Screening and evaluation of reliable traits of upland cotton (Gossypium hirsutum L.) genotypes for salt tolerance at the seedling growth stage

SIKDER Ripon Kumar, WANG Xiangru, JIN Dingsha, ZHANG Hengheng, GUI Huiping, DONG Qiang, PANG Nianchang, ZHANG Xiling* and SONG Meizhen*


correspondence: hainan1571@163.com; songmeizhenaas@163.com
State Key Laboratory of Cotton Biology, Institute of Cotton Research of Chinese Academy of Agricultural Sciences, Anyang 455000, Henan, China

Abstract

Background: Salt stress significantly inhibits the growth, development, and productivity of cotton because of osmotic, ionic, and oxidative stresses. Therefore, the screening and development of salt tolerant cotton cultivars is a key issue towards sustainable agriculture. This study subjected 11 upland cotton genotypes at the seedling growth stage to five different salt concentrations and evaluated their salt tolerance and reliable traits.

Results: Several morpho-physiological traits were measured after 10 days of salinity treatment and the salt tolerance performance varied significantly among the tested cotton genotypes. The optimal NaCl concentration for the evaluation of salt tolerance was 200 mmol·L\(^{-1}\). Membership function value and salt tolerance index were used to identify the most consistent salt tolerance traits. Leaf relative water content and photosynthesis were identified as reliable indicators for salt tolerance at the seedling stage. All considered traits related to salt tolerance indices were significantly and positively correlated with each other except for malondialdehyde. Cluster heat map analysis based on the morpho-physiological salt tolerance-indices clearly discriminated the 11 cotton genotypes into three different salt tolerance clusters. Cluster I represented the salt-tolerant genotypes (Z9807, Z0228, and Z7526) whereas clusters II (Z0710, Z7514, Z1910, and Z7516) and III (Z0102, Z7780, Z9648, and Z9612) represented moderately salt-tolerant and salt-sensitive genotypes, respectively.

Conclusions: A hydroponic screening system was established. Leaf relative water content and photosynthesis were identified as two reliable traits that adequately represented the salt tolerance of cotton genotypes at the seedling growth stage. Furthermore, three salt-tolerant genotypes were identified, which might be used as genetic resources for the salt-tolerance breeding of cotton.

Keywords: Cotton genotypes, Salt stress, Screening, Membership function value, Cluster analysis

Background

Salinity is the second-most prevalent abiotic stress after drought, which not only limits plant growth but also progressively decreases the optimal yield of crops worldwide (Gao et al. 2016). Globally, more than 800 million hectares are invariably affected by salinity (Munns and Tester 2008). The salinity-affected zone increases as a result of climate change, sea level rise, and disproportionate surface irrigation in combination with inappropriate drainage systems (Kumar et al. 2013). It has been projected that if the current increase of salinity continues, 50% of the existing cultivated land will be affected by salinity stress by 2050 (Wang et al. 2003). In China, about 36 million hectares (4.88% of the available land) are affected by salinity stress (Li et al. 2014).
Currently, the global cultivable land is decreasing at an alarming rate and competition between cereal and fiber crops is progressively increasing. As a result, the cotton-growing area is increasingly relocated to saline and alkaline soils (Peng et al. 2018).

In general, salt-resistance is considered multifarious phenomena and plants have developed several mechanisms to counteract this form of stress at the cellular, subcellular and organ levels. The primary effect of salinity stress is osmotic stress, which starts by increasing Na\(^+\) and Cl\(^-\) concentrations in the root zone. Consequently, plants suffer ionic toxicity after the accumulation of salt in mature leaves as a result of transpiration (Munns and Tester 2008; Abdelraheem et al. 2019). Ion toxicity not only affects cellular metabolic processes but also impedes the photosynthetic unit of plants and induces oxidative damage. In case of prolonged stress, ion toxicity causes an initial biomass loss that may lead to programmed cell death (Sade et al. 2017). Moreover, the over-accumulation of toxic ions in a saline environment may enhance the disorder of K\(^+\)/Na\(^+\) ion homeostasis, accelerate the generation of reactive oxygen species (ROS), and thus, accelerate the proliferation of lipid peroxidation (Gill and Tuteja 2010; Sehar et al. 2019). Plants naturally cope with this condition by enhancing osmotic adjustment, regulating stomatal aperture, balancing ionic disequilibrium, increasing antioxidant activities, and consuming. Moreover, the salt tolerance of cotton represents a multifaceted quantitative trait, and thus, any single index may not be sufficient to reflect the tolerance level. Membership function value (MFV) analysis was applied to evaluate the salt tolerance of Brasica napus (Wu et al. 2019) and a number of scholars have also suggested the regression-based analysis for salt-tolerance evaluation of different crop species, e.g., wheat (Chen et al. 2012) and sorghum (Ding et al. 2018). Therefore, to shorten the selection period, identify the procedure, and improve the reliability of evaluation, 11 cotton materials with differing biological characteristics (selected from preliminary work) were selected and studied to establish a salt-tolerance evaluation system under the condition of hydroponics. This study is (1) to determine the optimum salt concentration of selected cotton genotypes; (2) to identify reliable morpho-physiological traits for salt tolerance at the early seedling growth stage; and (3) to explore the genetic potentiality of cotton genotypes and to cluster these into different salt-tolerance groups.

**Methods**

**Plant materials, growth environments, and treatments**

The seeds of Gossypium hirsutum were collected by Professor Wuwei Ye’s team of Institute of Cotton Research, Chinese Academy of Agricultural Sciences, Anyang, Henan, China. Genotypes were selected based on a previous selection experiment under salt pond conditions (Table 1). It is generally assumed that emergence and young seedling stages are very salt-sensitive stages in cotton (Peng et al. 2014), so this study investigated the traits at the seedling stage. Healthy and uniform seeds were selected and disinfected by 10% sodium hypochlorite solution for 5 min and carefully washed three times with deionized water. The seeds were sown at about 3 cm depth in sterilized wet sand 7 in plastic pots (10 cm × 10 cm × 10 cm). After 7 days of germination, vigorous and identical seedlings were enclosed in foam bands at the root-shoot junction and were transplanted...
to 7 L plastic boxes (12 plants per pot) filled with an aerated, modified half-strength Hoagland nutrient solution (2.0 mmol·L\(^{-1}\) Ca (NO\(_3\))\(_2\), 1.0 mmol·L\(^{-1}\) KCl, 0.25 mmol·L\(^{-1}\) KH\(_2\)PO\(_4\), 1.0 mmol·L\(^{-1}\) MgSO\(_4\), 0.05 mmol·L\(^{-1}\) EDTA·Fe·Na, 23.1 μmol·L\(^{-1}\) H\(_3\)BO\(_3\), 4.55 μmol·L\(^{-1}\) MnCl\(_2\)·4H\(_2\)O, 0.4 μmol·L\(^{-1}\) ZnSO\(_4\)·7H\(_2\)O, 0.15 μmol·L\(^{-1}\) CuSO\(_4\)·5H\(_2\)O, 0.5 μmol·L\(^{-1}\) (NH\(_4\))\(_6\)Mo\(_7\)O\(_24\)·4H\(_2\)O) in triplicate for each treatment. Seedlings were grown in a growth chamber under 16 h light and 8 h dark cycle, at a temperature of 28–30 °C, and (60 ± 5)% relative humidity. The nutrient solution was replaced once per week throughout the experimental period to maintain the appropriate composition. The pH of the growth medium was maintained between 5.8 and 6.2. Evaporated water was refilled with de-ionized (dH\(_2\)O) water every day to retain the target volume.

Seedlings were grown under non-saline conditions until they had three true leaves. Salinity treatment was initiated by providing various levels of NaCl (100, 150, 200, and 250 mmol·L\(^{-1}\)) along with controls (only nutrient solution). To avoid early lethal damage, NaCl was dissolved in the nutrient solution and the salt concentration was increased stepwise (25% of the maximum concentration on every alternative day) over a 7-day period. The salt stress treatment under hydroponic cultivation continued for 10 days, and then, plants were harvested and sampled (Fig. 1). The leaf samples were instantly frozen in liquid nitrogen at −80 °C for biochemical analysis. Different growth parameters, as well as ionic and physiological indices were recorded accordingly.

**Measurement of morpho-physiological parameters**

**Growth traits**

Growth was evaluated in terms of shoot length (SL), leaf numbers (LN), leaf area (LA), and total biomass (TB) of cotton seedlings. Ten days after salt treatment, seedlings from

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**Table 1** Plant materials information

| Names   | Background                                      | Salt response in the sand pond |
|---------|-------------------------------------------------|-------------------------------|
| Z9807   | This line derived from CCRI 7                   | Tolerant                      |
| Z0710   | This line derived from YU 2067                 | Tolerant                      |
| Z1910   | A hybrid progeny crossing between Ji851 and GKZ19| Tolerant                      |
| Z0228   | A hybrid offspring breeding between Ji Mian228 and 9807 | Tolerant                      |
| Z7514   | A hybrid offspring crossing between Ji Mian616 and Ji Mian228 | Tolerant                      |
| Z7516   | The line obtained from crossing between Yu Zao 97-1335 and Yu 2067 | Tolerant                      |
| Z7526   | Derived from Zhong 75-y-17                      | Tolerant                      |
| Z0102   | Crossing between Zhong 9807 and Zhong 9612 (F2 generation breeding). Recombinant Inbred Lines formed by multi-generation selfing | Sensitive                      |
| Z9612   | Selected from CCRI 12                           | Sensitive                      |
| Z7780   | The progeny of GK50                             | Sensitive                      |
| Z9648   | Breeding between GK50 and SGK36                | Sensitive                      |

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![Fig. 1 Schematic representation of the experimental method](image)
control and salt-treated genotypes were harvested. The morphological traits of five seedlings per replicate were measured. Shoot length was determined by computing the length from the cotyledon to the apex of the emerging leaf. The number of leaves was counted manually. For dry mass estimation, the fresh plant materials were cleaned with deionized water and divided into two parts (shoots and roots). Shoot and root parts were oven-dried at 105 °C for 30 min followed by 80 °C for 24 h until they maintained a constant weight. Shoot length was expressed in cm whereas g shoot⁻¹ was used to measure the biomass. A handheld leaf-area meter was used to calculate the leaf area (LI-3100, LI-COR, Inc., Lincoln, NE, USA).

**Determination of leaf relative water content**
The leaf relative water content (LRWC) was determined according to Barrs and Weatherley (1962). After collection, fresh leaves were immediately weighted as fresh weight (FW), then placed in a paper bag, and immersed in distilled water for 24 h under ambient light condition. The turgid weight (TW) was measured by removing all excess water from the leaves with the help of blotting paper. The dry weight (DW) was recorded after 48 h of oven drying at 80 °C. Finally, the LRWC was calculated based on the following equation:

\[
\text{LRWC} \% = \left( \frac{\text{FW} - \text{DW}}{\text{TW} - \text{DW}} \right) \times 100
\]

**Gas exchange parameters**
After 10 days of salt stress, gas exchange parameters, such as photosynthetic rate (A) and stomatal conductance (gsw), were recorded in each plant by using a handheld photosynthesis system (LI-6800, LI-COR, Inc., Lincoln, NE, USA). All gas exchange parameters were measured with 6 cm² of total leaf area from the top third fully spread leaf. The data were automatically collected after 4–5 min with at least four replicates.

**Assay of lipid peroxidation**
Lipid peroxidation as malondialdehyde (MDA) was measured from the third fully expanded frozen leaf (Shi et al. 2010). Approximately 0.3 g of fresh leaf samples were homogenized in 8 mL of 10% trichloroacetic acid (TCA) followed by centrifugation at 4000xg for 10 min at 4 °C. To 2 mL aliquot of the supernatants, 2 mL 0.67% 2-thiobarbituric acid (TBA) were added and the mixture was placed in boiling water (100 °C) for 15 min and then immediately cooled in an ice bath. Finally, the absorption of the supernatant was measured by a spectrophotometer (UV-1280, Shimadzu, Japan) at 600, 532, and 450 nm, respectively. The MDA content was calculated based on the following equation:

\[
C_{\text{MDA}} = 6.45 \cdot (A_{532} - A_{600}) - 0.56 \cdot A_{450} \text{ MDA} \\
= C_{\text{MDA}} \cdot \frac{N}{(W)}
\]

where \(C_{\text{MDA}}\) represents the colorimetric MDA (μmol·L⁻¹), \(N\) represents the total volume of the supernatant (mL), and \(W\) represents the fresh sample mass (g). Finally, the MDA concentration was expressed as μmol·g⁻¹ FW.

**Tissue elemental ion analysis**
Sodium and potassium ion contents were analyzed using the acid digestion method. Dried shoot (leaf and stem) and root samples were ground and passed through a 2-mm mesh sieve. Approximately 0.12 g of the ground powders were completely digested with 3 mL concentrated H₂SO₄ at 200 °C, supplemented with 1.0–1.5 mL H₂O₂ (30% v/v). After digestion, every sample was filled to 50 mL by adding distilled-deionized water to the final volume. Finally, the Na⁺ and K⁺ contents were analyzed by flame spectrophotometer (FP 6410, Shanghai Precision Instruments Co., Ltd., China).

**Salt tolerance evaluation**
The salt tolerance level of the investigated cotton genotypes was assessed by MFV. The MFV of a fuzzy set is a simplification of the indicator function of ordinary sets and exemplifies the extent of reality as an extension of the evaluation (Chen et al. 2012).

The MFV value was calculated via the salt tolerance index (STI), using the following formula:

\[
\text{STI} = \frac{\text{STP}}{\text{CP}}
\]

where stressed plants (STP) represents the mean value of a single trait under salt stress and control plants (CP) represents the mean value of a single trait under nonsaline treatment.

MFV was computed using the following equation:

\[
X_i = \frac{(X - X_{\text{min}})}{(X_{\text{max}} - X_{\text{min}})} \cdot 100\%
\]

where \(X_i\) represents the membership function value of the specific cotton genotype; \(X\) represents the definite measured value of the salt tolerance index in a specific genotype, and \(X_{\text{max}}\) and \(X_{\text{min}}\) represent the maximum and minimum values observed in all-cotton genotypes, respectively (Wu et al. 2019).

The salt tolerance of genotypes was evaluated according to the mean MFVs of each trait and the MFVs of all studied cotton genotypes, ranging from 0 to 1. For each cotton genotype, the mean of MFV was computed as the average of the MFVs of all morpho-physiological traits. A higher mean value of MFV indicated a higher salt tolerance level.

According to Wu et al. (2019), the salt damage index (SDI) was calculated according to the following equation: SDI = 1 - STI.

**Statistical analysis**
Data analysis was performed using SPSS 25.0 (SPSS Inc., USA) and differences between treatments were assessed by ANOVA (Analysis of variance), followed by
DMRT (Duncan’s multiple range test). Data were considered significantly different at $P < 0.05$.

**Results**

**Determination of optimum salt concentration**

In this study, 11 cotton genotypes were subjected to four different salt concentrations and one control treatment to identify the target salt concentration. The SDI of different morpho-physiological traits, such as SL, LN, LA, TB, LRWC, A, gsw, MDA, SK$^+$/Na$^+$, and RK$^+$/Na$^+$, were calculated accordingly. The results were interpreted by linear regression analysis (Fig. 2). When cotton seedlings were treated with 181.5 mmol·L$^{-1}$ NaCl, the SDI of SL decreased to 50% of the total SDI value. Similarly, other traits (e.g., LN, LA, TB, LRWC, A, gsw, MDA, SK$^+$/Na$^+$, and RK$^+$/Na$^+$) decreased to 50% of the total SDI in response to treatments with 186.0,

**Fig. 2** Determination of optimal salt concentration for assessing salt tolerance. The salt concentration of the SDI is the half of each morpho-physiological trait of cotton genotypes.
172.0, 173.5, 180.0, 175.2, 170.0, 155.0, 188.1, and 177.1 mmol·L\(^{-1}\) NaCl, respectively. The average salt concentration of all investigated parameters against SDI was 175.81 mmol·L\(^{-1}\). Therefore, a salt concentration of 200 mmol·L\(^{-1}\) was selected for the current study to compare the salt tolerance among the 11 tested cotton genotypes.

**Genotypic variation under salt stress**

Radar plot analysis based on STI values showed that the cotton genotype exhibited varying responses under salt treatment (Fig. 3a and b). The genotypes Z9807, Z0228, and Z7526 achieved the highest average STI values, while the genotype Z0102 exhibited the lowest STI compared with other cotton genotypes for all considered traits except for the MDA content (Table 2). The STI of the MDA content was high in Z0102, Z9648, Z9612, and Z7780 genotypes, while Z9807 accumulated the lowest MDA under salt stress (Fig. 3b). Additionally, the K\(^+\)/Na\(^+\) ratio assumed a pivotal role in ion homeostasis under salt stress conditions. Genotype Z9807 retained a higher K\(^+\)/Na\(^+\) ratio than all other investigated cotton genotypes by maintaining a higher STI value in both shoot and root tissues.

**Correlation analysis under salt stress**

The correlation coefficient of all studied morphophysiological traits under salt treatment was analyzed by Pearson’s correlation (Table 3). All examined parameters related to salt tolerance indices exhibited significant and

**Table 2** Salt tolerance verification with their mean MFVs

| Genotypes | SL | LN | LA | TB | LRWC | A | gsw | MDA | SK\(^+\)/Na\(^+\) | RK\(^+\)/Na\(^+\) | Mean of MFV |
|-----------|----|----|----|----|-------|---|-----|-----|-----------------|-----------------|-------------|
| Z9807     | 0.783 | 0.921 | 0.819 | 0.671 | 0.861 | 0.739 | 0.707 | 0.400 | 0.866 | 0.754 | 0.897 |
| Z9648     | 0.645 | 0.807 | 0.667 | 0.503 | 0.750 | 0.425 | 0.351 | 0.673 | 0.570 | 0.570 | 0.418 |
| Z0228     | 0.770 | 0.913 | 0.792 | 0.653 | 0.847 | 0.724 | 0.686 | 0.425 | 0.792 | 0.627 | 0.803 |
| Z7526     | 0.767 | 0.925 | 0.783 | 0.632 | 0.843 | 0.692 | 0.647 | 0.490 | 0.779 | 0.620 | 0.756 |
| Z7780     | 0.658 | 0.830 | 0.688 | 0.539 | 0.770 | 0.449 | 0.369 | 0.650 | 0.526 | 0.453 | 0.307 |
| Z0710     | 0.741 | 0.906 | 0.745 | 0.610 | 0.817 | 0.649 | 0.645 | 0.514 | 0.689 | 0.553 | 0.653 |
| Z0102     | 0.625 | 0.769 | 0.625 | 0.483 | 0.728 | 0.395 | 0.314 | 0.729 | 0.490 | 0.397 | 0.100 |
| Z1910     | 0.748 | 0.907 | 0.756 | 0.610 | 0.828 | 0.651 | 0.679 | 0.542 | 0.778 | 0.607 | 0.690 |
| Z7514     | 0.723 | 0.919 | 0.761 | 0.605 | 0.822 | 0.637 | 0.637 | 0.533 | 0.779 | 0.566 | 0.688 |
| Z9612     | 0.652 | 0.820 | 0.667 | 0.499 | 0.757 | 0.439 | 0.338 | 0.656 | 0.542 | 0.441 | 0.220 |
| Z7516     | 0.759 | 0.915 | 0.760 | 0.624 | 0.833 | 0.667 | 0.666 | 0.556 | 0.801 | 0.584 | 0.718 |
positive correlations with each other, except for the MDA content of leaves. Among all traits, LRWC showed the highest positive and negative correlation with most of the traits; e.g., between LRWC and MDA, the strongest (−0.803) negative correlation was observed. In contrast, the most significant positive correlations were observed among LRWC and SK+/Na⁺ (0.918), RK+/Na⁺ (0.917), A (0.900), and LN (0.819). Moreover, A and gsw showed a strong positive correlation (0.939) with each other along with total biomass production (0.830 and 0.865) and LRWC (0.876) of salt-stressed cotton seedlings. The results of Pearson’s correlation coefficient also supported the result of cluster analysis.

**Determining reliable salt tolerance traits**

To identify the most suitable salt tolerance traits, a linear regression fit was performed using the STI of distinct traits and the average membership function value (Fig. 4). The results of this analysis indicated that the mean MFV value was affected by the STI value, and the higher the STI value of each trait, the higher the MFV value (except for MDA). The highest regression value was observed in the LRWC ($R^2 = 0.9921$) followed by A ($R^2 = 0.9873$) between the average of MFV and STI value, while RK+/Na⁺ had the lowest regression value ($R^2 = 0.9196$). The regression values of TB, LA, SL, gsw, SK+/Na⁺, LN, and MDA were 0.986 7, 0.983 9, 0.978 5, 0.965 4, 0.953 5, 0.949 3, and 0.922 3, respectively. Overall, these results indicated that both LRWC and photosynthesis can be considered as two potential traits for the salt tolerance evaluation of cotton genotypes at the seedling growth stage.

**Cluster heat map analysis**

To categorize the genotypes into a homogenous group based on their morpho-physiological traits, cluster heat map analysis was performed, using the Euclidean squared distance metric (Fig. 5). According to the potential characteristics of the genotypes, the studied 11 cotton genotypes were clustered into three statistically significant clusters. Cluster I represented the salt-tolerant group, which was composed of the genotypes Z9807, Z0228, and Z7526. Genotypes Z7516, Z1910, Z7514, and Z0710 were clustered into cluster II, which represented a moderately salt-tolerant group. Cluster III included the genotypes Z0102, Z7780, Z9612, and Z9648, and represented the salt-sensitive group. However, genotype Z9807 showed the highest STI, while genotype Z0102 showed the lowest STI within cluster I and cluster III, respectively.

**Discussion**

Salinity significantly affects growth-related traits by decreasing shoot length, leaf numbers, and leaf area, which collectively decreased the photosynthetic rate and subsequently the total biomass. Leaf area has been identified as an instantaneous response to stress condition (Zhang et al. 2014). The decreased leaf area under salt stress changes the leaf cellular structure, which results in a decrease of the net photosynthetic rate (Munns and Tester 2008). Moreover, biomass production under stress condition represents a prime attribute with which to assess the extent of the stress (Gong et al. 2013). In the current study, all cotton genotypes significantly decreased their total biomass production under salt stress, and significant decrease was found in genotypes Z0102, Z9648, Z9612, and Z7780. These genotypes were identified as salt-sensitive ones (Fig. 3a). Several previous reports have suggested that salt tolerant plants showed less biomass reduction and better growth under salt stress compared with salt-sensitive plants (Ahmed et al. 2013; Singh and Sarkar 2014; Chiconato et al. 2019). The reason may be primarily attributed to the functional impairment of the osmotic potential, followed by ionic imbalance, predictably leading to a nutritional disproportion in plants (Meloni et al. 2001; Alharby et al. 2019). Another possible mechanism for the lower biomass accumulation may be the diversion and/or exchange of potential energy from plant growth to sodium ion exclusion. Alternative mechanisms are the synthesis of

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**Table 3** Pearson’s correlation among different traits of cotton genotypes in the presence of 200 mmol·L⁻¹ NaCl treatment

| Traits | SL     | LN     | LA     | TB     | LRWC   | A      | gsw    | MDA    | SK+/Na⁺ | RK+/Na⁺ |
|--------|--------|--------|--------|--------|--------|--------|--------|--------|---------|---------|
| SL     | 1      |        |        |        |        |        |        |        |         |         |
| LN     | 0.868a | 1      |        |        |        |        |        |        |         |         |
| LA     | 0.675a | 0.705a | 1      |        |        |        |        |        |         |         |
| TB     | 0.767a | 0.778a | 0.869a | 1      |        |        |        |        |         |         |
| LRWC   | 0.721a | 0.819a | 0.728a | 0.713a | 1      |        |        |        |         |         |
| A      | 0.775a | 0.834a | 0.770a | 0.830a | 0.900a | 1      |        |        |         |         |
| gsw    | 0.783a | 0.844a | 0.798a | 0.865a | 0.876a | 0.939a | 1      |        |         |         |
| MDA    | -0.676a| -0.781a| -0.571a| -0.492a| -0.803a| -0.762a| -0.673a| 1      |         |         |
| SK+/Na⁺| 0.756a | 0.758a | 0.797a | 0.785a | 0.918a | 0.856a | 0.833a | -0.707a| 1       |         |
| RK+/Na⁺| 0.757a | 0.755a | 0.785a | 0.788a | 0.917a | 0.869a | 0.839a | -0.711a| 0.995a  | 1       |

*aCorrelation is significant at the 0.01 level
compatible solutes (Munns and Tester 2008) or the direct effect of decreased photosynthesis (Haider et al. 2019). Similar inhibitions of growth and development induced by salt-stress have been reported for salt-stressed tomato (Gong et al. 2013), wheat (Elkelish et al. 2019), melon (Sarabi et al. 2017), and maize (Hessini et al. 2019).

The relationship between photosynthesis and salt stress is remarkably complex, and depends on the salt concentration, the duration of salt stress in the growth medium, and species (Chiconato et al. 2019). The potential reduction of stomatal conductance may also inhibit the photosynthetic capacity in most glycophytes, including cotton (Brugnoli and Lauteri 1991). In the present study, the photosynthetic rate decreased significantly with increasing salt concentration in the growth media. The genotypes Z9807, Z0228, and Z7526 showed the lowest photosynthesis reduction compared with control (Fig. 3b).

Fig. 4 The linear fit regression between STI of each morpho-physiological trait and mean MFV of cotton genotypes
This result indicated that these three genotypes were able to maintain a higher photosynthetic rate by up-regulating their rubisco activity. Similar findings were reported in salt-stressed cotton genotypes, e.g., the salt-tolerant genotype CCRI-79 maintained the maximum photosynthetic rate compared with the salt-sensitive genotype Simian-3 (hang et al. 2014). The gsw also decreased under salt treatment (Kawakami et al. 2013) and a significant decrease was also observed in the present study. The possible mechanism of gsw reduction under salt stress may be the increase of abscisic acid levels (He and Cramer 1996) and/or the decrease of stomatal aperture (Ahmed et al. 2013). This effect has also been reported for salt-stressed cotton (Kawakami et al. 2013; Zhang et al. 2014), pepper (Penella et al. 2015), date palm (Al Kharusi et al. 2017), melon (Sarabi et al. 2019), and sugarcane (Chiconato et al. 2019).

The LRWC is a key physiological trait that can also be used as an important indicator to discriminate the salt tolerance ability among genotypes (Sinclair and Ludlow 1986; Sánchez-Rodríguez et al. 2010). Under salt stress, the water potential decreases and inhibits the uptake of water by the roots, thus decreasing the LRWC (Munns and Tester 2008; Hassanvand et al. 2019). In the current study, the LRWC decreased prominently in the salt-sensitive group (Z0102, Z9648, Z9612, and Z7780) compared with the salt-tolerant group. Previous studies corroborated this result, suggesting that salt-sensitive genotypes may not be able to uphold the water status as they accumulate more soluble salts compared with salt-tolerant genotypes (Al Kharusi et al. 2017; Sarabi et al. 2017; Aghaie et al. 2018). Furthermore, a highly significant positive correlation was found between the LRWC and the shoot-root K+/Na+ ratio (Table 3). This indicated that plants with a higher K+/Na+ ratio retained more water in their leaf tissues under salt stress.

Salt stress primarily affects plants by osmotic stress, which is followed by ionic stress and perturbed ion homeostasis, resulting in a change of metabolic and cellular acclimation in plant physiology (Fig. 6). Na+ is often considered as the most toxic ion in a saline environment because of its excessive cytosolic concentration; moreover, frequent Na+ accumulation is also deleterious for plants (Munns and Tester 2008; Munns et al. 2016). To avoid excessive Na+, salt exclusion and/or ion sequestration into vacuoles represent prominent mechanisms of salt-stress adaptation (Liu et al. 2018; Wang et al. 2019). In the present study, all cotton genotypes accumulated inorganic Na+ at different extents. Salt tolerance genotypes maintained higher K+/Na+ ratios compared with salt-sensitive genotypes such as Z0102. Specifically, salt-tolerant genotypes exhibited the highest K+/Na+ ratios in the shoot and root tissues. This might be the result of the induction of salt-related genes, such as SOS1 and HKT1, which prevented excessive Na+ loading in the xylem sap, thus reduced the accumulation of lethal Na+ and maintaining the proper K+ ratio in cells (Assaha et al. 2017). This result corroborates other research findings (Tester and Davenport 2003; Chen et al. 2005; Rahneshan et al. 2018).
Apart from inorganic ion homeostasis, salt stress generates ROS, thus causing oxidative damage as secondary damage in plant cells. Excessive accumulation of ROS in salt-treated cotton seedlings might accelerate the Haber-Weiss reaction and consequently amplify both lipid peroxidation and cell membrane damage (Gill and Tuteja 2010; Borzouei et al. 2012). As expected, the leaf MDA content was lowest in genotype Z9807, whereas the highest induction was observed in genotype Z0102 grown under salt stress for 10 days (Fig. 3b). The lowest MDA accumulation under salt stress was also observed in the salt-tolerant CSSL line of rice compared with the salt-sensitive CSSL line (Nounjan and Theerakulpisut 2012). This result agrees with the results reported for other plant species, such as strawberry (Garriga et al. 2015), melon (Sarabi et al. 2017), rice (Wang et al. 2018), and mungbean (Alharby et al. 2019). Moreover, a highly significant negative correlation was observed between MDA content and LRWC as well as photosynthesis (Table 3), suggesting lipid peroxidation, caused by salt stress, as one of the main reasons for the decrease of the tissue-water content and photosynthetic rate in cotton genotypes.

Salt stress perturbs the morphological, physiological, and biochemical changes of plants at the seedling growth stage. A number of parameters should be assessed for an appropriate evaluation of salt tolerance; however, not all traits may be effective or reliable for the screening of genotypes. It is also well known that the evaluation process for reliable salt-tolerance traits is the crux of a successful salt-tolerance breeding program (Zeng et al. 2002). Therefore, such evaluation process is needed to identify a reliable salt-tolerance screening method for the effective evaluation of the salt tolerance of cotton genotypes at the seedling growth stage. The current study performed a multiple regression analysis to identify reliable salt-tolerance traits between the mean MFV and the STI of each individual morphophysiological trait of the studied cotton genotypes. The mean MFV was calculated based on the average of

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**Fig. 6** Schematic summary of salinity mediated physiological changes
MFVs of shoot length, leaf numbers, leaf area, total biomass, photosynthesis, stomatal conductance, malondialdehyde content, as well as shoot and root K⁺/Na⁺ ratios of each genotype. The average MFV was considered as multiple indicators to assess the salt tolerance of plants, higher MFV value indicates a higher salt-tolerance level (Table 2). According to the data obtained by the present study, the STI of LRWC ($R^2 = 0.992$) and $A$ ($R^2 = 0.982$) represent reliable traits compared with the other eight morpho-physiological traits, based on the regression value between the mean of MFV and multiple indexes (Fig. 4). Thus, both the LRWC and photosynthesis traits might be used as reliable indicators for the salt-tolerance assessment of cotton genotypes at the seedling stage. In a previous study, Suriya-Arunroj et al. (2004) suggested that LRWC could be used as an effective indicator for the salt-tolerance evaluation of rice at the seedling stage. Another study recommended the root and shoot length as primary indicators for the salt tolerance of Brassica napus (Long et al. 2013).

Cluster heat map analysis was performed to classify the genotypes into homogenous groups based on the STI value of all studied traits. According to the results of this cluster analysis, the 11 cotton genotypes were divided into three major clusters with respect to their salt-tolerance levels (Fig. 5). In earlier studies, cluster analysis distinguished 12 rice cultivars (Chunthaburee et al. 2015) and eight almond species (Sorkheh et al. 2012) into four and three major clusters, respectively, based on their physiological traits. The result of the present study showed that the genotypes of cluster I exhibited superior performance under salt stress and achieved higher LRWC, photosynthesis, total biomass production, and higher shoot-root K⁺/Na⁺ ratios compared with other genotypes (grouped within the other two clusters). Moreover, lower MDA content caused less oxidative damage in these genotypes, which could also be used as an indicator for salt-tolerant genotypes. This result also suggests cluster analysis as an important data-mining tool for discriminating cotton genotypes according to their salt-tolerance levels based on their morpho-physiological features. This should be further investigated at different growth stages of cotton plants to explore the genotypic discrimination in response to salt stress.

Conclusions

In conclusion, the results of this study identified substantial effects of salt stress on the studied traits, and a noteworthy genotypic difference was found. Screening clearly identified three distinct groups: salt tolerant (showing superior tolerance), moderately salt tolerant (showing intermediate tolerance), and salt sensitive (showing susceptibility to salinity). Within the salt-tolerant groups, Z9807 showed the best tolerance potential, followed by Z0228 and Z7526. This suggests that these genotypes are prominent resources because of their salt tolerance, and could thus be used as genetic materials for the further breeding framework. In addition, LRWC and $A$ were identified as two reliable salt-tolerance traits with which the cotton genotypes at the seedling growth stage can be evaluated. Therefore, these findings contribute to the breeding program of salt-tolerant cotton genotypes.

Abbreviations

$A$: Photosynthetic rate; gsw: Stomatal conductance; LA: Leaf area; LN: Leaf number; LRWC: Leaf relative water content; MDA: Malondialdehyde; MFV: Membership function value; $R^2$: Root potassium sodium ratio; SDI: Salt damage index; $SK^+/Na^+$: Shoot potassium sodium ratio; SL: Shoot length; STI: Salt tolerance index; TB: Total biomass

Authors’ contributions

Sikder RK conducted the experiment and drafted the manuscript. Wang XR performed part of the statistical analysis and revised the manuscript. Jin DS assisted in data collection and part of the biochemical analysis. Zhang HH helped to prepare the graphs. Gui HP, Dong Q, and Pang NC helped with data collection. Song MZ and Zhang XL conceived the study, participated in its design, and helped to revise the manuscript. All authors read and approved the final manuscript.

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Availability of data and materials

All data generated or analyzed during this study are included in this published article.

Ethics approval and consent to participate

Not applicable.

Consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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