Salt-tolerance screening in *Limonium sinuatum* varieties with different flower colors

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*Limonium sinuatum*, a member of Plumbaginaceae commonly known as sea lavender, is widely used as dried flower. Five *L. sinuatum* varieties with different flower colors (White, Blue, Pink, Yellow, and Purple) are found in saline regions and are widely cultivated in gardens. In the current study, we evaluated the salt tolerance of these varieties under 250 mmol/L NaCl (salt-tolerance threshold) treatment to identify the optimal variety suitable for planting in saline lands. After the measurement of the fresh weight (FW), dry weight (DW), contents of Na\(^+\), K\(^+\), Ca\(^{2+}\), Cl\(^-\), malondialdehyde (MDA), proline, soluble sugars, hydrogen peroxide (H\(_2\)O\(_2\)), relative water content, chlorophyll contents, net photosynthetic rate, and osmotic potential of whole plants, the salt-tolerance ability from strongest to weakest is identified as Pink, Yellow, Purple, White, and Blue. Photosynthetic rate was the most reliable and positive indicator of salt tolerance. The density of salt glands showed the greatest increase in Pink under NaCl treatment, indicating that Pink adapts to high-salt levels by enhancing salt gland formation. These results provide a theoretical basis for the large-scale planting of *L. sinuatum* in saline soils in the future.

*Limonium sinuatum* L., a native Mediterranean plant, is widely distributed in Northern Africa, western Asia, and Europe\(^1\). *L. sinuatum*, a member of the Plumbaginaceae family, is a typical recratohalophyte that can grow in saline\(^2\) or drought environments due to the presence of salt glands in the epidermis and the surrounding thick cuticle to reduce the water evaporation\(^3\). Different *L. sinuatum* varieties are usually identified based on flower color. The flower petals are white, while the calyxes can be different colored (e.g., white, blue, pink, yellow, and purple)\(^4\). The calyxes remain long after the petals have disappeared, making the flowers attractive for long periods of time and excellent for use as fresh cut flowers or in dried arrangements\(^5\). The flower stems are approximately 40–50 cm tall, each flower stem has 2–3 branches with 5–6 flowers clustered together, and the diameter of each flower is ~0.5 cm\(^6\). *L. sinuatum* is commonly referred to as static, sea lavender, sea notchleaf\(^7\), or wavyleaf sea lavender when used for gardening or floral arrangements\(^8\). Moreover, the entire plant is used as a traditional Chinese medicine for hemostasis\(^8\). In China, *L. sinuatum* is widely distributed along the coast of the Yellow Sea and the Bohai Sea\(^8\).

Soil salinization is a major environmental factor affecting plant growth and development\(^9\). Plant salt tolerance is a highly complex trait involving many factors, such as tissue and organ structure and physiological and biochemical reactions\(^10\). Salt stress alters the contents of soluble sugars, ions, and proline in plants\(^11\), and it affects the synthesis of plant soluble substances\(^12\), leading to changes in salt tolerance\(^13\).

Most crops are non-halophytes, and their growth is inhibited in saline soil\(^14\). By contrast, halophytes, including euhalophytes, recratohalophytes, and psudohalophytes\(^15\), can grow and complete their lifecycles in the presence of ≥ 200 mmol/L NaCl\(^16\). *L. sinuatum* is a recratohalophyte with salt glands for excreting excess Na\(^+\) out of the plants to avoid salt stress. To date, 67 species with salt glands have been reported, belonging to 13 families\(^17\). Among Plumbaginaceae family members, many studies have been carried out on *L. bicolor*, including salt secretion measurements\(^18\), analysis of salt gland differentiation\(^19\), and transcriptomic analysis during leaf development and in response to NaCl treatment\(^18\). However, although *L. sinuatum* also belongs to Limonium, few studies of the salt resistance in different varieties of this plant have been reported.

Generally, *L. sinuatum* has greater potential than *L. bicolor* for use as a horticultural crop in saline soils\(^20\), because the growth cycle of *L. sinuatum* is short (do not need vernalization for flowering) and the flower color is various, and it is more suitable for use as fresh cut flowers or in dried arrangements\(^21\). Given that *Limonium* species are considered to be pioneer plants for transforming saline soils\(^22\), it is important to explore the use of these plants to maximize the utilization of saline lands to increase the economic and ecological value of these environments. Here, we selected five common
garden varieties of *L. bicolor* (named based on flower color), including *White, Blue, Pink, Yellow,* and *Purple,* and determined their salt-tolerance thresholds. We evaluated the salt tolerance of these varieties by comparing fresh weight, dry weight, ion, proline, soluble sugars, and chlorophyll content, net photosynthetic rate, and osmotic potential to identify the varieties with the strongest salt tolerance. Based on these salt-tolerance indicators, we identified the most suitable variety for planting in saline soil.

**Materials and methods**

**Plant materials and growth conditions.** The seeds of five *Limonium sinuatum* varieties with different flower colors (*White, Blue, Pink, Yellow,* and *Purple*) were purchased from Lanxiang Horticulture Seeding Co., Ltd. (China). This study complies with local and national regulations. The author Baoshan Wang had formally identified *L. sinuatum* and the seeds harvesting process is in full compliance with relevant government guidelines. Unfortunately, we were unable to find a voucher specimen of *L. sinuatum* stored in any publicly available herbarium. The dried seeds were stored in a refrigerator at < 4 °C. The seeds were sterilized in 6% NaClO for 15 min, washed with sterile distilled water, and sown in nutrient soil (soil: vermiculite: perlite, 3: 1: 1). The plants were grown in a growth chamber at 28 °C/23 °C (day/night) under 600 μmol/m²/s full-spectrum light (15 h photoperiod) and 60% relative humidity. In order to make accurate comparisons among the five varieties, all plants were cultured for six months under the above conditions for flowering (Fig. 1). Flower color can be used to distinguish among different varieties of *L. sinuatum,* including *White, Blue, Pink,* *Yellow,* and *Purple.*

**Measurement of salt-tolerance threshold.** To determine the salt-tolerance threshold, two-week-old seedlings with two expanded leaves were treated with different concentrations of NaCl (0, 100, 200, 300 and 400 mmol/L). After two weeks of treatment, the aerial parts of the plants were collected and used to measure fresh weight (FW) and dry weight (DW) according to Huang et al. They replicated were performed per variety, and the means among different varieties under each NaCl treatment were used to calculate the salt-tolerance threshold. In detail, a fit regression curve was established with different NaCl concentrations vs. FW or DW. The NaCl concentration at which the plants showed 50% FW or DW compared to the non-NaCl treatment group was considered to be the salt-tolerance threshold.

After the determination of salt-tolerance threshold, two-week-old seedlings of the five varieties with two expanded leaves were then re-treated with 0 mmol/L and 250 mmol/L NaCl (considered to be the salt-tolerance threshold in the following experiments) for two weeks and used to measure the physiological indicators.

**Determination of physiological indicators.** Determination of FW, DW and relative water content of leaf. After cleaning the leaves with 10 mM calcium chloride solution followed by deionized water, FW of the leaves was measured immediately and DW was obtained following incubation at 105 °C for 15 min and drying to constant weight at 70 °C for 2 days. Five replicates were performed per variety and treatment. The reduction rate was calculated as (FW under control condition—FW under saline condition)/FW under control condition × 100%. The same method was processed in calculating the reduction rate of DW. The relative water content is calculated as (FW—DW)/FW × 100%.

Determination of sodium ion (Na⁺), potassium ion (K⁺), calcium ion (Ca²⁺), and chloride ion (Cl⁻) contents. The ion contents in the samples were measured according to Higinbotham. In brief, leaf tissue (0.5 g FW) was collected from plants under both 0 and 250 mmol/L NaCl treatment for all five varieties. The tissues wereashed, dissolved in HNO₃, and the contents of Na⁺, K⁺, and Ca²⁺ measured using a Flame photometer (FP6440, Yuanxi, Shanghai, China). Cl⁻ content was measured by ion chromatography according to Wang. Briefly, after boiling for 30 min and filtering through a 0.22 μm filter membrane, the ion solution was injected into an ion chromatograph (ICS-90A, ThermoFisher, Massachusetts, USA) to measure Cl⁻ contents. The ion concentration is shown as mmol/g FW. Five replicates were performed for each variety and treatment. Given that Na⁺ content increased under NaCl treatment, the increase rate was calculated as (Na⁺ content under saline condition—Na⁺ content of control)/Na⁺ content of control × 100%. The same calculation method was applied in the increase rate of Cl⁻.

Determination of proline content, osmotic potential, malondialdehyde (MDA), hydrogen peroxide (H₂O₂), soluble sugars, and chlorophyll content, and photosynthetic rate. Proline content was determined in accordance with Demiral. The plant tissue was ground, and ground tissue (0.5 g FW) was added to 10 mL of 5% acetic acid and 40 mL of distilled water. After filtering, the filtrate (8 mL) was mixed with 0.8 g zeolite with shaking for 5 min and centrifuged for 10 min (1500 g). The supernatant (3 mL) was combined with glacial acetic acid (3 mL) and ninhydrin reagent (3 mL) and boiled for 1 h. Benzene (3 mL) was used for static layering, and the upper colored liquid was collected and used to measure optical density at 515 nm. The proline content was calculated from a standard curve based on the optical density. Five replicates were performed for each variety and treatment.

The osmotic potential was measured as described by Tomlinson. Fresh leaf tissue (0.5 g) was cut into small pieces, frozen in liquid nitrogen, and placed into a syringe to squeeze out and collect the cell sap. A freezing point osmometer (SMC 30C-1, Tianhe, Tianjin, China) was used to measure the osmotic potential of the plant cell sap. The formula used to calculate osmotic potential is $\Delta iCRT (R = 0.0083143 L Mpa\, Mol^{-1}\, K^{-1}, T = 298 K)$. Five replicates were performed for each variety and treatment.

The MDA content was determined as reported in Hong. Leaf tissue (0.5 g) was collected and homogenized in 5 mL 0.1% TCA. The homogenate was transferred to the test tube, combined with 5 mL 0.5% thiobarbituric acid solution, and boiled for 10 min. The sample was centrifuged at 1500 g for 15 min, and the optical density of the supernatant was measured at 532 nm and 600 nm. MDA content (mmol/g FW) = $\Delta A_{532} - \Delta A_{600}$, N is the total volume of the supernatant; 155 is the absorption coefficient of
1 mmol reaction product at 532 nm; W is the fresh weight of the plant material (g). Five replicates were performed for each variety and treatment.

The content of H$_2$O$_2$ was determined as described by Vergara$^{32}$. In brief, fresh leaves (0.3 g) was grinded in 5 mL precooled acetone before centrifuged at 500 g for 8 min. Afterward the supernatant (1 mL) was mixed with...
ammonia (0.2 mL) and 20% TiCl₄ (0.1 mL) for 2 min, the precipitate was washed with acetone for 3–5 times and dissolved in 2 M H₂SO₄ (5 mL) after centrifuged at 600 g for 7 min. Then the content of H₂O₂ was measured at 415 nm and calculated as H₂O₂ content (μmol/g FW) = CV₁/FW₁, C is the concentration of H₂O₂ in the sample checked on the standard curve (μmol), V₁ was the total volume of sample extract (mL), V₂ was the volume of sample extract (mL), and FW was the fresh weight of plant tissue (g). Five replicates were performed for each variety and treatment.

Soluble sugars were measured following the protocol of Prado33. Fresh leaf tissue (0.3 g) was dissolved in 10 mL of double distilled H₂O (ddH₂O) in a boiling water bath for 50 min, filtered, and brought to a volume of 25 mL. Afterward 0.5 mL of extract solution was combined with 1.5 mL distilled water, 0.5 mL ethyl anthrone acetate, and 5 mL concentrated sulfuric acid, shaken thoroughly, boiled in water bath for 1 min, and cooled. The optical density of the solution was measured at 630 nm. The soluble sugars content was calculated from a standard curve. Five replicates were performed for each variety and treatment.

Chlorophyll levels were determined referring to Maxwell34. Leaf tissue (0.3 g) was combined with 5 mL dimethyl sulfoxide in 5 mL 80% acetone and incubated in a 65°C water bath at 24 h (protected from the light) to fully decolorize. Afterward bring to 25 mL after filtration and the solution was used to measure the optical density at 663 nm, 645 nm, and 470 nm. Chlorophyll content (mg g⁻¹ or mg dm⁻²) = CV/1000A, C is chlorophyll concentration (mg L⁻¹ or mg dm⁻²); V is the total volume of extract solution (mL); A is fresh weight of the sample (g) or sampling area (dm²). The pigment concentration (mg/L) was calculated as Cₐ = 12.7A₆₆₃ − 2.69A₆₄₅; Cₙ = 22.9A₆₄₅ − 4.68A₆₆₃; Cₜₐₜ = 20.0A₆₆₃ + 8.02A₆₄₅; Cₜₐₜ = (1000A₆₇₀ − 3.27Cₐ − 104Cₙ)/229; Cₐ, Cₙ are the concentrations of chlorophyll a and b, Cₜₐₜ is the concentration of total chlorophyll; Cₜₐₜ is the total concentration of carotenoids. Five replicates were performed for each variety and treatment.

The photosynthetic rate was measured on the basis of Wang35. In this experiment, a photosynthetic instrument (LI-6400XT, LI-COR, Nebraska, USA) was used to measure the photosynthetic parameters of leaves. The photosynthetic effective quantum density, Uₚₑₙᵦ (μmol m⁻² s⁻¹), µ is 4.55³⁶, was measured at a temperature of 23°C, and the leaf area of each cultivar was 1 cm². Five replicates were performed for each variety and treatment.

Determination of salt gland density in different varieties. The density of salt glands was measured according to Yuan37. The leaves were fixed in a mixture of ethanol and acetic acid (3:1; v/v), rinsed with 70% ethanol to decolorize, and cleared in Hoyer’s solution. Afterward cleaned leaves were fixed on a glass slide for DIC microscopy (ECLIPSE 80i, Nikon, Tokyo, Japan). The salt gland density was calculated according to Ding38 and expressed as number per mm². Five replicates were performed for each variety and treatment.

Data analysis
Statistical analysis and correlation were performed using SPSS 13.0 software (SPSS Software Inc., USA). The results were subjected to a one-way analysis of variance (ANOVA), and Duncan’s test was used to determine significant differences between the means (P = 0.05). In the figures, the error bars represent the means ± standard deviations (n = 5) and different letters indicate significant differences at P = 0.05. Correlation is processed at P = 0.05 and 0.01 using Pearson correlation analysis. The figures were generated using SigmaPlot 12.5 (Systat Software, Chicago, IL, USA).

Results
Identification of the salt-tolerance threshold. Plant biomass is an important measure of salt tolerance. Figure 2 shows the biomass of the aerial parts of plants under a gradient of different NaCl concentrations (0, 100, 200, 300, and 400 mmol/L) after 2 weeks of treatment. FW and DW were measured in the five varieties of L. sinuatum seedlings and constructed a regression curve based on the means of five data under different treatments as independent variables. Most studies use the salt concentration at which plant growth or biomass decreases by 50% of the control value as the salt-tolerance threshold39. Here, when the FW and DW of the
upper parts of the seedlings were reduced by 50% of the non-NaCl treatment value, different varieties showed different salt-tolerance thresholds. The highest threshold was obtained for Pink (250 mmol/L) (Supplementary Fig. 1), suggesting that Pink is the most salt-tolerant variety. To identify the optimal salt concentration for further experiments, we calculated the average salt-tolerance threshold, i.e., 228 mmol/L for FW and 233 mmol/L for DW (Fig. 2). Therefore, a salt-tolerance threshold of 250 mmol/L was utilized in subsequent experiments.

Pink shows the best growth under 250 mmol/L NaCl treatment. All varieties showed inhibited growth under 250 mmol/L NaCl treatment (Fig. 3), but the changes in FW and DW showed no significant trends among varieties (Supplementary Fig. 2). In order to make effective comparison among different varieties, FW and DW reduction rate are calculated to compare the changes between control and saline condition (Fig. 4). Pink showed the least FW reduction, followed by Yellow, Purple, White and Blue, while White has the least DW reduction, afterward Pink, Yellow, Purple and Blue. Based on the reduction rate of FW and DW, Pink is considered the most salt tolerance variety, followed by Yellow, Purple, White, and Blue. Biomass can be used as a measure of plant growth, and various physiological processes could be responsible for the ability of Pink to maintain growth in the presence of salt. Therefore, we measured the physiological indicators of the different varieties under NaCl treatment in order to reveal the underlying salt-tolerance mechanisms.

Effect of NaCl treatment on different physiological indicators in five varieties. Comparisons of the Na⁺, K⁺, Ca²⁺, and Cl⁻ contents; MDA, soluble sugars, proline contents, H₂O₂ content and relative water content of leaf; chlorophyll contents; and osmotic potential and photosynthetic rate are shown in Figs. 5, 6, 7 and 8, respectively. Each variety showed significant changes under NaCl treatment.

Figure 5 shows a comparison of the relative Na⁺, K⁺, Ca²⁺ and Cl⁻ contents among varieties under 250 mmol/L NaCl treatment. Na⁺ and Cl⁻ content under NaCl treatment increased compared with the control in all varieties (Fig. 5a,c), while K⁺ and Ca²⁺ showed various trends in different varieties (Fig. 5e,f). Given that Na⁺ and Cl⁻ are considered the stress ion to protoplast and different varieties have various basal level under control, the increase rate of Na⁺ (Fig. 5b) and Cl⁻ (Fig. 5d) under NaCl treatment are calculated in each variety in order to make intuitive comparison among different varieties. Pink shows the least Na⁺ increase (1.87%) under saline condition, followed by Yellow, Blue, Purple and White. In the aspect of Cl⁻ increase, Purple (96.04%) has the least, afterward Pink, Yellow, Purple and Blue. Together with the increase rate of Na⁺ and Cl⁻, these results indicate that Pink accumulates less Na⁺ and Cl⁻ than the other varieties under high-salt conditions, which should lead to less injury than the other varieties.

Figure 6 shows a comparison of the relative MDA, soluble sugars, proline contents, H₂O₂ and relative water content of leaf among varieties. Though the MDA increase rate of Pink under saline condition (Fig. 6b) shows the most, the absolute value of MDA (Fig. 6a) is the least accumulation in Pink, indicating that Pink suffered the
least amount of damage under salt treatment, as MDA can be used as a measure of the degree of damage under NaCl treatment. High accumulation of MDA can be detected in White and Blue (Fig. 6a), which may explain the serious damage level.

To cope with the damage caused by NaCl treatment, cells usually accumulate organic osmotic regulating substance such as soluble sugars (Fig. 6c) and proline (Fig. 6e). High accumulation of soluble sugars is shown in Pink and Purple, and the increase rate (Fig. 6d) under saline treatment indicates the comparison between varieties. Highest increase rate is detected in Purple, followed by Pink and Yellow. Besides, high proline accumulation is shown in Blue, Yellow and Pink in descending order of actual value (Fig. 6e), while the increase rate has the opposite trend with the most in White, Blue and Pink (Fig. 6f). Proline reduces the osmotic potential in the cell, allowing it to resist external osmotic stress, thereby improving plant survival in adverse environments.

Proline content depends on the catabolism of sugar. Combined the absolute value and the increase rate, Proline is considered to accumulate a large amounts of soluble sugars and proline to improve the osmotic adjustment ability under salt treatment. The large accumulation of osmoregulation substances can effectively reduce the osmotic potential under NaCl treatment. Figure 8 (a) shows that the osmotic potential decline markedly in all varieties, and Yellow and Pink have the most reduction rate (Fig. 8b).

Moreover, H$_2$O$_2$, as a kind of superoxide, can cause oxidative stress to plants under various stresses. In Fig. 6g, under salt stress, the lowest H$_2$O$_2$ generation is detected in Pink, while higher in White and Purple, which indicates that Pink suffers the least oxidative stress under salt treatment and is more suitable for saline environment. In addition, relative water content is also measured (Fig. 6h) and no significant difference is detected among different varieties, indicating that all varieties of Limonium sinuatum can keep normal moisture condition to cope with physiological drought of NaCl.

Plants always produce large amounts of pigments under saline environment to maintain normal photosynthetic efficiency. In addition, a positive correlation was detected between chlorophyll content and net photosynthetic rate. Figure 7 shows a comparison of chlorophyll content among varieties. In order to show the changes in pigment content in more detail, the changes in total chlorophyll, chlorophyll $a$, chlorophyll $b$, and carotenoid contents are shown. The pigment contents of the Pink and Purple varieties were high, which help improve the photosynthetic rate. Under NaCl treatment, net photosynthetic rate is obviously inhibited in all varieties (Fig. 8c). Pink shows the highest under saline condition, and the reduction rate under salt treatment is also the lowest in Pink (Fig. 8d). These results further explain why Pink has the highest biomass under NaCl treatment, which may be due to high accumulation in osmoregulation substance and high photosynthetic efficiency.

**Effects of NaCl on salt gland density of five varieties of Limonium sinuatum.** Salt glands are structures for salt secretion that are specifically produced by recratohalophytes. We therefore performed statistical analysis of the salt gland densities of expanded leaves of the five Limonium sinuatum varieties under NaCl treatment.

Figure 9a shows the changes in salt gland density in the leaves of the five varieties of Limonium sinuatum under salt stress (images shown in Supplementary Fig. 3). Salt gland density increased in all varieties under NaCl treatment compared to the control. The density of salt glands in Pink increased by 225.86% (Fig. 9b), while in Purple only 7%. The increased salt gland can help the plants to excrete more Na$^+$ outside in order to further decrease the Na$^+$ accumulation in vivo.

Finally, given that Pink showed the greatest tolerance to NaCl treatment, in order to verify the optimal variety suitable for growing in field, we examined flowering in the five varieties grown in Yellow River Delta (salt content: 0.2%). After six months of growth, only Pink and Yellow plants flowered consistently, whereas the three other varieties rarely flowered and only showed vegetative growth (Fig. 10). These results suggest that Pink and Yellow...
are the optimal varieties for the development of saline horticulture and further planting in saline soil, which is consistence with the formal results in laboratory conditions.

Discussion

*L. sinuatum* is a pioneer plant that could be used for the improvement of saline lands due to the high salt resistance and various colors⁴⁸. Therefore, it is important to identify the best salt-tolerant varieties for cultivation in these areas. In the current study, *Pink* showed the highest biomass and the strongest salt resistance among the five varieties examined. Our analysis of physiological indicators including *Na*⁺, *K*⁺, *Ca*²⁺, and *Cl*⁻ contents; MDA, soluble sugars, proline contents, *H₂O₂* content, relative water content and chlorophyll contents, osmotic

**Figure 5.** Effect of NaCl treatment on *Na*⁺, *K*⁺, *Ca*²⁺, and *Cl*⁻ contents. Contents of *Na*⁺ (a), *Cl*⁻ (c), *K*⁺ (e) and *Ca*²⁺ (f) under control and NaCl treatment in different varieties. The data are means ± SD of five replicates. Different letters indicate significant differences between two groups at *P* = 0.05 using Duncan’s test with SPSS. (b, d) The increase rate of *Na*⁺ and *Cl*⁻ under NaCl treatment in five varieties, which was calculated as (ion contents under NaCl treatment—ion contents under control)/ion contents under control × 100%.
Figure 6. Effect of NaCl stress on MDA, soluble sugars, proline contents, H$_2$O$_2$ and relative water content in the leaves of five varieties of *Limonium sinuatum*. Contents of MDA (a), soluble sugars (c), proline (e), H$_2$O$_2$ (g) and relative water content (h) of leaves under control and NaCl treatment in different varieties. The data are means ± SD of five replicates. Different letters indicate significant differences between two groups at $P=0.05$ using Duncan’s test with SPSS. The increase rate of MDA (b), soluble sugars (d) and proline (f) are calculated using (value under NaCl treatment—value under control)/value under control × 100%.
potential, photosynthetic rate, and salt gland density under 250 mmol/L NaCl (salt-tolerance threshold) treatment explained why *Pink* has the greatest salt tolerance. Moreover, analysis of plants grown in the field (Fig. 10) confirmed the superior salt resistance of *Pink*. *Pink* is recommended as the optimal varieties for extensive planting and greening in saline soil, followed by *Yellow*.

Plant biomass is an important indicator of salt tolerance: the greater the increase in biomass under salt stress, the higher the salt tolerance. As shown in Fig. 4, *Pink* showed the minimal reduction under 250 mmol/L NaCl treatment, and photosynthetic efficiency of *Pink* was also the highest among five varieties (Fig. 8), indicating that *Pink* suffered the least damage under saline condition. The three basic components of salt stress are usually considered as ionic toxicity, osmotic stress and oxidative stress.

How *Pink* cope with high ionic toxicity? On the one hand, under salt stress, Na⁺ content increases, which affects the absorption of K⁺ and Ca²⁺. K⁺ plays an important role in the osmotic regulation of cells. Ca²⁺ regulates the ionic balance and reduces the absorption of Na⁺. In salt-tolerant plants, K⁺ efflux is significantly inhibited under salt stress to maintain high intracellular K⁺/Na⁺ levels, thereby reducing the damage from salt stress. After salt treatment, the *Pink* variety had the relatively low Na⁺ and Cl⁻ contents among the five varieties, whereas K⁺ and Ca²⁺ showed the opposite trend. Therefore, *Pink* regulates ionic balance under salt stress, maintaining high K⁺/Na⁺ levels, thus showing strong salt tolerance. On the other hand, salt gland is the typical and specific epidermal structure of recetohalophytes, which can excrete the excessive Na⁺ out of the plants to avoid damage. The most salt gland was induced in *Pink* under salt treatment (Fig. 9), so it is speculated that Na⁺ can be effectively transferred out of the cell to further avoid ionic toxicity.

NaCl can induce the physiological drought due to the osmotic stress. Compatible media was always generated to deal with the osmotic stress, such as proline and soluble sugars. Proline is an important compatible solute in plant cells that protects enzymes from inactivation by NaCl and reduces the osmotic potential in the cells, thereby helping plants resist external osmotic stress and tolerate adverse environments. Sugar content under stress condition is intricately associated with carbohydrate content of plant. Though not always the highest...
Figure 8. Effect of NaCl stress on the osmotic potential and net photosynthetic rate in the leaves of five varieties of *Limonium sinuatum*. (a, c) Osmotic potential and net photosynthetic rate under control and NaCl treatment. The data are means ± SD of five replicates. Different letters indicate significant differences between two groups at *P* = 0.05 using Duncan’s with SPSS. (b, d) The reduction rate of osmotic potential and net photosynthetic rate under NaCl treatment. The reduction rate was calculated as (the value under control—the value under NaCl treatment)/the value under control × 100%.

Figure 9. Effect of NaCl stress on salt gland density in the leaves of five varieties of *Limonium sinuatum*. (a) The data are means ± SD of five replicates. Different letters indicate significant differences between two groups at *P* = 0.05 using Duncan’s with SPSS. (b) The increase rate of salt gland density was calculated using (value under NaCl treatment—value under control)/value under control × 100%.
Figure 10. Growth of five varieties of *Limonium sinuatum* in saline soil.

Table 1. Correlation analysis between fresh weight (FW) and other 14 indicators including dry weight (DW), Na⁺ (Na), K⁺ (K), Ca²⁺ (Ca), Cl⁻ (Cl), MDA, soluble sugars (Sugar), proline (Proline), hydrogen peroxide (H₂O₂), relative water content (RWC) and chlorophyll contents (Chl), osmotic potential (Os), photosynthetic rate (Pr), and salt gland density (SG) using Pearson correlation analysis. **Correlation is significant at the 0.01 level (2-tailed). *Correlation is significant at the 0.05 level (2-tailed).
accumulation in proline and soluble sugars, Pink keep the relatively high level of osmotic adjustment substance to reduce the osmotic potential, which allowed the plant to re-absorb water from the NaCl solution to maintain normal growth.

Oxidative stress is inevitable induced by salt stress due to the generation of superoxide, with the typical representative H₂O₂. Pink suffered the smallest oxidative stress under salt treatment and is more suitable for saline environment. MDA content can reflect the degree of membrane damage and the effects of stress on plants. The least accumulation of MDA also insist the opinion that Pink suffered less oxidative stress.

Given that 16 physiological indicators were measured under salt treatment in addition to DW and FW, which one is more closely related to biomass? Correlation analysis was performed between FW and the other 14 indicators. As shown in Table 1, photosynthetic rate showed the strongest positive correlation with salt resistance in L. sinuatum.

Together, the results of our experiments show that the Pink variety has the strongest salt tolerance. Based on these results, the level of salt tolerance of the five varieties of L. sinuatum from high to low is Pink, Yellow, Purple, White, and Blue. Further analysis of the performance of these varieties in saline soil also validate the results of these preliminary experiments. Nonetheless, our findings provide a theoretical basis for the cultivation of L. sinuatum in saline-alkali areas, which could be widely planted to facilitate the greening and transformation of saline soils.

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F.Y. designed the research; X.X. and Y.Z. performed the research; X.X. and P.M. analyzed the data; X.X. wrote the paper; F.Y. and B.W. revised the paper.

Competing interests
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