Cloning of *SjCA* gene and its expression analysis on upland cottons

Jingjing Kong, Xuke Lu, Xiaojie Zhao, Zujun Yin, Shuai Wang, Delong Wang, Junjuan Wang, Weili Fan, Wuwei Ye

State Key Laboratory of Cotton Biology / Institute of Cotton Research, Chinese Academy of Agricultural Sciences, Anyang, Henan, P. R. China.

Correspondence: Wuwei Ye. Address: Institute of Cotton Research, Chinese Academy of Agricultural Sciences, Anyang, Henan, 455000, China. Email: yew158@163.com

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Abstract

Acquisition of salt-tolerant genes from exogenous plants to improve cotton salt resistance has always been a hotspot of research on cotton salt resistance. However, the information regarding the method of conversion of living cotton pollens by portable gene gun technology is still little. Complete sequence information of the *Carbonic Anhydrase (CA)* gene was obtained from NCBI database, and its full ORF length sequence was cloned by RT-PCR technology. After the construction of pBI121-CA::GFP fusion expression vector, we used cotton pollens from upland cotton varieties Y-2067, ZA-23, and GZ-2, which have weaker autofluorescence, to conduct the research on the transient expression of cotton pollens in vivo via particle bombardment (gene gun technology). The results indicated that green fluorescence enhancement of the three kinds of cotton pollens was realized after the CA gene transformation, meaning that the CA gene expression level was increased. Besides, the salt tolerance germination ability of transgenic T1 seeds was also improved. Our research initially established a transient expression system of cotton pollen in vivo via particle bombardment technology, laying the theoretical foundation for further research of cotton genetic transformation and creation of cotton salt-tolerant germplasm.

Key words
Transformation, *Carbonic Anhydrase (CA)*, *Saccharina japonica*, Cotton, Gene gun technology

1 Introduction

Due to the continuous land utilization for repeated agricultural activities and the excessive irrigation, the spread and use of saline-alkali soil has become a worldwide ecological problem. Nowadays, the area of global saline-alkali soil is increasing and restricting the world agricultural production. Therefore, the development and exploitation of saline-alkali land has a huge commercial value and important social significance. Planting salt-tolerant crops is one of the accepted ways to make use of saline-alkali soil. As an important pioneer and cash crop, cotton is one of the most appropriate plants for cultivation on saline-alkali soil. Consequently, the improvement of cotton salt tolerance by gene engineering and transgenic technology has been an urgent task and major direction of cotton breeding [1].
Recently, scientists conducted a large number of researches on the molecular mechanism of plant salt tolerance and found a great number of salt tolerance-related proteins, including enzymes involved in the synthesis of a small organic molecular penetrant, a protective agent, and a series of enzymes participating in salt absorption, translocation, and the regional distribution in cells\textsuperscript{[2-5]}. Besides, many salt-tolerant genes were cloned. Carbonic anhydrase (CA, EC 4.2.1.1) containing a zinc atom was cloned and identified to catalyze the reversible conversion between carbon dioxide and bicarbonate radical (\(\text{CO}_2 + \text{H}_2\text{O} \leftrightarrow \text{HCO}_3^-\))\textsuperscript{[6, 7]}, which plays an important role in many biological processes, such as ion exchange, respiration, pH stabilization, calcification, CO\(_2\) capture, and photosynthesis\textsuperscript{[8-10]}. Since 1933, when the CA gene was found in red blood cells, it has also been discovered in all types of mammalian histiocytes. In recent years, scientists have confirmed that CA gene exists in other living organisms, such as higher plants, prokaryotic and eukaryotic algae, bacteria, and other microorganisms\textsuperscript{[11-13]}. According to its amino acid sequence and crystal structure, the known CA could be divided into three forms: \(\alpha\)-CA, \(\beta\)-CA, and \(\gamma\)-CA. However, the sequence similarities between the three CA enzymes were so low that they were assumed to be formed through different evolutionary paths. Among them, \(\beta\)-CA exists mainly in the chloroplasts of bacteria, algae, and monocotyledon and dicotyledon plants\textsuperscript{[14-16]}.

According to some research findings, there is a relationship between CA response and environmental stress. Previous studies have documented that the change of cotton photosynthetic rate was consistent with the change of CA activity under various conditions (drought, high temperature, and high salinity), showing CA in cotton was a gene related with abiotic stresses\textsuperscript{[17, 18]} . Previous reports showed that the CA gene expression level and enzyme activity in triticale increased under salt stress, suggesting that CA played a part in the response to salt stress\textsuperscript{[19]}. According to the examination findings of Yu et al., the similar result was also found in rice\textsuperscript{[20]}. Fisher et al. cloned the CA gene of strong salt tolerance from Dunaliella salina and found that the expression levels of mRNA and protein were enhanced along with the increase of the concentration of sodium chloride in the nutrient solution\textsuperscript{[21]}.

With the advance of research on plant genome, increasing more genes have been included in studies conducted across the world, and the gene transient expression technology has become a fast and effective way for gene expression investigation. Agrobacterium tumefaciens is a soil phytopathogen that could infect plant by the wound sites and causes crown gall disease via delivery of T-DNA from bacterial cells into host plant cells through a bacterial type IV secretion system\textsuperscript{[22]} . Although Agrobacterium is a wound-associated pathogen, it is still considered a better way to transfer DNA into diverse host cells or tissues under unwounded conditions\textsuperscript{[23-25]}. Compared with the traditional transgenic process, transient expression has several advantages via gene gun method, such as simple operation, short cycle period, and high transformation and expression efficiency. Gene gun is a method in which small metal particles, for example gold powder, wrapped with vector DNA are used to bombard the recipient cells to realize the transformation of DNA, which is also termed as particle bombardment technology. What is more, the stably expressed transgenic plants may cause problems of food chain pollution and gene flow. On the other hand, transgenic expression is an independent process that does not produce hereditable offspring, which contributes to its high bio-safety and reliable results. But the transient expression in cotton pollens was a new, easier and more effective method to study gene expression patterns and obtain transient offspring. Until now, no report has been published concerning the transgenic expression of cotton pollen genes; thus our study provided a new approach to investigate transgenic expression of pollen genes.

The typical, highly salt-tolerant alga, Saccharina japonica, was chosen because it exhibits high homology with the same type of protein from other species, including Dunaliella salina and rice. We speculated that may play a positive role in the response to salt stress. Thus, we tried to transform the CA gene from Saccharina japonica into cotton pollens via the Gene Gun technology to obtain transgenic salt-tolerant plant material. After transforming the cotton pollen, we tested the expression level and established a system of transgenic expression for cotton pollen transformation via particle bombardment, providing the basis of further engineering of salt-tolerant genetic material.
2 Materials and methods

2.1 Experimental materials
Gametophyte material from *Saccharina japonica* was provided by the Ocean University of China, Qingdao, Shandong Province, China. *Saccharina japonica* is belonged to Laminariales, Phaeophyta, one of the most valuable seaweeds. *Saccharina japonica* female and male gametophyte clonal lines were preserved as the previous methods in the PES culture medium [26], and the conditions were below: Culture temperature was 17±℃, light intensity was 40 μmol photons /m2•s, and photoperiod was 16 h (day) and 8 h (night). *Saccharina japonica* female and male gametophytes were collected and crushed with grinder (Philips) into suspension and then centrifuged to collect gametophytes in 100 ml conical flask containing 15 g fresh gametophytes each. Fresh gametophytes were grinded into superfine powder to extract RNA. The total RNA of *Saccharina japonica* was extracted using EASY spin Plant RNA Kit (Aidlab), and the first chain of cDNA was completed with TransScript all in one kit (TRANS). First strand cDNA was obtained in a 20 μl reaction system, containing 3 μl RNA (330 ng/μl), 4 μl All in one supermix, 1 μl gDNA remover and 12 μl RNase-free water, and the reaction was completed at 42°C for 15 min. At last the tube was subjected to 85°C to end the reaction. Products were kept at -20°C for use.

2.2 Obtaining the complete ORF of CA gene
The complete information of cDNA from *Saccharina japonica* was downloaded through searching the existing nucleic acid database in NCBI. To amplify the completed ORF sequence, we designed primers by use of Oligo 6.0 software, CA-F1: 5’-CATCACCATGACTTTCACG-3’ and CA-R1: 5’-GTCCACTACTAATGCTCTC-3’. By using cDNA as a template and through the action of LA Taq DNA Polymerase, we obtained the complete ORF sequence. The target bands (TaKaRa, Takara Bio Inc) in the gel were recovered and ligated to the vector pGEM-T Easy (Promega, USA). The ratio of vector and PCR product adapted in the ligation system was 1: 3. The ligation system was composed with pGEM-T Easy vector 1 μl, PCR product 3 μl, T4 DNA ligase 1 μl and T4 DNA ligase 2×quick ligase buffer 5 μl. After the transformation of competent *Escherichia coli* cells DH5α (TaKaRa), the positive clones were selected and sequenced for verification.

2.3 Bioinformatics analysis
The sequencing results were analyzed by DNAStar (DNAStar, Madison, WI, USA). The multiple protein sequences were compared by using the DNAMAN software (Lynnon LLC., San Ramon, CA, USA), and the phylogenetic tree of CA was analyzed via MEGA5.0 and built through the neighbor-joining method. Secondary structure prediction was realized by an analysis of the protein function via the DNAStar software.

2.4 Construction of fluorescent fusion protein expression vector
As a receptor vector, we used pBI121-GFP, provided by the research group working on cotton early-ripening traits at the Institute of Cotton Research of CAAS, China, and chose two enzyme sites, *Xba* I and *Sma* I. The complete ORF sequence was regarded as the target fragment. Infusion primers were designed by the utilization of the online primer design software of the Clontech company.

After amplifying and recovering the target fragments, the fusion protein transient expression vector pBI121-CA::GFP was constructed with in-fusion technology. The primers used were designed with online program http://bioinfo.clontech.com/infusion. And the forward primer was CA-F2: 5’-CAGGGGGAGCTC TAGAATGACTT TCACGGCGCACCT-3’, and the reverse primer was CA-R2: 5’-AGGGACTG ACCACCGGGGTAGTGGACGAAGAAGATCT-3’. Ligation system adapted 50 μl, containing cDNA 0.5 μl, up/downstream primers 1μl each, dNTP 1μl, taq polymerase 1 μl and ddH2O 45.5 μl. The competent cells of *Escherichia coli* DH5α were used to perform transformation before further sequencing and identification of the positive clone results.
2.5 Green auto-fluorescence detection of the cotton pollen

Twenty-six cotton varieties (were identified by our laboratory for many years) were chosen, and their pollens were collected. Green auto-fluorescence detection was performed by confocal laser scanning microscopy (FV1000, Olympus, Japan) at a wavelength of 488 nm (excitation light was green at the wavelength of 488 nm). We only selected cotton varieties with weak green auto-fluorescence from the 26 cotton varieties for the interference of strong green auto-fluorescence.

2.6 Transformation of the cotton pollen via in vivo particle bombardment technology

Gold powder (1 µm diameter of the particles) was washed with 70% ethanol and sterile purified water. Gold powder suspension liquid was prepared by adding sterile purified water. First, 30 mg was added into a 1.5 ml tube containing 1ml 70% ethanol, shaking for 5 min; then centrifuged for 30 s at 12,000 rpm after standing for 15 min and discarded the supernatant; then 1 ml ddH2O was added into the tube to re-suspend the gold powder and re-centrifuged for 15 s at 12,000 rpm and discarded the supernatant. Ethanol and ddH2O were used to re-cycle the steps above. At last, gold power washed with Ethanol and ddH2O was suspended in 500 µl ddH2O.

Aliquots of 100 µl gold powder suspension liquid were subjected to shaking after adding the plasmid pBI121-CA::GFP. Meanwhile, pre-cooling 40 µl 0.1 mol·l-1 spermidine and 100 µl 2.5 mol·l-1 CaCl2 were added into the suspension liquid, dropwise. After mixing and centrifugation, the supernatant was discarded. Then 200 µl ethanol was used to wash and re-suspend the mixture of plasmid and gold powder. The products were saved at -20°C for further use.

The gene delivery tool GDS-80 (Wealtec corp) was utilized to bombard the cotton pollens. The pressure of the helium tank was adjusted to 1,350 psi, and 10 µl of particle suspension wrapped in DNA was added into the center of GDS-80 before the bombardment. The pollens were collected the day before and stored at room temperature for the bombardment. Finally, the pollens bombarded were stored overnight without light at 25°C. Then a brush was used to dip in the pollens bombarded and dropped the pollens on a new sterile glass slide. Then a drop of ddH2O was added on the pollens to make a temporary slide to be observed. The gene transient expression was observed with Laser Confocal Scanning Microscope (FV1000) at a wavelength of 488 nm.

3 Results

3.1 The cloning of CA gene from Saccharina japonica and its bioinformatics analysis

By PCR amplification, complete CA gene ORF was obtained: a product with a size of 873 bp, which encoded 290 amino acids (see Figure 1A). Another 10 species were chosen for a multiple sequence comparison. By performing BLAST in NCBI for CA gene-coded amino acid sequence, homology with the proteins of other species was observed. Among them, the similarity between CA and Dunaliella salina, Oryza sativa Japonica, Sinorhizobium meliloti, and Oncorhynchus mykiss reached 89%, 79%, 78%, and 70%, respectively. The ten pre-selected representative species for a multiple sequence comparison were as follows: Chlorella sorokiniana (BAA28217), Gossypium hirsutum (AAM22683), Sinorhizobium meliloti (NP_435267), Dunaliella salina (AAO83593), Oryza sativa Japonica Group (BAD29283), Pasteurella multocida (AAK03989), Human Carbonic Anhydrase Iv (pdb|1ZNC|A), Oncorhynchus mykiss (AAR99330), Tetraodon nigroviridis (CAG08972), and Gallus gallus (XP_415893) (see Figure 1B). An evolutionary tree was constructed to analyze the evolutionary relationship of CA gene in different species (see Figure 1C). The sequence comparison results indicated high conservation with other species and a low similarity with the cotton sequence, meaning
that the CA proteins of cotton and Saccharina japonica belonged to different kinds. The phylogenetic tree findings showed that the CA of Saccharina japonica had a short evolutionary distance from the CA of Dunaliella salina, meaning that a closer genetic relationship was existent. The two CAs belonged to one type, and we speculated that CA may play a vital role in the salt stress environment. However, the CA of Saccharina japonica and the CA of upland cotton had a long evolutionary distance, and we assumed that they pertain to different kinds. We imported the CA gene sequence of a gametocyte from Saccharina japonica into the Protean (DNASTar, Madison, WI, USA) in DNASTar for a secondary structure prediction and the results revealed that the structure included α-helices, β-sheets, torsion angle, and a random coil. The prediction result of CA gene location with online program PSORT (http://psort.hgc.jp/form.html) showed that the gene was located on plasma membrane or endoplasmic reticulum (membrane), so we speculated this protein was a membrane protein. Besides, online program ProtScale (http://web.expasy.org/protscale/) was used to predict the hydrophilic-hydrophobic property of the CA protein and the result indicated this protein was a hydropathicity protein. We also conducted the signal-protein prediction (see Figure 2A), and the results indicated that there was a typical signal-protein digestion site at 20-21 (Gly-Val), which site would be important for the function of CA gene.

![Figure 1](http://jbei.sciedupress.com)

**Figure 1.** Amplification, multiple alignment and phylogenetic analysis of CA gene. (A) Amplification of CA gene ORF. 1: ORF frame gene PCR product; M: DNA marker SM. (B) Alignment of the deduced amino acid sequence of SjCA and that from other species. Identical residues are shaded in black and similar residues in grey, the box indicates the conservative domain and the red arrow shows the Zn-binding sites. (C) Phylogenetic tree of putative CA gene from Saccharina japonica and other species.

### 3.2 The construction of the expression vector of fluorescent fusion protein

We inserted the CA gene into the plant expression vector pBI121-GFP to construct a fusion protein expression vector. After conversion to the competent cells of Escherichia coli DH5α, the positive clone was screened by kanamycin and was selected by colony PCR identification and sequencing. The results of the sequence comparison and further digestion insertion were correct, demonstrating that the expression vector was constructed successfully. We named the constructed expression vector pBI121-CA::GFP whose detailed description was displayed in Figure 2B.
**Figure 2.** Signal-proteins prediction and the construction of fluorescence fusion expression vector of CA gene. (A) The prediction results of signal-proteins of CA gene. (B) The construction of fluorescence fusion expression vector of CA gene with pBI121-GFP.

**Figure 3.** Spontaneous green fluorescent of cotton pollen and transient expression analysis of CA gene by gene gun transformation in cotton pollen (wavelength: 488 nm). (A) Green fluorescent spontaneously of cotton pollen. a) H-109; b) H-177; c) ZA-23. (B) Transient expression of CA in cotton. A1, B1, C1, D1, E1, F1: Fluorescence images; A2, B2, C2, D2, E2, F2: Bright-field pollen grain images; A3, B3, C3, D3, E3, F3: Merged images of A1 and A2, B1 and B2, C1 and C2, D1 and D2, E1 and E2, F1 and F2, respectively.
3.3 Detection of green auto-fluorescence emission of the cotton pollen

The clone of the salt-tolerance gene CA and the construction of the green fluorescent protein (GFP) fusion expression vector were accomplished in preliminary work. The results indicated that green auto-fluorescence interference existed in every cotton varieties. Until now, there have been no reports establishing whether the pollen emitted green auto-fluorescence. Therefore, in our study, we tried to verify whether the green auto-fluorescence of the pollen existed, and whether different pollens had different green fluorescence intensities. So we selected cotton varieties with weak green fluorescence to verify the expression level of the foreign gene CA of Saccharina japonica in the cotton pollens.

Table 1. Materials and the results

<table>
<thead>
<tr>
<th>Varieties</th>
<th>Spontaneous fluorescence intensity of Pollen</th>
</tr>
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<tbody>
<tr>
<td>Z-07</td>
<td>strong</td>
</tr>
<tr>
<td>Z-12</td>
<td>strong</td>
</tr>
<tr>
<td>Z-41</td>
<td>strong</td>
</tr>
<tr>
<td>Z-221</td>
<td>strong</td>
</tr>
<tr>
<td>Z-9409</td>
<td>strong</td>
</tr>
<tr>
<td>Z-9806</td>
<td>strong</td>
</tr>
<tr>
<td>H-109</td>
<td>strong</td>
</tr>
<tr>
<td>H-177</td>
<td>strong</td>
</tr>
<tr>
<td>BPA-68</td>
<td>strong</td>
</tr>
<tr>
<td>TM-1</td>
<td>strong</td>
</tr>
<tr>
<td>GK-50</td>
<td>strong</td>
</tr>
<tr>
<td>GK-19</td>
<td>strong</td>
</tr>
<tr>
<td>X-2</td>
<td>strong</td>
</tr>
<tr>
<td>L-1138</td>
<td>strong</td>
</tr>
<tr>
<td>HU-749513</td>
<td>strong</td>
</tr>
<tr>
<td>NG-1</td>
<td>strong</td>
</tr>
<tr>
<td>NG-2</td>
<td>strong</td>
</tr>
<tr>
<td>KHW-164</td>
<td>strong</td>
</tr>
<tr>
<td>Z-701</td>
<td>strong</td>
</tr>
<tr>
<td>Z-710</td>
<td>strong</td>
</tr>
<tr>
<td>ZNG-191</td>
<td>strong</td>
</tr>
<tr>
<td>ZNG-192</td>
<td>strong</td>
</tr>
<tr>
<td>ZA-16</td>
<td>strong</td>
</tr>
<tr>
<td>Y-2067*</td>
<td>weak</td>
</tr>
<tr>
<td>ZA-23</td>
<td>weak</td>
</tr>
<tr>
<td>GZ-2</td>
<td>weak</td>
</tr>
</tbody>
</table>

Note. *Y-2067 was used as a control to categorize the fluorescence intensities. Cotton varieties with stronger fluorescence intensities than Y-2067 were termed as strong and others were weak.

We observed the green auto-fluorescence of 26 cotton pollens by FV1000 at a wavelength of 488 nm. The results indicated that all of the pollens had a different degree of green auto-fluorescence. Table 1 summarizes the different fluorescence intensities. For example, H-109 (see Figure 3A-a) and H-177 (see Figure 3A-b) were representative materials with strong auto-fluorescence. By contrast, ZA-23 was a typical material with weak green auto-fluorescence (see Figure 3A-c). Among them, Y-2067, ZA-23, and GZ-2, which were three upland cotton materials, had weaker green auto-fluorescence. Hence, we chose the pollens of Y-2067, ZA-23, and GZ-2 for pBI121-CA::GFP conversion to investigate the transient expression. The remaining 23 materials with strong auto-fluorescence were no longer used.

3.4 Transient expression of CA gene in cotton pollen analysis

We performed particle bombardment of cotton pollens with gold powder and plasmid suspension. Then we observed them via confocal laser scanning microscopy (see Figure 3B) after overnight culture at room temperature. We found that green
auto-fluorescence intensities of all of the three pollens were enhanced, meaning the gene expression was successful and the foundation was laid for the development of salt-tolerant materials via particle bombardment of cotton pollens. The results indicated that Y-2067, ZA-23, and GZ-2 normally had weaker green auto-fluorescence (A1, A2, A3; C1, C2, C3; E1, E2, and E3). However, after the particle bombardment, the green auto-fluorescence was enhanced obviously (B1, B2, B3; D1, D2, D3; F1, F2, and F3), revealing that the expression of CA gene in the pollens was successful, and the gene particle bombardment is a practicable and reliable method.

3.5 Gene gun living conversion of CA gene in cotton

Gene living conversion in plants has become a hotspot these years. In this study, Bt cotton varieties, Hu749513 and F12, and non-Bt cotton varieties, GK50 and Yu 2067, were selected to conduct the gene gun (GDS-80) living conversion of CA gene in the fields. The vector pBI121-CA mixed with gold powder was suspended in absolute ethyl alcohol. The cotton flowers of the selected varieties were harvested on the previous day and kept on a super clean experimental table at room temperature. Cotton pollens were collected from the flowers, harvested into sterile petri dishes, and bombarded by the portable gene gun GDS-80 containing 10 µl vector mixture. Then, the bombarded pollen was used to pollinate other flowers, which were subsequently marked (see Figure 4A). When the seeds matured, we harvested the transgenic seeds and performed a molecular analysis.

![Image of gene gun transformation and molecular detection](http://jbei.sciedupress.com)

**Figure 4.** The flow chart of gene gun transformation in the field and molecular detection result of transgenic cotton plants. (A) The flow chart of gene gun transformation in the field. a) the cotton in the field; b) getting rid of stamens, putting on the candle tube and marking the flowers; c) the flowers collected; d) the pollen collected; e) gene gun bombardment; f) pollination. (B) Molecular detection result of the CA gene.
3.6 Salt tolerance detection of T1 seeds of transgenic cotton

We obtained 162 T1 seeds after the CA gene conversion into non-insect-resistant cotton (Hu 749513) and compared them with the non-transgenic seeds of Hu 749513. We took 40 full and uniform seeds and subjected them to the sandwich filter paper cultivation method. After the treatment with 0.8% NaCl solution, the seeds were cultivated and left to germinate in an illumination incubator with intervals of 14 h in the light at 28°C and 10 h in the dark at 25°C. The process was repeated three times, and the germination data were calculated. As shown in Table 2, the transgenic seeds had higher salt-tolerance ability during germination than those in the control group, meaning that the CA gene improved the salt tolerance of seeds.

Table 2. Transgenic salt resistance testing

<table>
<thead>
<tr>
<th>Materials</th>
<th>Seeds in total</th>
<th>Number of shoots</th>
<th>Germination rate (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Transgenic seeds of CA gene</td>
<td>120</td>
<td>27**</td>
<td>22.5</td>
</tr>
<tr>
<td>Control</td>
<td>120</td>
<td>1</td>
<td>0.08</td>
</tr>
</tbody>
</table>

We planted the germinated seeds in a greenhouse and performed DNA detection assays in young leaves. Aiming at the identification of the CA gene, we designed primers (F: ATGACTTTCAACGCGCACCTAC; R: TTAGTAGTGA CGAAGAAGATC) and detected the salt tolerance gene, which confirmed the successful transgenosis (see Figure 4B).

4 Discussions

The response of plants to salt stress is controlled by complicated polygenes, involving many physiological, biochemical, and cellular changes, as well as protective, defensive, and salt tolerance-related gene expression. Extensive research on salt-tolerance genes has been conducted in the world. Under the condition of salt stress, plant genes are selectively expressed in order to improve the resistance ability. The studies on salt-tolerant genes would improve the knowledge and understanding of the underlying mechanisms of action and, by their contribution to discovering substantial theoretical evidence, they will facilitate the realization of further transgenic research [27, 28].

α-, β-, and γ-CA have been reported to be distributed widely in microalgae and higher plants, are zinc-metalloenzymes that catalyze the reversible interconversion of CO₂ and HCO₃⁻ [29]. In higher plants, CAs play a vital role in the process of photosynthesis, respiration, ion absorption [30]. In the study, germination capacity of T1 transgenic seeds increased, which may be because that CA gene enhanced the respiration of transgenic respiration. The enzyme activity of CA is kind of photosynthetic carbon metabolism enzyme in adapting stress conditions and could be induced by the substrate, playing an important function in keeping stable photosynthetic carbon metabolism [31]. Previous studies showed that AtaCA1 protein may be transported into chloroplast through the secretory pathway [32] and β-CA6 was located in mitochondria [33]. This suggested that different CAs have different locations, which may lead to different functions.

Nowadays, there are a large number of reports concerning genetically modified (gm) events. GhMAPK is the first MAPK gene cloned in Gossypium hirsutum, which plays an important role in salt stress response. Yamada et al. cloned the salt-tolerant gene mangrin from Bruguiera sexangula (Lour.) Poir. and converted it in Escherichia coli, yeasts, and tobacco, improving the salt tolerance of the obtained transformant [34]. Yu et al. converted the salt-tolerant gene mangrin into Eucalyptus and also acquired a transformant with enhanced salt tolerance [35]. Gao et al. obtained wheat plants with higher salt and drought tolerance by GmDREB gene transgenosis [36]. Lee et al. converted TPS gene of Escherichia coli into tobacco to improve its salt tolerance [37]. Besides, Ganesan et al. isolated the AmMYB1 gene from grey mangrove and incorporated it into tobacco to acquire high-salt-tolerant plants [38]. Russell et al. bombarded bean stem tips with exogenous DNA covered with Tungsten particles and obtained transgenic plants [38]. Nowadays, there are few reports about foreign gene insertion into cotton. El-Shemy et al. transferred the V3-1 gene into soybean, which increased the content of globulin and improved its nutritional quality [40]. Jun Wu et al. developed a modified particle bombardment method for plant transgenesis incorporating foreign DNA into the plant genome with an increased transformation frequency, as compared to the conventional particle bombardment method [41]. The research on the related endogenous gene of salt
tolerance from cotton is emerging. Some genes, including \textit{GhAKT1} and \textit{GhWRKY3}, were cloned and inserted into onion for transient expression studies\cite{42, 43}, while similar research in cotton has not been reported. The germination ability of \textit{CA} transgenic seeds under salt stress is improved, which leads to the conclusion that \textit{CA} is a plant salt-tolerance gene. In recent years, the inclusion of foreign genes into tobacco, tomato, wheat, rice, corn, soybean, and cotton has been studied extensively\cite{44-50}. The transgenic plants were mainly acquired by \textit{Agrobacterium tumefaciens}-mediated transformation and bombardment of embryo or callus via a desktop gene gun. In the process of \textit{Agrobacterium}–mediated in planta transformation, the genotype is generally found to influence T-DNA transfer and transgene expression in Arabidopsis and grapevine\cite{51-54}. Studies have reported in other plants species, the choice of the \textit{Agrobacterium} strain appeared critical and specific because of specific host-bacteria inactions\cite{52, 55}. In our work, we cloned a exogenous gene \textit{CA} from \textit{Saccharina japonica}, which was an extreme salt-tolerant algae, enlarging our range of experiment materials.

In our investigation, firstly, we have obtained 3 cotton varieties with weak green fluorescent spontaneously from many upland cottons and particle bombardment in vivo was applied for the transformation of cotton pollens. Second, we established a system of transient expression of cotton pollens via the particle bombardment technology, which is characterized by easier manipulation, lower pollution, and higher survival rates compared with the pollen tube injection, and provided theoretical evidence and technical support for new methods of transgenesis and creation of salt-tolerant materials.

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\section*{Conflict of interest}
Wuwei Ye, Jingjing Kong and Xuke Lu conceived and designed the research. Jingjing Kong and Xuke Lu conducted the experiments. Xiaojie Zhao, Zujun Yin, and Shuai Wang analyzed the data. Jingjing Kong and Xuke Lu wrote the manuscript. Delong Wang, Junjuan Wang and Weili Fan edited the manuscript. Jingjing Kong and Xuke Lu contributed equally to this work. All of the authors declare that there is no conflict of interest.

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