EFFECTS OF GENIPOSIDE ON THE REGULATION MECHANISMS OF PHOTOSYNTHETIC PHYSIOLOGY, ROOT OSMOSIS AND HORMONE LEVELS IN MAIZE UNDER SALINE-ALKALI STRESS

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Abstract. The aim of the study was to monitor the effects of exogenous geniposide (GD) on maize (Zea mays L.) and explore the mechanisms through which it mitigates saline-alkali stress to provide a theoretical basis for revealing the growth physiology and chemical regulation processes in maize. The control (CK) received nutrient solution culture while the group under saline-alkali stress (SAS) received 150mmol L−1 saline-alkali solution (molar concentration ratio NaCl:NaSO3:NaHCO3:Na2CO3=1:9:9:1). ‘Jilong2’ (saline-alkali tolerant) and ‘Xinxuan58’ (saline-alkali sensitive) were chosen as experimental materials and were cultivated with hydroponic method. Maize received the saline-alkali solution 12 hours after GD treatment and saline-alkali solution was administered 3 times, every 12 hours. The GD and saline-alkali solution group constituted the S+G treatment which promoted maize and a significant increase was observed in fresh and dry weights, plant height and photosynthetic enzymes in the leaf along with a significant decrease in the root to shoot ratio compared with saline-alkali treatment. The indoleacetic acid (IAA), zeatin-riboside (ZR), gibberellin (GA), superoxide radical (O2−) generation rate, malonaldehyde (MDA) and hydrogen peroxide (H2O2) levels in roots were significantly lower than in the S+G treatment under saline-alkali stress, while the level of abscisic acid (ABA) and the activity of antioxidative enzymes increased.

Keywords: maize seedlings, photosynthetic enzyme, osmotic substance, antioxidant enzyme, hormone

Introduction

Enrichment of soil soluble saline-alkali can change soil physical and chemical properties, even brings about land salinization. The salinization and alkalization in soil often occurs together in natural conditions. High concentrations of salts and alkali causes injury to plants not only with ionic, osmotic and oxidative stresses to plants, but also through high...
pH leading to a block in the transportation of substances required for nutrition (Mukhtar et al., 2019). Salinization is a severe trouble that threatens agricultural development worldwide, the area of saline-alkali soil is increasing year by year, most of which are not exploited and utilized (Zhao et al., 2020). The total area of saline-alkaline soils is $9.5 \times 10^8$ ha$^{-2}$ worldwide. Especially in China, saline-alkaline soils were widely distributed in several provinces of Northwest China, Northeast China, North China and Central China. Saline-alkaline land with agricultural development potentially occupies over 10% of the total cultivated area of China. Songnen Plain is one of the most widely distributed area in saline-alkali soils all over the world and it is located in the northeast China. The saline-alkali area is estimated at 3.73 million hectares and was still increasing at a speed of 1.4% per year (Litalien et al., 2020).

Maize is considered to be particularly sensitive to saline-alkali stress during seedling stage. Therefore, revealing the adaptive and protective mechanisms of maize to saline-alkali stress is of great theoretical significance and practical value. Assimilation of inorganic carbon by photosynthesis is the main source of CO$_2$ organic matter in plant assimilation, plant growth and biomass accumulation (Huang et al., 2019). The primary pathway of photosynthetic carbon fixation in higher plants contains three kinds that is C$_3$ pathway, C$_4$ pathway and Crassulacean acid metabolism (CAM) pathway. C$_4$ pathway is the physiological basis of C$_4$ crops such as maize of high photosynthetic efficiency, biomass potential and crop yield formation under saline-alkali stress (Hughes et al., 2019).

The main physiological function of PEPCase in C$_4$ plant photosynthesis is to use PEP as substrate to fix CO$_2$ and produce oxaloacetatate, and oxaloacetate(OAA) is catalysed to malic acid by MADP-ME. Malate is metabolized by NADP-MDH to produce CO$_2$ and pyruvate, PEP use pyruvate as a substrate to be produced by PPDK catalysis and returned to the C$_4$ cycle. Studies have shown that plants convert carbon dioxide via C$_4$-pathway and Calvin cycle immobilization under the influence of the activity of related metabolic photosynthetic enzymes, such as phosphoenolpyruvate carboxylase (PEPcase), pyruvate dkinase (PPDK), NADP-malatease (NADP-ME), malate dehydrogenase (NADP-MDH) and ribon glycol diphosphate carboxylase (RuBPCase) (Lei et al., 2017). This indicated that saline-alkali stress slowed down the CO$_2$ assimilation rate in crops, resulting in decreased enzyme activity, which further disrupted the balance of consumption and accumulation of metabolites in the C$_4$ pathway and Calvin cycle, and reduces the content of assimilates transported to roots, which affect root growth (Zhao et al., 2019). The production of reactive oxygen species is affected by saline-alkali stress, which affect hormone metabolism and distribution, and destroys hormone balance. There are synergistic or antagonistic effects between endogenous hormones. Hormones can be used as an endogenous signal in response to abiotic stress. Plant hormones can be signals involved in physiological functions and metabolism. IAA play a major role in root growth of maize seedlings, GA induces cell elongation, ZR is an important form of CTK transport in xylem, and ABA is a growth inhibitor to slow seedling growth under saline-alkali stress. The production of reactive oxygen species (ROS) was led by salt-alkali stress, which in turn affects hormone metabolism and distribution, and destroys hormone balance (Li et al., 2017). The accumulation of soluble sugar and proline can effectively maintain the normal metabolism of plant roots because of reducing the osmotic potential of plant cells, eliminating free radicals and ROS, thus relieving oxidative stress (Amira et al., 2020). At present, some progress has been made in the application of physiological control measures of exogenous substances in agricultural production, such as amendments (Wang et al., 2016), trehalose (Rohman et al., 2019), abscisic acid (Lu et al., 2019), humic acid (Kaya et al., 2018) and...
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salicylic acid (Taha et al., 2019), which can effectively alleviate the injury of crops under saline-alkali stress. GD is a class of cyclic ether terpenoids (iridoids) that exhibit a wide variety of biological activities and are widely distributed in the plant kingdom. GD content in gardenia is about 5%. As a natural extract, GD is non-toxic and has high biological activity, soluble in water, safe and convenient to use, mainly includes Gentiopicroside, Vanilloylicatalpol and geniposide. GD has wide-ranging beneficial applications in the avoidance of neuroprotection and anti-diabetes as well as GD has the effect of protecting the liver and the gallbladder (Wu et al., 2019). In agricultural production, GD also plays an important role in the growth of crops, and the compound product-enhancing agent prepared by GD can effectively increase the yield of cucumber, cowpea, wheat and cotton (Zhang et al., 1998a,b; Zhang et al., 1999a,b). It can also promote the growth and development of maize seedlings, improve the chlorophyll content and chlorophyll light energy conversion efficiency in the leaves of green vegetables, promote the accumulation of soluble sugar and soluble protein, and enhance the photosynthetic ability of plant leaves. GD could promote root growth, thus increasing plant biomass under saline-alkali stress.

At present, effects of GD on growth and development of maize seedlings under saline-alkali stress have not been reported. Effects of exogenous GD on growth morphology, seedling biomass, leaf photosynthetic enzyme activity, root osmotic regulation, antioxidant system and endogenous hormones under saline-alkali stress of different saline-alkali tolerant maize seedlings were investigated.

In this study, the effects of exogenous GD on growth morphology, seedling biomass, leaf photozyme activity, root osmotic regulatory substance content, antioxidant system and endogenous hormones of saline-alkali resistant maize seedlings under saline-alkali stress were investigated with two varieties as test materials, and it provides theoretical and experimental basis for the application of exogenous GD in maize field stress tolerance production.

Materials and methods

Study design and experimental procedures

In the present study, we choose maize (Zea mays L.) Jilong2 and Xinxuan58 cultures to be transplanted into nutrient solution after seedling stage. The maize cultures were provided by Heilongjiang julong seed industry co., LTD and Shenyang xinxuan agricultural science and technology co., Ltd. The concentrations and treatment duration of the agents used in this study were determined after preliminary experiments. GD was provided by Guangxi Shan Yun Biochemistry Technology Co., Ltd, the treatment concentration was 5 mg L⁻¹, and its molecular weight was 388.37, and the purity was > 98%. The saline-alkali solution was prepared, in which the major constituents were NaCl: Na₂SO₄: NaHCO₃: Na₂CO₃ with a molar ratio of 1:9:9:9:1. Light absorbance was measured by a UV-visible spectrophotometer (UV-5500, manufactured by Shanghai Metash Instruments Co., Ltd.).

A hydroponic experiment was performed under different treatment conditions in the maize cultivation physiology laboratory of the Agricultural College of Northeast Agricultural University in June 2019. Maize seeds were surface-sterilized by rinsing them for 2 min in a 10% NaClO solution, then the seeds were rinsed with distilled water for several times, and soaked in distilled water for 8 h. Germination tests were carried out in an incubator at 25±1°C with 75% humidity. The uniformly germinated seeds were selected after 48 h incubation and cultivated in a test pot (60 cm in length, 30 cm in width, 12 cm in high) containing 150 L 1/2 Hoagland nutrient solution (pH 6.0±0.1) with 25 seedlings in
each pot with nutrient solution supply (CK), saline-alkali solution supply (SAS), saline-alkali solution and GD supply (S+G) or GD solution supply (GD). The cultures were maintained in standard conditions of day room temperature (28±1°C), night room temperature (25±1°C) with 65%−75% humidity, on a 12-h light and 12-h dark cycle with a light intensity of 4000 lx. The nutrient solution was changed in every 3 days, and was supplied with oxygen by air pump every day. When the plants had 3 true leaves, the seedlings were divided into four groups before further treatment. Fifty seedlings were used for each set of experiments. The seedlings were randomly assigned to one of the four groups: control group (CK), saline-alkali group (SAS), GD and saline-alkali group (S+G) or GD group (GD). The CK treatment received 1/2 Hoagland nutrient solution. The SAS treatment received 1/2 Hoagland nutrient solution and 150 mmol L⁻¹ saline-alkali solution. The S+G treatment received 1/2 Hoagland nutrient solution, 150 mmol L⁻¹ saline-alkali solution and 5 mg L⁻¹ GD. The GD treatment received 1/2 Hoagland nutrient solution and 5 mg L⁻¹ GD. To avoid the impact of saline-alkali stress on maize seedlings, saline-alkali solution was added to the maize 12 hours after GD treatment and saline-alkali solution was administered 3 times, every 12 hours. Saline-alkali solution reaching 150 mmol L⁻¹ marked Day 0. Seedlings were sampled at 4th d of germination for biochemical and physiological measurements. All experiments were repeated a minimum of 3 times.

**Agronomic characters of maize seedlings**

Three plants of each treatment were harvested on the 4th day. Plants were rinsed with distilled water and, after removing the excess water with filter paper the fresh and dry weight of the seedlings was measured by a balance. After the plant samples were heated for 30 min at 105°C to halt metabolism, and dried at 80°C to constant weight for assays of dry weight and the average value was taken as the final result, the fresh and dry weight has units of g 3plant⁻¹ (take the total weight of three seedlings as the unit). Height was measured from the base of the seedlings to its highest point and the average value was taken as the final result, the plant height has units of cm. Root dry weight was divided by stem dry weight to calculate the root-shoot ratio (R/S).

**Measuring photosynthetic activity of maize leaves**

Photosynthesis was measured during the study at day 4. The third true leaves from three plants of maize seedlings were cut for measurements. The enzyme extraction and activity of PEPCase and RuBPCase was performed according to a modified method of Sayre (1979). The concentration of the standard substance and the OD value at 490 nm of each sample were determined sequentially on the enzyme-linked immunoassay spectrophotometer. RuBPCase determination: 0.1 ml of the enzyme solution was mixed with 0.3 ml 0.1 mol L⁻¹ Tris-HCl (pH 8.0), 0.3 ml 0.1 mol L⁻¹ MgCl₂, 0.3 ml 50 mmol L⁻¹ adenine nucleoside triphosphate (ATP), 0.3 ml 50 mmol L⁻¹ dithiothreitol (DTT), 0.3 ml 2.0 mmol L⁻¹ NADH, 0.3 ml 1.0 mmol L⁻¹ EDTA-Na, 0.1 ml 200 μmol L⁻¹ NaHCO₃, 1.0 ml distilled water and 0.1 ml 3-phosphoglycerate kinase (PGK)/ 3-glyceraldehyde phosphate dehydrogenase (GAP-DH) (15 u/15 u) buffer preheated in 30 constant temperature water bath for 10 min. The absorbance E₀ of the samples was measured at a wavelength of 340 nm with distilled water as a reference wavelength. Then 0.1 ml 9 mmol L⁻¹ 1,5-ribose diphosphate (RuBP) was added to initiate the reaction, the changes in light absorption was determined in 15 s intervals. PEPCase determination: the enzyme solution 1 ml was mixed with 1 ml 0.1 mol L⁻¹ H₂SO₄ (pH 9.2), 0.1 ml 10 mmol L⁻¹ MgCl₂, 0.1 ml 10 mmol L⁻¹ NaHCO₃, 0.2 ml 40 mmol L⁻¹ phosphoenolpyruvate (PEP), 0.3 ml 1 mg ml⁻¹ NADH, 0.2 ml malic dehydrogenase (MDH)
then pre-heated by water bath at 28°C for 10 min, the reaction was initiated with 0.2 ml PEPC extract, and the decrease of sample absorbance was measured rapidly at 340 nm. To determine the activity of NADP-MDH, NDAP-ME and PPDK a kit was used according to the operating procedures of the instructions. At least three replicates for each assay were performed. The average of test result was calculated and analyzed. Experiments were repeated thrice and three repeats were performed each time.

**Measuring endogenous hormone content of maize roots**

Endogenous hormone content was measured during the study at day 4. The contents of IAA, ZR, GA and ABA in maize root were analyzed using an enzyme-linked immunosorbent assay. 0.5 g tissue samples (maize leaves) were ground in 2 ml sample extract under ice bath to form homogenate, and samples were transferred into the test tube and extracted under 4°C for 4 h, then centrifuged at 3500 r/min for 8 min at 4°C, and the supernatants were collected into a 10 ml centrifuge tube. 1 ml of extract was added in the precipitate and laid still at 4°C for 1 h, then centrifuge, the supernatant was taken merge with the previous supernatant and record the volume, the residue is discarded. Use a C-18 solid phase extraction column to purity the supernatant. After passing through the column, the sample was transferred into a 5 ml plastic centrifuge tube. The methanol in the extract was removed by vacuum, concentrated and dried or blown dry with nitrogen and the volume was fixed with the sample diluent. After extracting the hormone, the maximum concentration of the ABA is 500 ng ml⁻¹, IAA and ZR is 100 ng ml⁻¹, GA is 10 ng ml⁻¹, and then diluted 2 times, 4 times, 8 times, 16 times, 32 times, 64 times until there are 8 concentrations (including 0 ng/ml). The series of standard samples were added to the first two rows of a 96-well enzyme standard plate, add samples to be tested to the remaining holes, and add the same sample every two holes, 50 μl per hole. A certain amount of anti-mixing was added to the 5 ml sample diluent, then the enzyme label plate was put into the wet box to start the competition. The competitive conditions were 37°C and 0.5 h. Washing boards: get rid of the reaction fluid on the board and pat it clean on paper. Get rid of it immediately after adding washing liquid to the board, repeat four times. Appropriate enzymatic II antibodies were added in 10 ml sample diluents, each hole was100 μl, then the enzyme labeling board was incubated in wet box at 37°C for 0.5 h. Wash again. 10 mg o-phenylenediamine (OPD) was completely dissolved in 10 ml substrate buffer and mixed with 4 μl 30% H₂O₂, each hole was 100 μl, and the enzyme label plate was placed into the wet box until appropriate color development, then 50 μl 2 mol L⁻¹ sulfuric acid was added to each hole to terminate the reaction. The concentration of the standard substance and the OD value at 490 nm of each sample were determined sequentially on the enzyme-linked immunoenzyme spectrophotometer. Experiments were repeated thrice and three repeats were performed each time.

**Osmotic content in maize roots measurement**

Osmotic content was measured during the study at day 4. The anthrone colorimetric method was used to determine the soluble sugar content in maize roots. The standard curve was established using sucrose. The 0.3 g root was cut into pieces and put into the scale test tube together with 5 ml distilled water and sealed with plastic film. Samples were extracted in boiling water for 30 min the extraction was performed two times. The extract was filtered into 25 ml volumetric flasks, the test tube was washed with distilled water, and the residues were transferred into volumetric flasks and diluted to 25 ml. 0.5 ml extract was transferred by 1.5 ml of distilled water to a 20 ml scale tube, 0.5 mL anthrone and ethyl acetate were
added into the mixture. The samples were placed in a boiling water bath for 1 min then cooled to room temperature. The absorbance of the samples was measured at a wavelength of 630 nm with blank space as a reference wavelength. Soluble sugar content was calculated with the following formula:

\[ \text{Soluble sugar content [mg g}^{-1}] = \frac{(2500 \times C)}{0.3 \, \text{g}}, \]

where \( C \) is soluble sugar content of determination solution on standard curve. Soluble sugar content. Proline content in maize root was determined according to ninhydrin coloring method. Before sample measurement, standard curves were made with different concentrations of 2 ml standard solutions of proline. 0.5 g of fresh leaves and 5 ml 3% sulfosalicylic acid were added to the test tube and then transferred to boiling water for 10 min. After cooling, the extract was filtered. After that, 2 ml filtrated extraction mixed with a 2 ml ninhydrin reagent and 2 ml glacial acetic acid. The mixture was heated in a boiling water bath for 30 min. 4 ml of toluene was added after cooling and shaken for 30 s and the supernatants were collected into a 10 ml centrifuge tube. The supernatant was poured into a colorimetric cuvette and water was used as blank control. The absorbance values were read at 520 nm. Proline content was calculated with the following formula:

\[ \text{Proline content [μg g}^{-1}] = 2500 \times C, \]

where \( C \) is 2 ml proline content of determination solution on standard curve.

**Determination of superoxide anion production rate and hydrogen peroxide content in maize root system and antioxidant enzyme system**

Photosynthesis was measured during the study at day 4. \( \text{O}_2^- \) generation rate was measured using a hydroxylamine oxidization method. For determining \( \text{O}_2^- \) generation rate, 5 g tissue samples (maize root-tip) were ground in 6 ml of 65 mmol L\(^{-1}\) phosphate buffer solution (PBS) (pH 7.8) into a homogenate, after filtration through four layers of gauze, centrifuged at 5000 r/min for 10 min at 4°C. Then, 1 ml (about 0.5 mg protein) supernatant was mixed with 0.9 ml PBS and 0.1 ml 10 mmol L\(^{-1}\) hydroxylamine chloride for reaction for 20 min at 25°C. Supernatant (0.5 mL) was mixed with 0.5 ml 7 mmol L\(^{-1}\) 3-aminophenylsulfonic acid and 0.5 ml 7 mmol L\(^{-1}\) \( \alpha \)-naphthylamine, and incubated for 20 min at 25°C. After the reaction, the same volume of n-butanol was added and shaken vigorously, and laid still for layer separation. The intensity of the chromogenic agent in the butanol layer was determined at 530 nm and sample solution was replaced with phosphate buffer. Both supernatants were combined and centrifuged for 5 min at 1500 \( \times \) g and absorbance was measured at 530 nm.

The H\(_2\)O\(_2\) content was determined spectrophotometrically. First, 2 g of fresh sample was weighed, precooled at 4°C, acetone with quartz sand was added at a ratio of 1:1, and the homogenate was ground. Then, it was centrifuged at 3000 rpm for 10 min and the supernatant (\( V_s \)) was taken. Finally, 5% titanium sulfate and ammonia water were added to the supernatant. When the solution became turbid, it was centrifuged again 5000 rpm for 10 min. After discarding the clear supernatant, the precipitates were washed with acetone to remove plant pigments. Subsequently, 5 mL of concentrated sulfuric acid was added to the precipitates. When the precipitate dissolved, the sample volume was made up to 10 ml, the absorbance was measured at 415 nm. H\(_2\)O\(_2\) content was calculated with the following formula:
H₂O₂ content [nmol⁻¹ FW] = (C × V_t)/(2 g × 1 ml),

where C is the H₂O₂ content from the standard curve, V_t: total volume of extracting enzyme solution.

The crude enzyme extraction for SOD (Superoxide dismutase), POD (Peroxidase), CAT (catalase), MDA was similar: 4 ml 0.1 mol L⁻¹ PBS (pH 7) was added to 2 g of each sample to crush in precooled mortar and pestle. The final volume was made up to 200 ml. The 1.5 ml homogenate was centrifuged at 10,000 rpm at 4°C for 15 min, and the crude enzyme extract was the supernatant. SOD activity was assayed by measuring the ability of inhibit photochemical reduction of NBT. 0.6 ml 50 mmol L⁻¹ PBS, 0.12 ml 130 mmol L⁻¹ Met, 0.12 ml 750 μmol L⁻¹ NBT, 0.12 ml 100 μmol L⁻¹ EDTA-Na₂, 0.12 ml 20 μmol L⁻¹ riboflavin, 0.02 mL of an enzyme solution (sections applied with PBS instead of enzyme solution were studied as the control) were mixed with 0.1 mL of distilled water in a 10 mL centrifuge tube and mixed gently. One control centrifuge tube was placed in the dark and the other samples reacted at 25°C under 4000 lx sunlight for 20 min (each tube was light consistent). One unit of SOD activity was defined as the quantity of enzyme required for 50% inhibition of NBT reduction at 560 nm. POD activity was estimated using the guaiacol method. The reaction mixture contained 0.05 ml of enzyme extract, 2.2 ml PBS, 0.15 ml of 1% H₂O₂ and 0.6 mL 1% guaiacol solution and were warmed for 5 min at 37°C in a water bath. Reaction buffer without crude enzyme solution was used as the control. The change in absorbance at 470 nm was measured over a period of 3 min at 1 min intervals. We determined enzyme activity by measuring the change in absorbance per minute, and one unit (U) of enzyme activity was defined as the amount of the enzyme that changes the absorbance by 0.01 per minute. CAT activity was assayed using hydrogen peroxide as substrate. A 0.1 ml enzyme sample and 2.865 mL PBS were added to a test tube, mixed, and placed in a boiling water bath for 10 min. All the reaction mixture was incubated at 25°C for 5 min in the water bath. Then 0.035 ml 3% H₂O₂ was added and mixed well, the tube was quickly poured into the colorimetric cup and start to calculate the time. The colorimetric reaction was read at 450 nm, each record lasted 3 min and was set at 1 min intervals, and enzyme volume of 0.01 absorbance per minute was regarded as one unit (U) of enzyme activity. The content of MDA was determined by thiobarbituric acid (TBA) reaction chronometry. Two milliliter of the enzyme solution was mixed with 2 ml of 0.6% TBA, and the mixture was placed in a boiling water bath for 10 min. The absorbance of the resulting supernatant was measured respectively at the wavelength of 532, 600, and 450 nm after cooling and centrifuge.

For SOD activity determination the absorbance values of the control tube and the measuring tube were measured by a spectrophotometer at 560 nm to be A0 and A, respectively. Percentage of inhibition was calculated as P [%] = (A0 − A)/A0 × 100, and SOD activity as [U g⁻¹] = 114 × P/(1 − P). For POD activity determination the absorbance value A1 at 1 min and the absorbance value A2 at 2 min at 470 nm were measured with a spectrophotometer. POD activity was calculated as [U g⁻¹] = 20,000 ΔA, where ΔA = A2 − A1. CAT activity was calculated as [U g⁻¹] = Δ240×V/Va/W. For MDA content determination the absorbance value D₆₀₀ at 600 nm, value D₅₃₂ at 532 nm and value D₄₅₀ at 450 nm were measured with a spectrophotometer. MDA content was calculated as (nmol g⁻¹ FW) = (6.45 × (D₃₅₂−D₆₀₀)−0.56D₄₅₀) × 0.015/W. Experiments were repeated thrice and three repeats were performed each time.
Data calculation and analysis

The experimental data were statistically analyzed by Excel2010 software, variance analysis by SPSS17.0 software, and difference significance analysis by LSD method (Wu et al., 2014).

Results

Effect of GD on maize growth phenotype under saline-alkali stress

As shown in Table 1, a significant growth inhibition in maize seedlings has been caused by saline-alkali stress. Compared with the seedlings under the CK treatment, the seedlings under the SAS treatment had short plants and significantly decreased dry and fresh weight and root-shoot ratio increased. And compared with the seedlings under the SAS treatment, the seedlings under the S+G treatment had long plants and significantly increased dry and fresh weight and root-shoot ratio decreased. This shows that exogenous GD had a better alleviating effect on dry-fresh weight and height of maize seedlings. Especially the GD treatment can have significant promoting effect on the shoot and the change of Xinxuan58 root whitening, and the area of yellowing in Jilong2 was larger than that of Jilong2. It was observed that GD treatment greatly promoted dry-fresh weight, plants height and the growth of both maize cultivars compared with the CK. Seedling growth of Jilong2 and Xinxuan58 was significantly inhibited by saline-alkali stress (Fig. 1). Compared with the seedlings under the CK treatment, the seedlings under the SAS treatment had smaller leaves and roots, significantly decreased height and seedlings under the SAS treatment were wilting and some mature leaves showed chlorosis, the fibrous roots reduced, and seedlings growth was relatively inhibited. In both maize cultivars leaves of ecotype yellowed, and the area of yellowing in Jilong2 was larger than that in the Xinxuan58. The S+G treatment delayed leaf wilting in maize seedlings subjected to saline-alkali and fibrous roots increased. GD can effectively promote seedling growth, increase leaf width, promote root whitening, and increase fibrous roots.

Table 1. Effects of geniposide on the dry and fresh weight, plant height and root-shoot ratio of maize under saline-alkali stress on the fourth days

| Variety | Treatment | Fresh weight (g 3^1 plant) | Dry weight (g 3^1 plant) | Plant height (cm) | Root-shoot ratio |
|---------|-----------|----------------------------|--------------------------|------------------|-----------------|
|         |           | above ground | underground | above ground | underground |                |                |
| Jilong2 | CK        | 4.81±0.37^b | 2.59±0.13^b | 0.31±0.074^b | 0.22±0.039^b | 52.40±2.48^b | 0.54±0.04^b |
|         | SAS       | 1.63±0.18^a | 1.95±0.06^c | 0.20±0.059^b | 0.21±0.054^b | 41.90±1.93^d | 1.21±0.17^a |
|         | S+G       | 2.44±0.15^c | 2.55±0.13^c | 0.34±0.058^a | 0.37±0.032^b | 48.40±1.40^c | 1.04±0.02^b |
|         | GD        | 5.89±0.19^a | 2.95±0.12^a | 0.37±0.036^a | 0.22±0.042^a | 71.60±1.70^a | 0.50±0.04^b |
| Xinxuan58 | CK       | 2.70±0.17^b | 1.89±0.12^b | 0.18±0.060^c | 0.41±0.061^a | 51.80±1.69^a | 0.70±0.08^b |
|         | SAS       | 0.66±0.13^c | 1.50±0.13^d | 0.11±0.038^c | 0.31±0.042^a | 43.50±1.94^b | 2.19±0.48^a |
|         | S+D       | 2.59±0.25^b | 2.34±0.31^d | 0.28±0.066^b | 0.20±0.044^b | 54.50±2.12^a | 0.92±0.20^b |
|         | GD        | 4.96±0.12^a | 2.64±0.21^d | 0.39±0.020^a | 0.21±0.056^b | 54.40±1.91^a | 0.53±0.04^b |

Different letters within the same variety and column indicate significant difference at 0.05 level. The same as below
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Figure 1. Effect of GD on growth morphology of maize under saline-alkali stress on the fourth day

Effect of GD on photosynthetic enzyme activity of maize leaves under saline-alkali stress

The main physiological function of PEPCase in C_4 plant photosynthesis is to use PEP as a substrate to fix CO_2 and produce oxaloacetate, and oxaloacetate(OAA) is catalysed to malic acid by MADP-ME. Malate is metabolized by NADP-MDH to produce CO_2 and pyruvate, PEP use pyruvate as a substrate to be produced by PPDK catalysis and returned to the C_4 cycle. An important carbon fixation in photosynthesis is the Calvin cycle. CO_2 reacts with RuBP under RuBPCase catalysis in the Calvin cycle. Thus, photosynthetic enzyme reflects the photosynthetic activity of maize seedlings under saline-alkali stress (Fig. 2). Compared with the seedlings under the CK treatment, in the seedlings under the SAS treatment the activity of PEPCase, RuBPCase, NADP-ME, NADP-MDH and PPDK significantly decreased, the change of Xinxuan58 was greater than that of Jilong2. Compared with SAS treatment, there was no significant change in NADP-ME activity in Jilong2 under S+G treatment, while NADP-ME activity in Xinxuan58 increased. Compared with SAS treatment, in the seedlings under the S+G treatment the activity of NADP-MDH, PPDK and RuBPCase significantly increased. This suggests that saline-alkali stress inhibits the photosynthetic pathway and GD treatment resulted in increased C_4 photosynthesis activity.

Effect of GD on endogenous hormone content and its ratio in maize roots under saline-alkali stress

The plant adaptations to saline-alkali stress is reflected in changes of endogenous hormones and their ratios in maize under saline-alkali stress. Compared with CK, SAS treatment significantly inhibited the content of IAA in Jilong2, the relative content of endogenous IAA in Xinxuan58 was significantly increased and reached a notably high level (Table 2). The IAA concentration in Jilong2 and Xinxuan58 significantly increased under S+G treatment compared to the SAS treatments, respectively. Compared with CK, SAS treatment significantly inhibited the content of GA and ZR, the change of GA in Xinxuan58 was greater than that in Jilong2. After S+G treatment, the content of GA and ZR in roots of Jilong2 and Xinxuan58 was significantly greater than that in the SAS treatment. Compared with CK, SAS treatment significantly
increased the content of ABA in Jilong2, and in Xinxuan58, ABA content decreased. The ABA content in S+G treatment decreased relative to SAS treatment. Saline-alkali stress reduced IAA, GA and ZR contents in root of maize and inhibited plant growth. Under saline-alkali stress, increased levels of ABA may help to improve saline-alkali tolerance of plants. The accumulation of ABA content is conducive to plants to adaptation to stresses, and GD treatment can effectively alleviate the harm of saline-alkali stress and promote hormone balance recovery.

Figure 2. Effect of GD on photosynthetic enzyme activity in leaves of maize under saline-alkali stress on the fourth day. Different letters within the same variety and column indicate significant difference at 0.05 level

Hormone interaction in plants shows synergism or antagonism, the ratio of IAA/ABA, ZR/ABA and GA/ABA indicate different hormone impact on each other. The IAA/ABA and GA/ABA ratio in Jilong2 in SAS treatments decreased relative to CK, that in Xinxuan58 increased. And ZR/ABA ratio has no significant change. Compared with SAS, S+G treatment the ratio of IAA/ABA, ZR/ABA and GA/ABA significantly increased in Jilong2 and Xinxuan58. The ratio of IAA/ABA, ZR/ABA and GA/ABA under SAS treatment was greater, indicating that growth-promoting hormone inhibits ABA action. Plants regulate energy distribution through hormones to support growth and development under stress. GD can change the hormone content under saline-alkali stress and keep the relative balance of hormone content in plants.
Table 2. Effect of GD on endogenous hormones and hormones ratio in roots of maize under saline-alkali stress on the fourth day.

| Variety   | Treatment | IAA (ng g⁻¹) | GA (ng g⁻¹) | ZR (ng g⁻¹) | ABA (ng g⁻¹) | IAA/ABA | GA/ABA | ZR/ABA |
|-----------|-----------|--------------|-------------|-------------|--------------|---------|--------|--------|
| Jilong2   | CK        | 253.15±      | 376.47±     | 125.44±     | 146.21±      | 1.73±   | 0.89±  | 2.58±  |
|           | SAS       | 196.98±      | 364.16±     | 116.26±     | 155.48±      | 1.27±   | 0.75±  | 2.34±  |
|           | S+G       | 249.37±      | 395.31±     | 148.29±     | 106.43±      | 2.34±   | 1.39±  | 3.71±  |
|           | GD        | 274.07±      | 372.58±     | 146.69±     | 128.70±      | 2.13±   | 1.14±  | 2.90±  |
| Xinxuan58 | CK        | 189.37±      | 409.56±     | 160.86±     | 161.24±      | 1.17±   | 1.00±  | 2.54±  |
|           | SAS       | 244.33±      | 385.94±     | 142.38±     | 144.48±      | 1.70±   | 0.99±  | 2.69±  |
|           | S+G       | 269.25±      | 407.80±     | 159.03±     | 123.77±      | 2.18±   | 1.29±  | 3.30±  |
|           | GD        | 250.81±      | 453.18±     | 202.28±     | 101.76±      | 2.47±   | 1.99±  | 4.46±  |

Different letters within the same variety and column indicate significant difference at 0.05 level. The same as below.

Effects of GD on antioxidant system and osmotic regulation of maize roots under saline-alkali stress

In order to maintain the normal physiological function of the cells, the water potential of the cells was reduced by the accumulation of proline and soluble sugar under saline-alkali stress. The proline content in SAS treatment increased relative to CK, the change of GA in Xinxuan58 was greater than that in Jilong2 (Fig. 3). And S+G treatment could further increase proline content. Compared with SAS treatment, soluble sugar content in maize cultivars under S+G treatment significantly increased. This indicates that the accumulation of proline and soluble sugar content can effectively alleviate the saline-alkali stress on the root of Jilong2, relatively, Xinxuan58 saline-alkali tolerance was weaker. GD treatment is beneficial to the accumulation of proline and soluble sugar content to maintain the normal function of root system under saline-alkali stress. The H₂O₂ contents and O₂⁻ generation rate in the root of Jilong2 and Xinxuan58 under SAS treatment increased compared with that of CK. The S+GD treatments had lower H₂O₂ contents and O₂⁻ generation rate than the CK. SOD, POD and CAT activity and MDA content under SAS treatment higher than that of CK. Under SAS treatment, the POD and CAT activity of maize roots were consistent with the trend of SOD activity. And S+G treatment significantly increased SOD, POD and CAT activity and MDA content compared with SAS treatment. GD can significantly alleviate the damage caused by saline-alkali stress on two maize cultivars, the change of antioxidant capacity Jilong2 was greater than that of Xinxuan58.
Figure 3. Effect of GD on permeating substances, antioxidase activity and MDA content in roots of maize under saline-alkali stress on the fourth day. A: Proline content; B: Soluble sugar content; C: \( \text{H}_2\text{O}_2 \) content; D: \( \text{O}_2^{-} \) generation rate; E: SOD activity; F: POD activity; G: CAT activity; H: MDA content. Different letters within the same variety and column indicate significant difference at 0.05 level.
Discussion

The accumulation of biomass is the mitigation and adaptation of plants to saline-alkali stress, and the difference of accumulation reflects the resistance of different plants (Razieh et al., 2019). Maize seedling dry-fresh weight and plant height decreased during saline-alkali stress, root-shoot ratio increased. This indicates that the growth of maize seedlings was inhibited under saline-alkali stress, the photosynthesize produced was preferentially transferred to the root to maintain absorption of water and nutrients. Normal physiological function of root system is the basis of seedling growth. GD treatment markedly improved seedling growth under saline-alkali stress conditions, increased dry-fresh weight and plant height and decreased root-shoot ratio. Photosynthates in the growing period were normally allocated to each organs to increase the saline-alkali resistance of maize seedlings. Simple GD can promote root growth, increase plant dry-fresh weight and plant height, that can be used as growth promoters. Former study has shown that GD and its compound can promote dry-fresh weight of cucumber and cowpea (Zhang et al., 1998a). 25 mg L^{-1} GD could increase dry-fresh weight of radish, and improve the root-shoot ratio (Qian et al., 2016). It is proved that GD was propitious to increase plant biomass accumulation, and promote roots growth by promoting the transport of photosynthetic products to roots. Our conclusions are in line with these studies.

Photosynthesis is the physiological metabolism that provides energy for plants, photosynthetic efficiency is closely linked with biomass accumulation in plants. And photosynthetic efficiency is the foundation of plant growth. Photosynthetic enzyme activity can reflect the effect of saline-alkali stress on carbon assimilation pathway in maize plants. PEPCase play important roles in the regulation of responses to abiotic stresses and organic acid metabolism (Ueno et al., 2020). NADP - ME not only can catalyse reversible reactions of oxidative decarboxylation of malate, but also regulate pyruvate sources in plants through synergism between PEPCase and NADP-MDH (Borba et al., 2018). PEPCase has the function of maintaining the stability of cytoplasmic pH and osmotic potential. Research shows that NADP - ME activity of plants is increased by saline-alkali stress, and improve stress tolerance (Zhao et al., 2019). The present study shows that PEPC, PPDK, NADP-ME, NADP-MDH and RuBPCase activities were significantly decreased under saline-alkali stress compared to CK. Carbon assimilation rate in C_{4} photosynthesis was inhibited by saline-alkali stress, carbon assimilation rate reduced and assimilates translocation decreased (Kandoi et al., 2018; Dong et al., 2019). GD treatment significantly increased photosynthesis activity in maize leaves under saline-alkali stress, and GD could enhance the activity of photosynthesis to alleviate the inhibition of photosynthetic metabolism of maize seedlings under saline-alkali stress.

Plant endogenous hormone refers to the trace organic matter in plants, which can regulate plant physiological metabolism and directly or indirectly regulate plant growth. The level and distribution of endogenous hormones in plants follow the changes of external environment, so plants adapt to stress by regulating hormones. Regulation of hormone levels under abiotic stress can mediate physiological metabolic responses in plants and improve plant adaptability to adversity (Joshua et al., 2019). Studies have shown that saline-alkali stress may increase the level of reactive oxygen species (ROS), and ROS was capable of inhibiting the hormone synthesis, which leads to decreased hormone content (Jan et al., 2019). ABA is an endogenous signal to regulate stomatal closure to avoid water loss during saline-alkali stress treatment (Zhang et al., 2019b).
The IAA, ZR and GA contents in SAS treatment decreased relative to CK, and the ABA content was higher in Jilong2 and lower in Xinxuan58. There was different saline-alkali tolerance among cultivars. Because the energy distribution of physiological metabolism in plants to osmotic regulation and antioxidant regulation is suppressed by ABA (Li et al., 2017, 2020), Jilong2 can alleviate their own osmotic stress and antioxidant stress injury through induced ABA synthesis, and then improve maize resistance under stress. IAA/ABA, ZR/ABA and GA/ABA ratios directly reflect the relationship of endogenous hormones, and changes of endogenous hormone levels under saline-alkali stress affect the balance between hormones (Amany et al., 2019). According to the change of IAA/ABA, ZR/ABA and GA/ABA ratios under CK treatment and SAS treatment is explained ABA in Jilong2 is more sensitive to saline stress than IAA and GA, and ZR and ABA sensitivities are similar. Root system of Jilong2 by secreting ABA slows root growth and adaptation to saline-alkali stress. The response sensitivity of Xinxuan 58 to IAA was higher than that of Jilong 2, and ABA content decreased, so the tolerance was poor (Xie et al., 2019). This suggests that GD treatment effectively regulates endogenous hormone balance and promotes IAA, ZR and GA content. IAA, ZR and GA as ABA antagonists increased, which inhibited the production of ABA, and alleviated the inhibition of root growth, improved the saline-alkali tolerance of maize roots.

The saline-alkali stress significantly decreased water potential and osmotic pressure, but increased cell expansion pressure in seedlings, which resulted in osmotic stress. Osmotic stress caused direct damage to maize plants and affected various physiological activities in plants (Amany et al., 2019). Studies have shown that accumulation of proline and soluble sugar content can reduce the cell permeability potential in maize roots and play an important role in maintaining water absorption in plant roots under saline-alkali stress. This study showed that the content of proline and soluble sugar was significantly higher than that of control in Jilong2 roots and lower than that in Xinxuan 58 root under saline-alkali stress, saline-alkali tolerant varieties can maintain the osmotic balance of plant cells by accumulating osmotic substances. GD can effectively promote the content of osmotic substances in maize roots under saline-alkali stress and improve the capacity of osmotic adjustment, which plays an important role in maintaining the water balance in roots of maize. Former study shows that 100 mg L\(^{-1}\) GD significantly increased soluble sugar content in willow leaves, which was consistent with the conclusion of this study (Zhang et al., 2018, 2019a). Plant cells accumulate a large amount of reactive oxygen species under saline-alkali stress, which leads to membrane lipid peroxidation and destruction of cell membrane structure and function, which leads to dysregulation of intracellular metabolism. Membrane lipid peroxidation under saline-alkali stress is closely related to active oxygen accumulation, indicating a disruption in the structure and function of the cell membrane, that leads to disturbance of lipid metabolism (Ahmad et al., 2019). SOD, PDD and CAT are a system of antioxidant enzymes that scavenge ROS in plants, the change of enzyme activity showed the plant resistance (Li et al., 2020).

The main function of SOD is to clear O\(_2^−\) generate to form H\(_2\)O\(_2\), and then H\(_2\)O\(_2\) was decomposed into O\(_2\) and H\(_2\)O under POD and CAT catalysis. The antioxidant enzymes maintain a certain balance with ROS, the product of membrane lipid peroxidation is MDA that can reflect the degree of oxidative stress damage to plants (Jing et al., 2019). Compared with CK, O\(_2^−\) generation rate, H\(_2\)O\(_2\) content, MDA content, the activity of SOD, POD and CAT increased significantly under SAS treatment. During saline-alkali stress condition, the root of maize was affected by oxidative stress, the change of
antioxidant enzyme activity in Jilong2 was greater than that in Xinxuan58. ROS accumulation cause damages to membrane structure and the imbalance between pro-oxidant and anti-oxidant systems and then following oxidative stress (Liu et al., 2017). Compared with SAS treatment, \( O_2^- \) generation rate, \( H_2O_2 \) content, MDA content, the activity of SOD, POD and CAT decreased significantly under GD treatment. GD treatment alleviates the oxidative stress damage caused by saline-alkali stress on maize roots, and effectively improves the saline-alkali tolerance of maize, Xinxuan58 is more sensitive to GD effects. The increased activity of antioxidant enzymes can effectively scavenge ROS (Abdelgawad et al., 2016; Yao et al., 2016; Elrys et al., 2020). 25 mg L\(^{-1}\) GD significantly reduced SOD and POD activity in green vegetable leaves (Yan et al., 2016). 1 mg L\(^{-1}\) GD effectively reduced SOD activity in maize leaves (Qian et al., 2015). In this study the GD treatment could significantly improve the activity of SOD, POD and CAT, which appear to be at odds with that conclusion, due to different GD concentrations on antioxidant oxidase activity.

Conclusions

In conclusion, this study demonstrated that exogenous GD increased tolerance to saline-alkali stress in maize. Under saline-alkali stress, GD could reduce the harmful effects on the growth of maize by regulating osmotic metabolism, activating photosynthesis, enhancing the antioxidant system and increasing endogenous content. These results revealed a key role of GD in the relief of saline-alkali stress and indicated that it can be used as root-promoting agent. The physiological mechanism of GD on maize seedling growth promotion and saline-alkali stress tolerance is not clear. The next step is to determine the effect of GD on carbon and nitrogen metabolism of maize seedlings to determine the role of GD under saline-alkali stress.

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