Ca under salt stress increased
556	the soluble protein content of the first tiller leaves of Xiangliangyou900 by 4.30%, 0.25%, and
557	1.94% at the 7th , 14th , and 35th, respectively, and that of Huanghuazhan by $0.57%$ to $4.64%$
558	from the 7th to the 35th days (Fig. 15c and d). Foliar spraying of Pro-Ca increased the soluble
559	protein content of the second tiller leaves of Xiangliangyou900 by 0.55% to 4.38% from the 14th
560	to the 35th days and Huanghuazhan by 0.16% to 5.15% from the 7th to the 35th days after salt
561	stress (Fig. 15e and f).
562	
563	Effect of salt stress on soluble protein content in rice stems at each tiller position at tillering
564	stage and regulation by Pro-Ca
565	As can be seen from the Fig. 16a and b, the soluble protein content of the main stem of
566	Xiangliangyou900 increased by 1.92% to 9.30% from the 14th to the 35th days after salt stress,



and the soluble protein content of the main stem of Huanghuazhan increased by 1.03% to 7.41% from the 7th to the 28th days. Salt stress reduced the soluble protein content of the first tiller stems of Xiangliangyou900 by 0.31%, 5.92%, and 3.83% at the 7th, 28th, and 35th days, and reduced the soluble protein content of the first tiller stems of Huanghuazhan by 1.48% and 1.12% at the 21st and 35th days, respectively (Fig. 16c and d). The soluble protein content of the second tiller stems of Xiangliangyou900 decreased by 0.88% to 12.45% from the 7th to the 35th days after salt stress, and there was no significant difference at the 14th day, while that of the second tiller stems of Huanghuazhan decreased by 4.81% to 12.49% from the 14th to the 35th days (Fig. 16e and f).

Spraying Pro-Ca under salt stress conditions increased the soluble protein content of the

Spraying Pro-Ca under salt stress conditions increased the soluble protein content of the main stems of Xiangliangyou900 and Huanghuazhan by 0.97%~9.13% and 2.05%~15.38%, respectively, from the 7th to the 35th days after NaCl stress (Fig. 16a and b). Compared with the S treatment, the soluble protein content of the first tiller stems of Xiangliang900 under Pro-Ca+S treatment increased by 0.90% to 14.89% from the 7th to the 35th days, but there was no significant difference at the 21st day, and that of the first tiller stems of Huanghuazhan under Pro-Ca+S treatment increased by 3.89% to 22.64% from the 7th to the 35th days (Fig. 16c and d). Foliar spraying of Pro-Ca under NaCl stress increased the soluble protein content of the second tiller stems of Xiangliangyou900 and Huanghuazhan by 8.09% to 26.44% and 2.11% to 17.37%, respectively, from the 7th to the 35th days after salt stress (Fig. 16e and f).

Discussion

Salt stress, as a significant abiotic stressor, can greatly inhibit crop growth and development. It has the ability to trigger the production of ROS, an excess of which can harm cellular functions and even result in cell death. ROS also act as secondary messengers, activating molecular processes in response to environmental stresses. Therefore, maintaining an appropriate level of ROS is crucial for plant survival under stress conditions (Lee et al., 2022). However, excessive salt stress can disrupt the balance of internal ions in cells (Zulfiqar & Ashraf, 2021). High



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concentrations of salt stress also increase the production of reactive oxygen species (ROS), such as mono-linear oxygen (${}^{1}O_{2}$), superoxide (O_{2}^{-}), hydrogen peroxide ($H_{2}O_{2}$), and hydroxyl radicals (OH) (Ding et al., 2022). Plants have evolved complex mechanisms to combat salt stress-induced oxidative stress, such as antioxidant enzymes like superoxide dismutase (SOD), catalase (CAT), and peroxidase (POD). These enzymes are crucial for scavenging ROS, and an elevation in ROS levels triggers an upsurge in the activities of SOD, POD, CAT, and other enzymes during salt stress conditions. (Huihui et al., 2020). Salt stress damage to crops has been confirmed in various crops (Kousar et al., 2021; Jung et al., 2017; Feng et al., 2023). Jian et al. (2022) demonstrated that growth rate was reduced after 1 day of 100 mM NaCl stress in rice, and more severe wilting symptoms appeared on the tips of rice plant leaves after 15 days. 'IR29' was particularly damaged, which contained high levels of ROS. The results of this experiment indicated that the main stem leaves exhibited a stronger antioxidant capacity under salt stress due to increased SOD, POD, and APX activities. CAT activity decreased in all tillers under salt stress, while the main stem demonstrated higher salt tolerance compared to the tillers by enhancing SOD, CAT, and POD enzyme activities and experiencing a less significant reduction in APX. It was speculated that this discrepancy could be attributed to a lower level of stress in the main stem and a greater resilience in the main stem in contrast to the tillers. Yang et al. (2022) suggested that since tillers mature later than the main stem, the main stem consistently maintains a growth and development advantage. Prolonged exposure to stress further reinforces this superiority, leading to unequal competition for C and N resources between the main stem and tillers, ultimately diminishing seed yield (Tilley et al., 2017). Xie et al. (2019) observed that salttolerant wheat under salt stress conditions had higher activities of CAT, POD and APX thus more unfavourable H₂O₂ accumulation and reduced oxidative damage compared to sensitive varieties. Therefore, it is believed that the leaf tiller co-extension law leads to a pre-eminent advantage of temperature and light resource utilisation in the main stem and the first and second tillers, which contributes to the improvement of antioxidant capacity, nutrient accumulation and thus better growth and resistance.



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Foliar spraying of Pro-Ca under salt stress effectively mitigated the damage of salt stress on morphogenesis at the tillering stage of rice, as demonstrated in our previous studies. (Zhang et al., 2023; Zhang et al., 2023; Huang et al., 2023), In line with previous experiments, the results of this study indicated that foliar application of Pro-Ca had a more pronounced impact on tiller development compared to the main stem. Pro-Ca effectively alleviated the oxidative damage caused by salt stress on rice leaves. Foliar spraying of Pro-Ca increased the SOD and APX activities of tiller leaves and stems to a stronger extent than that of the main stems, but the effect on CAT activity of main stem leaves was more pronounced; moreover, Pro-Ca showed a better modulation in the alleviation of membrane damage and in the increase of soluble protein content. On the one hand, compared with the main stem, tillers will be subject to stronger oxidative damage, calcium regulator can increase SOD and APX activities as the first line of defence of the plant against oxidative stress (Shafi et al., 2015), on the other hand, it may be related to the spraying site of Pro-Ca, the treatment of the present experiment was in the seedling stage, and the sampling period was in the tillering stage, and the determined tillers were developed from the axils of the leaves that had been sprayed with Pro-Ca in the previous period, therefore, the regulator may have a better regulatory effect on the site to which it was sprayed. In prior trials, it was discovered that despite not being directly treated with ethylene and chlormequat chloride, oat T1 and T2 tillers exhibited delayed stem growth. This could be attributed to the transition from chlormequat chloride-treated to untreated sections of the plant, a phenomenon also observed in wheat. (Kang et al., 2010; Peltonen-Sainio et al., 2003).

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Conclusion

This experiment showed that salt stress hindered the growth of leaves and stems of rice tillers during the tillering stage. Additionally, the application of Pro-Ca through spraying effectively reduced the oxidative damage caused by salt stress on the tillers. Interestingly, the impact on the tillers was more pronounced compared to the main stems under similar conditions. This study offers new perspectives on the varying effects of salt stress on rice tillers and the regulatory role



648 of Pro-Ca.

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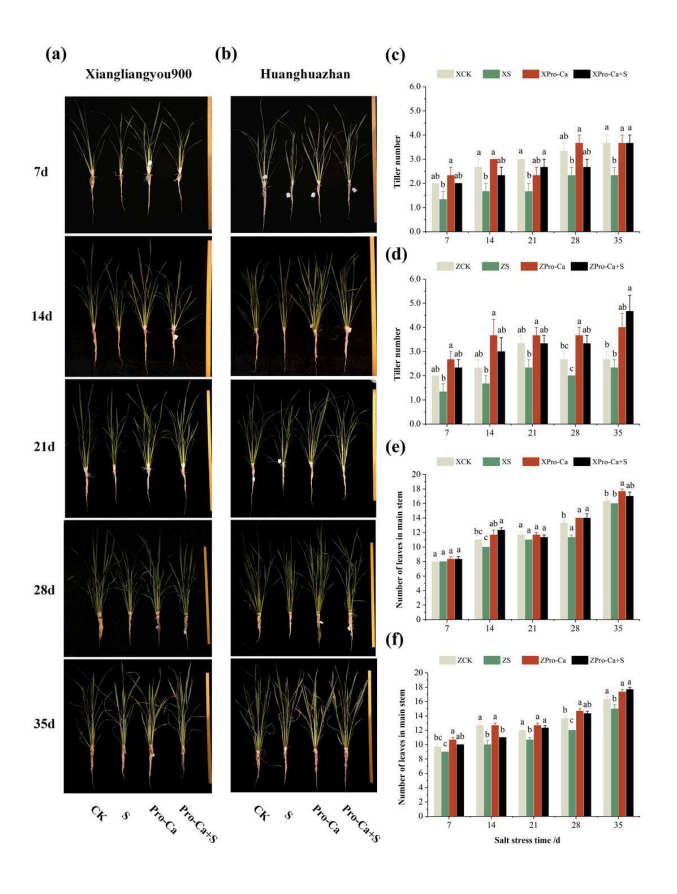
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Figure 1

Effects of Pro-Ca on rice growth under salt stress.

<!--[if !supportLists]-->(a-<!--[endif]-->b) Plant growth. Rice morphology after 7, 14, 21, 28, and 35d of salt stress. Figure (a) shows the form of Xiangliangyou900, and Figure (b) shows Huanghuazhan. From left to right, the plants were treated as follows: CK (distilled water), S (0.3% NaCl), Pro-Ca (100 mg•L⁻¹ Pro-Ca), Pro-Ca+S (100 mg•L⁻¹ Pro-Ca + 0.3% NaCl).(c-d) Tiller numbers of Xiangliangyou900 and Huanghuazhan in the main stem between different treatments.(e-f) Leaf numbers in the main stem. Comparison of tiller numbers and leaf numbers in the main stem between different treatments. Values are means \pm SD (n = 3) and bars indicate SD. Columns with different letters indicate significant difference at P < 0.05 (Duncan's test). Xiangliangyou900: XCK (distilled water), XS (0.3% NaCl), XPro-Ca (100 mg•L⁻¹ Pro-Ca), XPro-Ca+S (100 mg•L⁻¹ Pro-Ca + 0.3% NaCl), Huanghuazhan: ZCK (distilled water), ZS (0.3% NaCl), ZPro-Ca (100 mg•L⁻¹ Pro-Ca), ZPro-Ca+S (100 mg•L⁻¹ Pro-Ca + 0.3% NaCl).

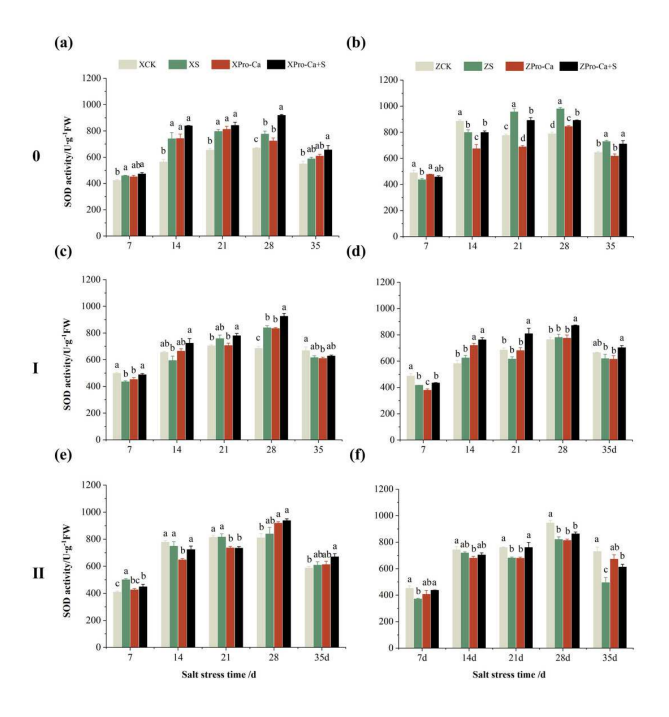






Effect of Pro-Ca on SOD activity of rice main stem (a, b), first tiller (c, d), and second tiller (e, f) leaves under salt stress.

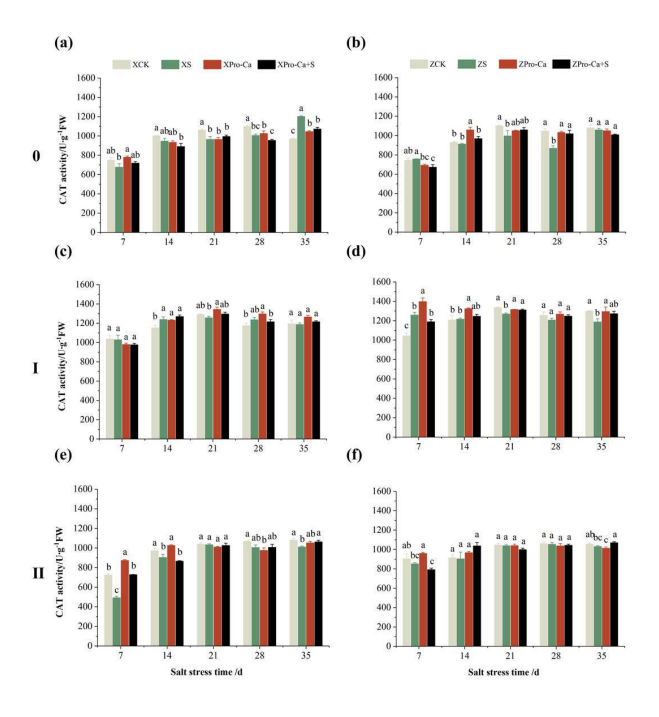






Effect of Pro-Ca on CAT activity of rice main stem (a, b), first tiller (c, d), and second tiller (e, f) leaves under salt stress.

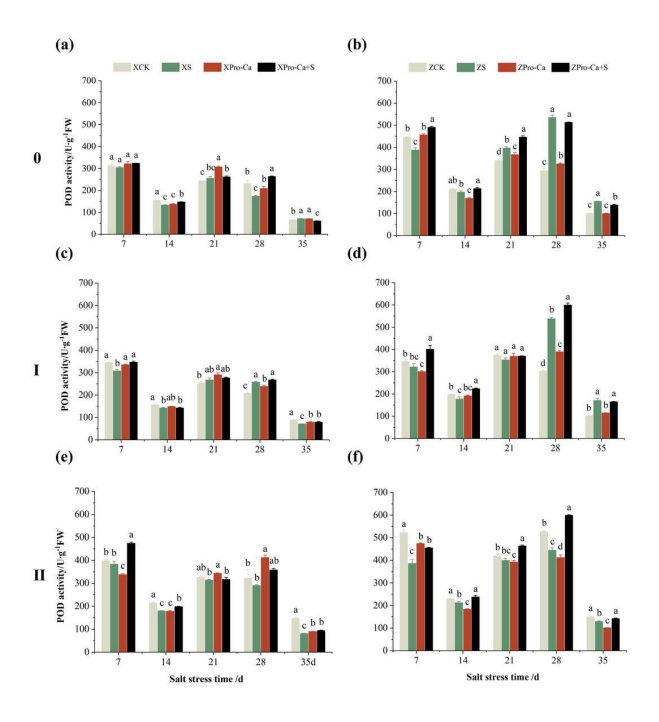






Effect of Pro-Ca on POD activity of rice main stem (a, b), first tiller (c, d), and second tiller (e, f) leaves under salt stress.

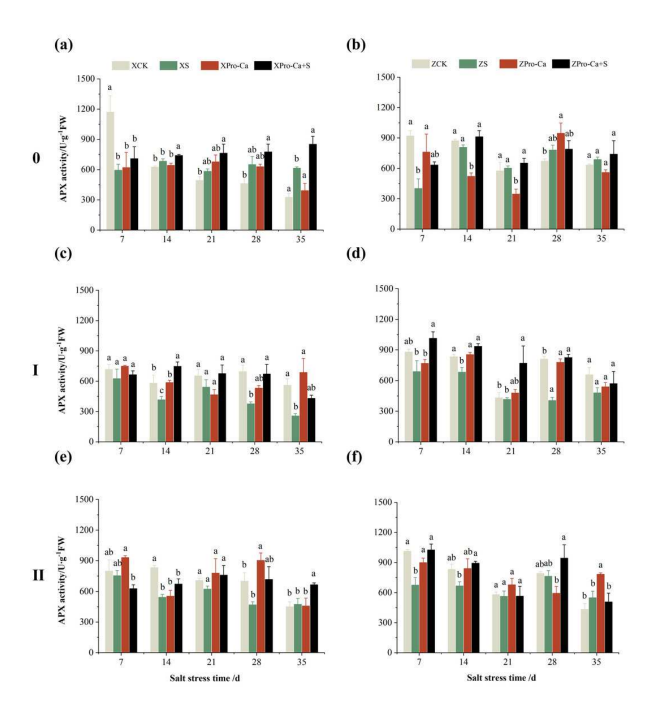






Effect of Pro-Ca on APX activity of rice main stem (a, b), first tiller (c, d), and second tiller (e, f) leaves under salt stress.

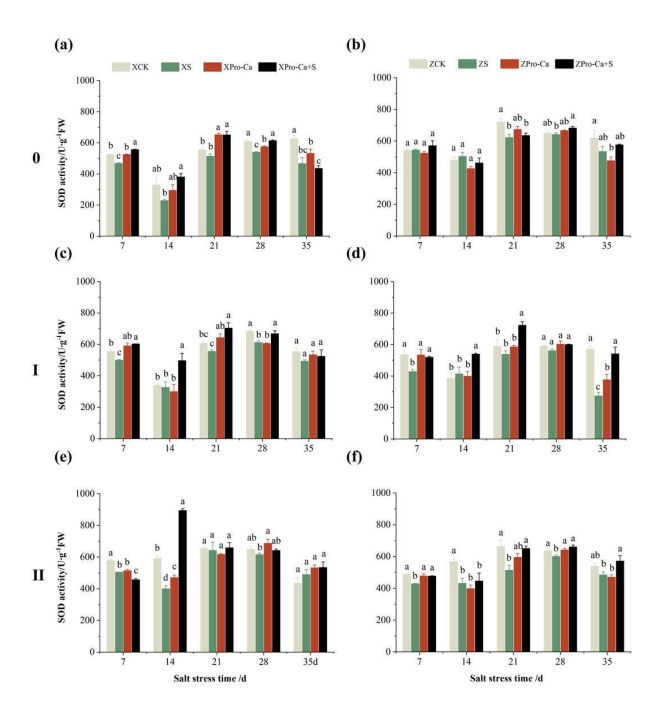






Effect of Pro-Ca on SOD activity of rice main stem (a, b), first tiller (c, d), and second tiller (e, f) stems under salt stress.

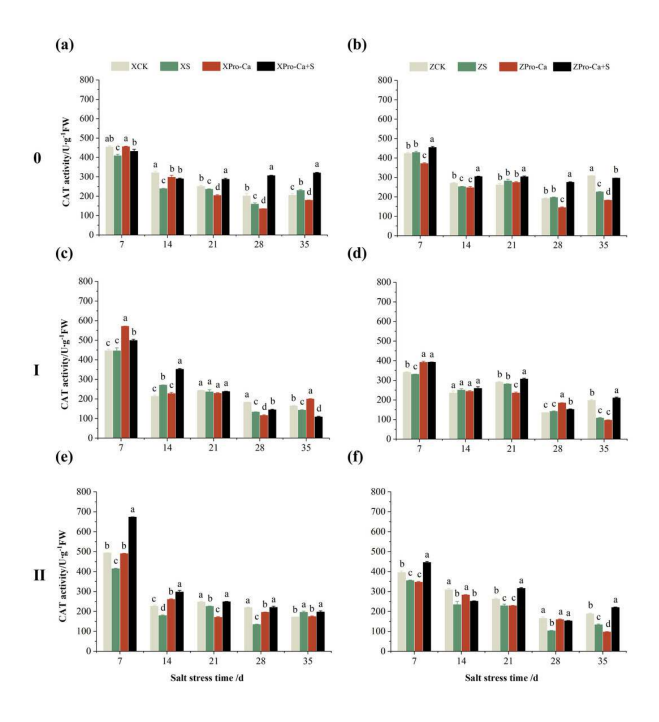






Effect of Pro-Ca on CAT activity of rice main stem (a, b), first tiller (c, d), and second tiller (e, f) stems under salt stress.

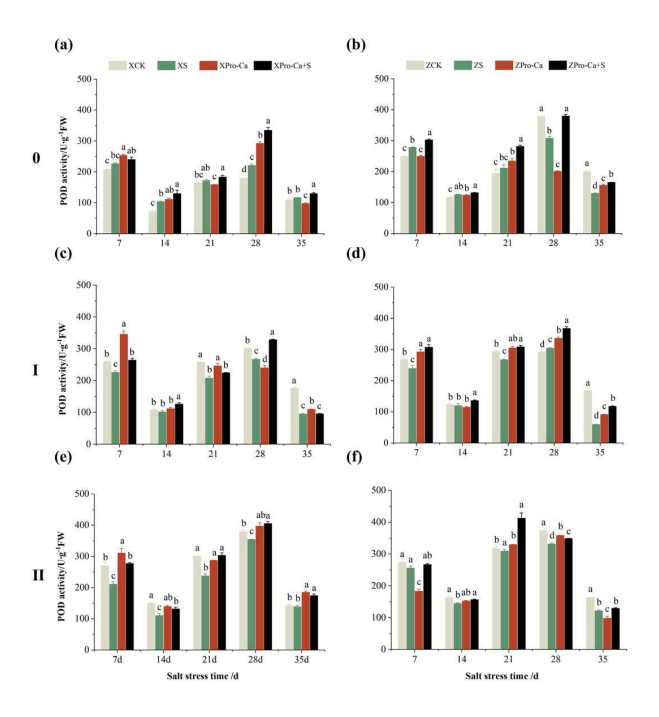






Effect of Pro-Ca on POD activity of rice main stem (a, b), first tiller (c, d), and second tiller (e, f) stems under salt stress.

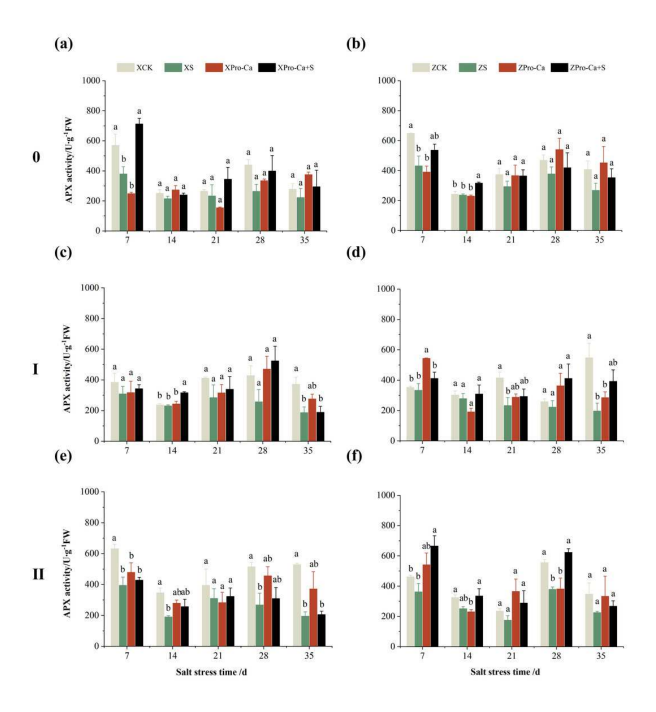






Effect of Pro-Ca on APX activity of rice main stem (a, b), first tiller (c, d), and second tiller (e, f) stems under salt stress.

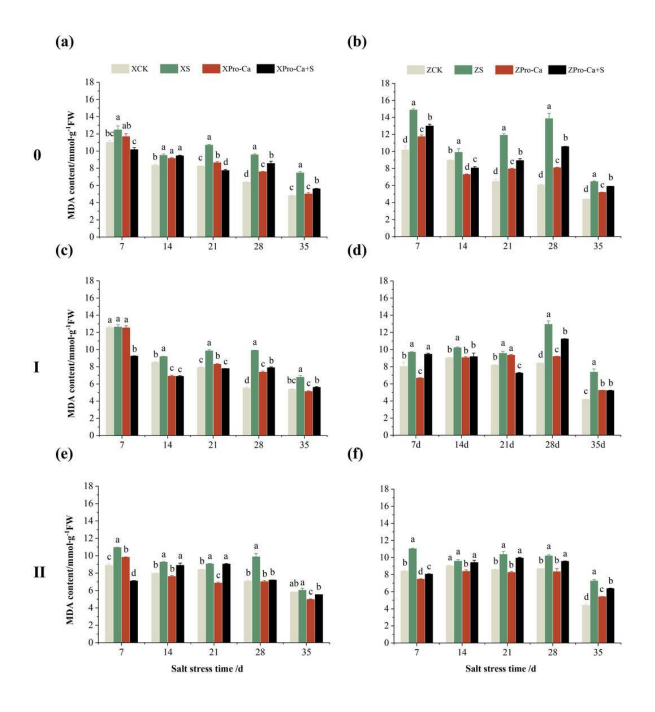






Effect of Pro-Ca on MDA content of rice main stem (a, b), first tiller (c, d), and second tiller (e, f) leaves under salt stress.

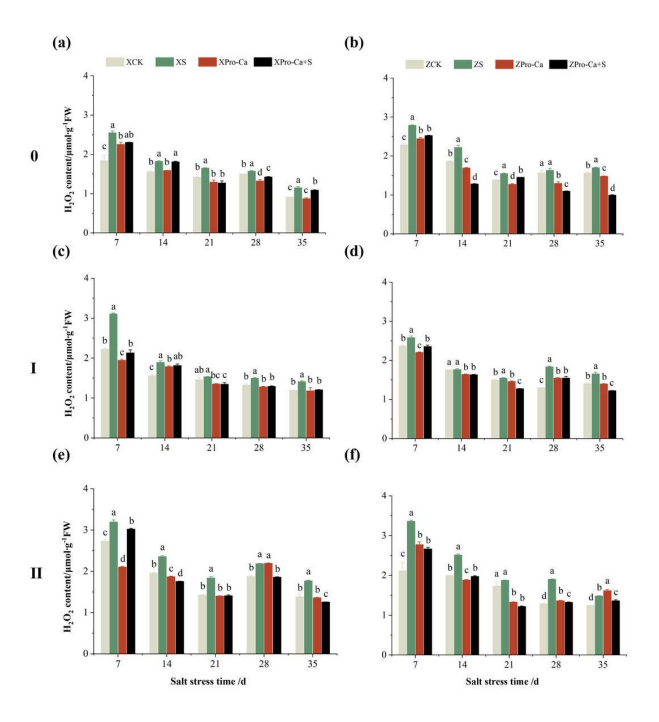






Effect of Pro-Ca on H_2O_2 content of rice main stem (a, b), first tiller (c, d), and second tiller (e, f) leaves under salt stress.



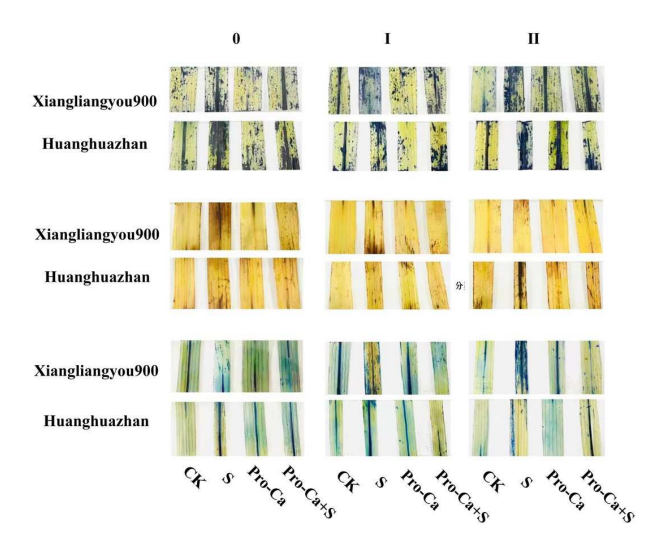




Effect of Pro-Ca on O_2 , H_2O_2 content and cellular activity of rice main stem leaves at each tiller position under salt stress.

From left to right, the three columns represent the tiller stem, the first node position, and the second tiller position. Each image shows the following treatments applied to the plants from left to right: CK (distilled water), S (0.3% NaCl), Pro-Ca (100 mg \cdot L $^{-1}$ Pro-Ca), Pro-Ca+S (100 mg \cdot L $^{-1}$ Pro-Ca + 0.3% NaCl). The spot area represents the degree of stress, and the larger staining area indicates that the more severe stress of leaves.

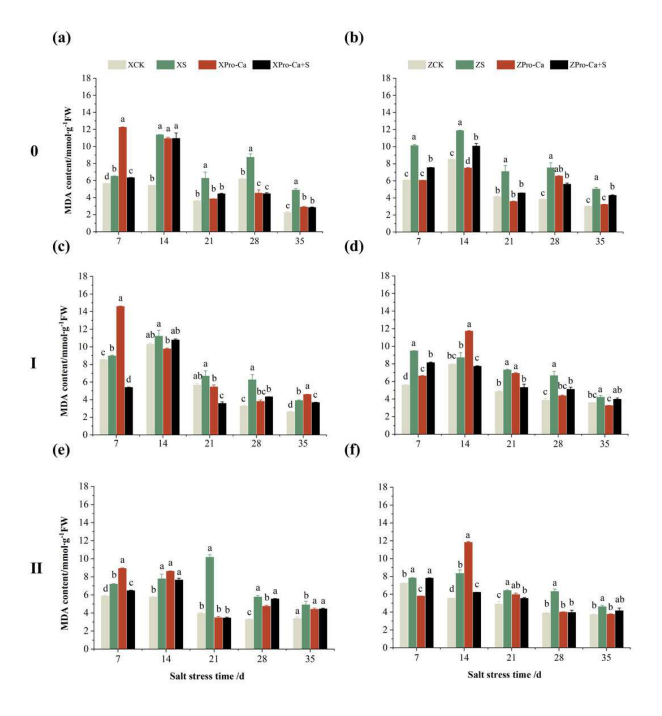






Effect of Pro-Ca on MDA content of rice main stem (a, b), first tiller (c, d), and second tiller (e, f) stems under salt stress.

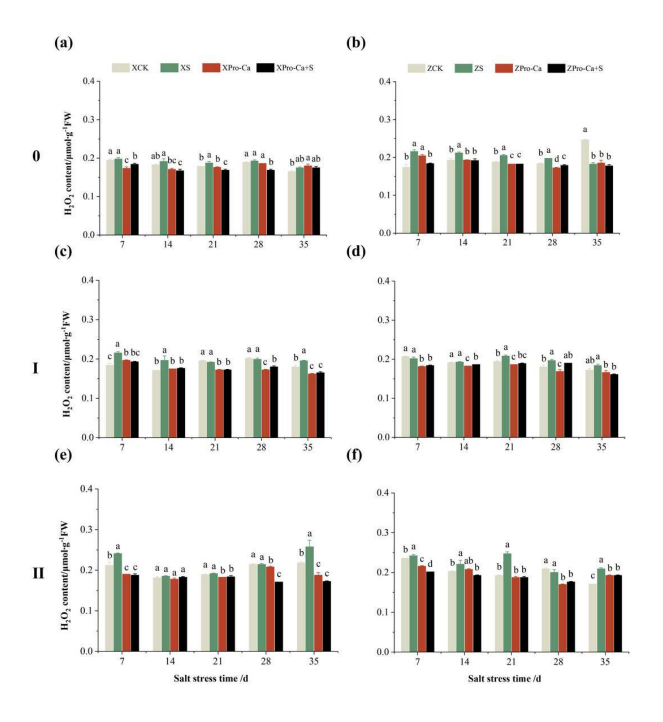






Effect of Pro-Ca on H_2O_2 content of rice main stem (a, b), first tiller (c, d), and second tiller (e, f) stems under salt stress.

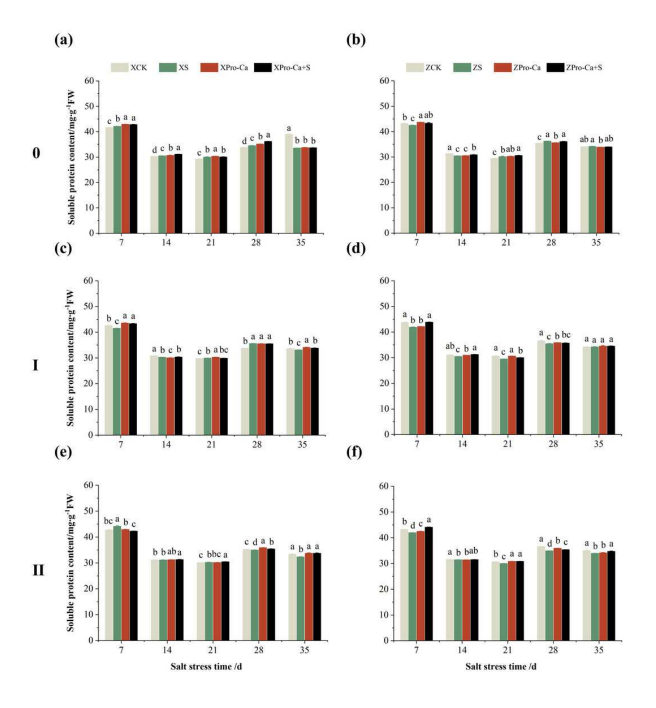






Effect of Pro-Ca on the soluble protein content of rice main stem (a, b), first tiller (c, d), and second tiller (e, f) leaves under salt stress.







Effect of Pro-Ca on the soluble protein content of rice main stem (a, b), first tiller (c, d), and second tiller (e, f) stems under salt stress.



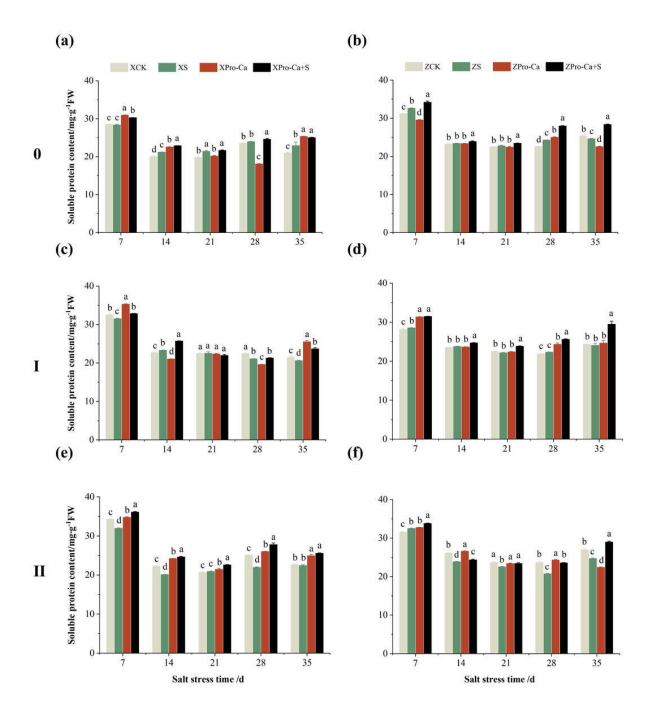




Table 1(on next page)

Effects of Pro-Ca on plant height and stem base width of rice main stem, first tiller, and second tiller stems at the tillering stage under salt stress.

Values described are the means±SE (n=3). Different letters denote significant difference from Duncan's LSD test (p<0.05).



1 Table 1

2 Effects of Pro-Ca on plant height and stem base width of rice main stem, first tiller, and

3 second tiller stems at the tillering stage under salt stress.

Time/ d	Treatment	Plant height/ cm			Stem base width/ mm		
Time/ u		0	Ι	II	0	I	II
	XCK	56.4±0.4a	41.3±0.9a	36.5±1.2a	9.4±0.6b	8.0±0.2a	5.2±0.1b
7	XS	52.4±0.9b	29.2±2.7b	27.3±1.9c	7.1±0.2c	5.6±0.4b	4.5±0.3b
	XPro-Ca	48.7±0.9c	28.9±0.4b	31.7±0.7b	11.2±0.5a	8.3±0.2a	$6.2 \pm 0.2a$
	XPro-Ca+S	51.2±0.4b	29.9±2.0b	35.0±1.0ab	9.4±0.3b	7.3±0.6a	$6.2 \pm 0.3a$
/	ZCK	$63.2\pm2.7a$	42.9±2.7a	45.2±5.0a	9.2±0.4a	$7.9 \pm 0.5 b$	6.4±0.1a
	ZS	45.5±1.6c	26.4±1.7c	$27.4 \pm 1.9b$	$6.3 \pm 0.2b$	$5.2\pm0.2c$	$3.6 \pm 0.3b$
	ZPro-Ca	51.6±0.6b	$34.1 \pm 2.3b$	$31.6 \pm 2.3b$	9.7±1.2a	8.9±0.1a	$6.3\pm0.1a$
	ZPro-Ca+S	56.5±1.3b	$36.5 \pm 1.2ab$	$35.2 \pm 2.0ab$	8.5±0.1a	$7.2\pm0.2b$	$6.0\pm0.4a$
	XCK	$66.5\pm2.1a$	46.9±1.0a	43.6±1.0a	10.8±0.7a	$6.9\pm0.4a$	$6.7 \pm 0.3a$
	XS	58.2±0.2b	45.8±1.6a	29.7±1.5c	$8.1 \pm 0.2b$	$5.8\pm0.1b$	$4.6 \pm 0.5 b$
	XPro-Ca	$56.5 \pm 0.7b$	$41.7 \pm 1.0b$	33.7±3.9bc	12.4±0.5a	$7.3\pm0.4a$	$6.7 \pm 0.1a$
14	XPro-Ca+S	52.3±0.3c	32.6±0.9c	37.4±0.6ab	11.7±0.9a	7.7±0.1a	6.8±0.1a
14	ZCK	67.3±1.9a	$56.3\pm2.0a$	51.8±2.6a	9.9±0.4a	8.0±0.6a	$6.8 \pm 0.2a$
	ZS	53.9±2.5b	$43.7 \pm 1.0b$	36.8±1.8bc	$6.5\pm0.1b$	$4.6 \pm 0.4 b$	$4.3 \pm 0.9b$
	ZPro-Ca	53.5±1.4b	$41.6 \pm 2.8 b$	31.6±2.8c	9.2±0.2a	9.6±0.2a	6.9±0.1a
	ZPro-Ca+S	51.8±3.5b	44.2±2.1b	$41.7 \pm 0.4 b$	9.6±0.6a	$8.2 \pm 0.5a$	$6.8 \pm 0.3a$
	XCK	83.3±1.5a	$58.1 \pm 5.4a$	49.0±0.8a	12.6±0.2a	11.4±0.7a	$7.8 \pm 0.6a$
	XS	73.7±1.2bc	48.0±2.6ab	38.0±1.0c	10.5±0.2b	$6.4\pm0.5c$	$5.0\pm0.6b$
	XPro-Ca	79.2±1.6ab	$43.5 \pm 2.8b$	$44.8 \pm 0.7 b$	12.7±0.3a	$9.8 \pm 0.8 ab$	$7.6\pm0.2a$
21	XPro-Ca+S	71.0±2.6c	41.7±4.8b	41.5±2.1bc	12.1±0.7a	$9.2 \pm 0.5b$	7.4±0.2a
	ZCK	76.0±0.6a	64.0±6.1a	56.6±1.0a	10.5±0.3a	10.0±1.0a	7.9±0.2a
	ZS	68.2±2.2b	55.7±0.3a	41.5±1.5b	$7.8 \pm 0.3b$	$5.6 \pm 0.5 b$	$5.2\pm0.2c$
	ZPro-Ca	73.3±3.2ab	57.7±3.3a	51.1±2.1a	10.9±0.8a	11.1±0.3a	$8.3 \pm 0.2a$
	ZPro-Ca+S	72.3±0.9ab	41.5±4.3b	50.5±2.6a	9.7±0.3a	8.9±0.6a	7.3±0.1b

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11 Table 1 continued

28	XCK	86.8±1.9a	71.6±3.9a	51.6±2.2a	16.0±0.1b	12.1±1.2a	8.5±0.5a
	XS	$83.2 \pm 1.5 ab$	58.8±2.8bc	45.0±1.2bc	11.9±0.5c	$8.2 \pm 0.7a$	5.9±0.6b
	XPro-Ca	81.4±1.7b	62.8±1.3b	48.1±1.6ab	17.6±0.5a	13.2±0.6a	7.9±0.2a
	XPro-Ca+S	$80.5 \pm 0.4b$	52.0±1.4c	$42.3\pm0.6c$	15.3±0.5b	11.9±2.6a	7.5±0.0a
	ZCK	$88.2 \pm 1.4a$	73.5±2.1a	61.3±0.9a	14.5±0.5a	10.6±0.8a	8.4±0.1b
	ZS	70.5±0.3c	58.4±1.4c	44.9±2.9b	8.9±0.5c	$6.1 \pm 0.4b$	$5.5\pm0.2c$
	ZPro-Ca	$76.9 \pm 3.2b$	$65.6 \pm 2.0 b$	57.3±3.5a	12.4±0.5b	10.7±0.3a	$8.8 \pm 0.3b$
	ZPro-Ca+S	74.2±0.7bc	$70.3 \pm 1.4 ab$	$56.0\pm3.5a$	$12.1 \pm 0.8b$	9.6±0.2a	9.7±0.1a
35	XCK	$86.2 \pm 2.6 ab$	70.0±3.9a	58.8±1.0a	19.8±1.0a	13.4±0.9a	9.9±0.5a
	XS	$80.3 \pm 1.6b$	64.4±0.8ab	51.3±0.2bc	15.6±0.4b	9.5±0.6b	7.5±0.3b
	XPro-Ca	$87.3 \pm 1.3a$	$63.8 \pm 2.4ab$	$52.8 \pm 0.4b$	20.0±1.1a	15.8±0.7a	10.0±0.9a
	XPro-Ca+S	82.0±2.0ab	55.8±4.9b	$49.2 \pm 1.4c$	18.9±0.3a	13.4±1.1a	$8.3\pm0.0ab$
	ZCK	90.0±0.3a	$84.0\pm2.0a$	75.8±2.0a	14.9±0.8a	$10.3 \pm 0.2b$	9.3±0.1a
	ZS	$80.2 \pm 0.6b$	70.3±6.6b	61.0±5.1b	12.3±0.5b	9.5±0.3b	$6.8 \pm 0.2b$
	ZPro-Ca	76.3±0.5bc	68.8±1.9b	61.9±2.8b	15.8±0.5a	13.3±0.9a	9.8±0.5a
	ZPro-Ca+S	$75.1\pm2.4c$	69.6±2.1b	63.8±0.2b	15.5±0.3a	10.6±0.2b	9.3±0.1a

Values described are the means \pm SE (n = 3). Different letters denote significant difference from

Duncan's LSD test (p < 0.05).

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Table 2(on next page)

Effects of Pro-Ca on root length of rice main stem and leaf area per stem of rice main stem, first tiller, and second tiller stems at the tillering stage under salt stress.

Values are the means±SE (n=3). Different letters denote significant difference from Duncan's LSD test (p<0.05).



- 1 Table 2
- 2 Effects of Pro-Ca on root length of rice main stem and leaf area per stem of rice main stem,
- 3 first tiller, and second tiller stems at the tillering stage under salt stress.

Time/ d	Treatment	Root length/ cm	Leaf area per stem/ cm ²			
Time/ a			0	I	II	
7	XCK	32.2±0.2a	5370.7±726.7ab	1817.1±158.7b	1297.3±249.1b	
	XS	16.7±2.1c	4305.7±149.7b	974.4±196.4c	863.1±169.3b	
	XPro-Ca	32.3±0.6a	6739.5±461.0a	2828.0±171.5a	995.5±141.1b	
	XPro-Ca+S	25.4±0.6b	6116.9±293.6a	2589.2±117.2a	1857.3±57.1a	
7	ZCK	28.0±0.4a	4316.7±701.1a	2139.5±320.7a	1812.1±29.7a	
	ZS	20.2±0.4c	2503.7±181.5b	573.0±47.9b	$628.0 \pm 114.9b$	
	ZPro-Ca	28.3±0.7a	4958.6±249.8a	2751.1±330.2a	1735.9±327.2a	
	ZPro-Ca+S	24.1±0.6b	4556.0±449.6a	2459.9±198.9a	1909.4±183.4a	
	XCK	34.0±0.7a	$8768.8 \pm 265.8ab$	4157.0±342.1a	2850.5±19.0a	
	XS	23.3±1.0d	5765.6±533.7c	2625.8±151.3b	$1610.0 \pm 138.4b$	
	XPro-Ca	30.4±0.1b	9411.7±502.4a	3800.7±95.0a	$2544.2 \pm 169.6a$	
14	XPro-Ca+S	26.5±0.9c	7524.2±54.7b	3416.4±357.4ab	2659.9±94.3a	
14	ZCK	31.6±0.5a	4542.4±699.9bc	3800.3±49.7b	2476.3±155.0b	
	ZS	24.0±0.5c	3191.6±310.9c	1884.3±108.6c	1335.7±146.5c	
	ZPro-Ca	$30.4 \pm 0.2a$	7599.9±557.3a	5933.6±537.9a	4297.4±297.2a	
	ZPro-Ca+S	$29.1 \pm 0.2b$	5725.1±198.5b	3424.3±163.0b	2854.3±261.0b	
	XCK	36.9±0.9a	13073.6±31.9b	9086.0±1632.7a	3487.7±242.7b	
	XS	$26.3\pm0.3c$	8518.4±824.1c	4431.7±473.8b	2343.2±156.9c	
	XPro-Ca	35.1±0.7a	16377.4±1268.1a	9106.6±655.3a	$4243.2\pm107.5ab$	
21	XPro-Ca+S	31.1±0.4b	13157.8±1026.7b	7390.5±261.9ab	4539.4±405.0a	
21	ZCK	31.3±0.3a	11510.1±169.6ab	$8480.1\pm214.0b$	4263.7±158.3b	
	ZS	25.9±0.3c	6699.3±469.0c	3349.3±104.3c	2419.2±134.3c	
	ZPro-Ca	31.9±0.4a	12499.1±591.7a	9743.8±430.3a	5790.4±431.2a	
	ZPro-Ca+S	28.1±0.5b	10685.9±115.6b	8323.6±267.9b	5174.6±300.1ab	

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10 Table 2 continued

	XCK	38.9±0.6a	19356.5±214.9a	10053.9±1078.9a	4877.8±232.1a
	XS	28.9±0.5c	11285.3±1893.9b	4555.5±186.5c	3383.9±246.4b
	XPro-Ca	35.5±0.2b	21134.8±1004.4a	11031.7±344.0a	5339.8±348.8a
28	XPro-Ca+S	35.7±0.6b	18674.2±442.4a	7893.0±501.8b	4676.2±44.4a
28	ZCK	34.3±1.0a	13441.2±333.7a	8997.6±286.3b	6685.3±605.7a
	ZS	27.6±0.6b	7290.5±219.1c	4316.1±337.7c	3482.6±238.6b
	ZPro-Ca	$34.9 \pm 0.4a$	12883.3±750.9a	9499.3±462.4b	$7186.9 \pm 142.7a$
	ZPro-Ca+S	34.8±0.3a	10919.9±694.5b	10803.5±466.6a	6794.2±293.9a
	XCK	41.9±0.7a	22136.7±1754.2a	15401.6±3314.5ab	7314.4±1013.4a
	XS	$30.0 \pm 0.4b$	14410.6±860.6b	6125.3±374.9c	4154.9±161.7a
	XPro-Ca	39.7±0.7a	22637.8±649.4a	18446.2±1148.1a	7497.9±1618.8a
35	XPro-Ca+S	41.0±1.0a	20386.9±589.8a	11051.6±981.7bc	5948.2±318.4a
33	ZCK	36.4±0.6a	17411.4±454.4a	11695.5±378.4b	9364.4±773.9a
	ZS	30.7±0.3b	10297.9±287.1d	8110.1±300.1c	4612.9±130.9b
	ZPro-Ca	$38.3 \pm 1.2a$	14462.8±419.4b	12506.7±509.8b	8107.0±497.7a
	ZPro-Ca+S	$37.8 \pm 0.5a$	12335.2±361.4c	14148.4±387.9a	7825.9±929.4a

Values are the means \pm SE (n = 3). Different letters denote significant difference from Duncan's

¹² LSD test (p < 0.05).



Table 3(on next page)

Effects of Pro-Ca on dry weight per stem of rice main stem, first tiller, and second tiller stems and root dry weight of rice main stem at the tillering stage under salt stress.

Values are the means±SE (n=3). Different letters denote significant difference from Duncan's LSD test (p<0.05).



- 1 Table 3
- 2 Effects of Pro-Ca on dry weight per stem of rice main stem, first tiller, and second tiller
- 3 stems and root dry weight of rice main stem at the tillering stage under salt stress.

Time/	T	D	D4 1		
d	Treatment	0	I	II	Root dry weight/ g
	XCK	0.470±0.017a	0.168±0.004a	0.103±0.005b	0.195±0.007a
	XS	$0.340\pm0.015b$	$0.062\pm0.011b$	$0.065\pm0.006c$	0.118±0.007d
	XPro-Ca	$0.478\pm0.045a$	$0.178\pm0.003a$	$0.133 \pm 0.005a$	$0.170\pm0.005b$
	XPro-Ca+S	0.466±0.011a	$0.156\pm0.015a$	$0.094\pm0.012b$	$0.146\pm0.004c$
7	ZCK	$0.426 \pm 0.053a$	$0.168 \pm 0.023b$	$0.129\pm0.004a$	$0.154\pm0.005b$
	ZS	$0.202\pm0.005b$	$0.058\pm0.004c$	$0.045 \pm 0.008b$	$0.098\pm0.002c$
	ZPro-Ca	0.392±0.011a	$0.212\pm0.009a$	$0.143\pm0.012a$	$0.189\pm0.007a$
	ZPro-Ca+S	$0.365 \pm 0.006a$	$0.159\pm0.008b$	$0.138\pm0.008a$	$0.139\pm0.013b$
	XCK	$0.626\pm0.020a$	$0.295\pm0.019b$	$0.198\pm0.002a$	$0.268 \pm 0.011ab$
	XS	$0.392 \pm 0.040b$	$0.188 \pm 0.004b$	$0.124\pm0.006c$	$0.140\pm0.000c$
	XPro-Ca	$0.573 \pm 0.028a$	$0.440\pm0.074a$	$0.206\pm0.014a$	0.318±0.053a
1.4	XPro-Ca+S	$0.532\pm0.016a$	$0.239\pm0.012b$	$0.168 \pm 0.002b$	$0.191\pm0.004bc$
14	ZCK	$0.464\pm0.017a$	$0.352\pm0.033a$	$0.191 \pm 0.013b$	$0.235 \pm 0.008b$
	ZS	$0.305 \pm 0.027b$	$0.148\pm0.012c$	$0.101\pm0.004c$	0.112±0.001c
	ZPro-Ca	$0.552\pm0.026a$	$0.415\pm0.032a$	$0.262\pm0.010a$	0.311±0.010a
	ZPro-Ca+S	$0.468 \pm 0.050a$	$0.245 \pm 0.007b$	$0.236 \pm 0.024ab$	$0.229 \pm 0.009b$
	XCK	1.145±0.066a	$0.382 \pm 0.003c$	$0.199 \pm 0.003b$	$0.405 \pm 0.020b$
	XS	$0.638 \pm 0.065b$	$0.345\pm0.029c$	$0.182 \pm 0.008b$	0.266±0.007d
	XPro-Ca	1.194±0.078a	$0.579\pm0.024a$	$0.250\pm0.020a$	0.266±0.007d
21	XPro-Ca+S	1.249±0.030a	$0.460 \pm 0.018b$	$0.259\pm0.016a$	0.266±0.007d
	ZCK	$0.988 \pm 0.049a$	$0.650\pm0.019a$	$0.291\pm0.012b$	$0.331 \pm 0.021c$
	ZS	$0.565 \pm 0.005c$	$0.256 \pm 0.014b$	$0.222\pm0.020c$	0.206±0.009d
	ZPro-Ca	1.008±0.062a	$0.681 \pm 0.047a$	$0.367 \pm 0.012a$	0.596±0.013a
	ZPro-Ca+S	$0.791 \pm 0.037b$	0.598±0.031a	$0.359\pm0.018a$	$0.493 \pm 0.035b$

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10 Table 3 continued

	XCK	2.160±0.134a	0.943±0.041b	0.344±0.009a	1.469±0.073a
	XS	$0.896 \pm 0.023c$	$0.617 \pm 0.033c$	$0.224 \pm 0.002b$	$0.428 \pm 0.046d$
	XPro-Ca	$1.828\pm0.070b$	1.373±0.117a	$0.326 \pm 0.030a$	$1.278 \pm 0.013b$
20	XPro-Ca+S	$1.571\pm0.045b$	$0.633 \pm 0.029c$	$0.346 \pm 0.022a$	$0.601 \pm 0.048c$
28	ZCK	1.387±0.121a	$0.767 \pm 0.051b$	$0.376 \pm 0.013b$	$0.954\pm0.012a$
	ZS	$0.654\pm0.019b$	$0.400\pm0.008c$	$0.278 \pm 0.011b$	0.284±0.011d
	ZPro-Ca	1.467±0.114a	$0.948\pm0.047a$	$0.689 \pm 0.030a$	$0.695 \pm 0.019b$
	ZPro-Ca+S	1.235±0.126a	$0.908 \pm 0.076 ab$	$0.824 \pm 0.077a$	$0.631 \pm 0.020c$
	XCK	2.817±0.131a	1.765±0.056a	$0.446 \pm 0.042ab$	$1.938 \pm 0.081ab$
	XS	$1.053\pm0.009c$	$0.725 \pm 0.011b$	$0.350\pm0.013b$	1.014±0.075c
	XPro-Ca	$2.802\pm0.072a$	1.547±0.258a	$0.603 \pm 0.086a$	2.153±0.219a
25	XPro-Ca+S	$2.502\pm0.099b$	$1.432\pm0.039a$	$0.449 \pm 0.036 ab$	$1.639 \pm 0.037b$
35	ZCK	2.243±0.104ab	1.265±0.139b	$0.544 \pm 0.076 b$	$1.446 \pm 0.059a$
	ZS	$1.575\pm0.054c$	$0.881 \pm 0.056c$	$0.429 \pm 0.012b$	$0.551\pm0.019c$
	ZPro-Ca	$2.321\pm0.144a$	$1.539\pm0.024a$	$1.341\pm0.052a$	$0.842 \pm 0.079b$
	ZPro-Ca+S	1.915±0.121bc	$1.421 \pm 0.018ab$	1.212±0.017a	$0.767 \pm 0.035b$

Values are the means \pm SE (n = 3). Different letters denote significant difference from Duncan's

¹² LSD test (p < 0.05).