Exogenous Melatonin Promotes the Salt Tolerance by Removing Active Oxygen and Maintaining Ion Balance in Wheat (*Triticum aestivum* L.)

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Melatonin (MT) is a small molecule indole hormone that plays an important role in the regulation of biological processes and abiotic stress resistance. Previous studies have confirmed that MT promotes the normal development of plants under stress by mediating physiological regulation mechanisms. However, the physiological mechanism of exogenous MT regulating seed germination and seedling growth of wheat under salt stress is still unclear. In this study, NaCl stress decreased germination rate and inhibited seedling growth of wheat, but shoot length, root length, and plant weight of SM15 did not change significantly. The addition of 300 μM MT in the cultivation solution directly promoted the germination rate of SM15 and ZM18, and lateral root production, but decreased the germination rate of JM22 and inhibited the length of germ and radicle of three varieties under salt stress. For wheat seedling, application of MT could increase proline content, soluble protein, soluble sugar, Ca²⁺ content, and vital amino acid content in leaves to keep high water content, low level of H₂O₂ content, and low [K⁺]/[Na⁺] ratio. MT increased root vigor and [K⁺]/[Na⁺] ratio and decreased H₂O₂ content in root induced by salt stress. In conclusion, MT enhanced salt tolerance in wheat seeds and seedlings by regulating the synthesis of soluble protein and sugar, ion compartmentation in roots and leaves, enhancement of enzymatic systems, and changes in amino acid levels. Salt resistance varied with different varieties under the same environmental condition. SM15 was a higher salt-resistant variety and JM22 was a salt-sensitive one. In wheat production, the application of exogenous MT should consider the differences among varieties of wheat during the sowing and seedling stages.

**Keywords:** salt stress, melatonin, wheat, germination, antioxidative activity
INTRODUCTION

Soil salinization, a geologically environmental problem all over the world, is one of the main factors resulting in the decrease in grain production (Liang et al., 2018). According to incomplete statistics, the salinization of arable land globally is increasing at a rate of 0.3–1.5 million hectares year⁻¹ (Harper et al., 2021). The total area of salinized soil in China is about 99.13 × 10⁶ hm², which accounts for 1.03% of the total Chinese land area (Yang and Wang, 2015). It is expected that 50% of arable land will be under saline-alkali stress by 2050 due to environmental pollution, lack of freshwater, improper irrigation methods, and other factors (Dou et al., 2019; Yang et al., 2021). Wheat (Triticum aestivum L.), one of the medium salt-tolerant crops and the second-largest food crop in the world, also faces soil salinization in its growing area. Salt stress leads to the significant inhibition of seed germination and seedling growth (Shah et al., 2015). Under salt stress, excessive reactive oxygen species (ROS) are accumulated which will cause cell damage, DNA damage, and oxidative stress to plants, consequently disrupting the physiological balance and leading to reduced photosynthesis and production yield (Zhang et al., 2016; Tripathi et al., 2020). High Na⁺ accumulation in plant cells leads to osmotic stresses and limits uptake and utilization of other nutrition ions, which causes moisture loss and electrolytic leaching due to cell membrane damage (Abbasi et al., 2015; Ahmed et al., 2015; Rehman et al., 2017).

Melatonin (MT), an amine hormone, has been confirmed extensively for enhancing plant tolerance to various abiotic stresses (Zhang et al., 2013; Turk et al., 2014; Ren et al., 2020). There are multiple mechanisms verified in MT helping plants to enhance abiotic stress tolerance (Huangfu et al., 2021; Sun et al., 2021). The application of MT has shown stimulating antioxidant enzymes and scavenging ROS in plants under stress conditions (Park et al., 2013; Li et al., 2018), preventing chlorophyll degradation and increasing photosynthetic efficiency, reducing ion toxicity (e.g., heavy metals, Na⁺, and so on) and increasing osmotic substances to maintain water status (Gao et al., 2016; Cai et al., 2017; Siddiqui et al., 2019a, 2020; Al-Huqail et al., 2020), and enhancing secondary metabolite biosynthesis and upregulating defense genes to decrease cell injury (Li H. et al., 2017; Zhao et al., 2017; Elsayed et al., 2021). Under salt stress, MT alleviates directly ROS burst and cell damage for bermudagrass (Shi et al., 2015). Also, MT alleviates the inhibitory effects of NaCl stress on germination of cucumber seeds mainly by increasing antioxidant enzymes, affecting phytohormone biosynthesis and catabolism, and regulating storage protein degradation to vital amino acids during seed germination (Zhang H. J. et al., 2014; Zhang et al., 2017). MT has been reported to extend longevity by significantly reducing chlorophyll degradation, delaying the leaf senescence, and preventing cell death in rice plant and tomato seedlings under salt stress (Siddiqui et al., 2019b; Zhao et al., 2021). During wheat seedling growth, MT accelerates the transformation from arginine and methionine to polyamines in wheat seedlings (Ke et al., 2018). Salt tolerance enhanced in maize with MT is most likely due to the improvement in photosynthetic capacity, antioxidative capacity, and ion homeostasis (increased K⁺ contents and K⁺/Na⁺ ratios) in leaves (Jiang C. et al., 2016). Also, Liu et al. (2020) demonstrate that MT improves salt tolerance in rice by reducing K⁺ efflux induced by high NaCl concentrations in roots. Subsequent researches have partly revealed the mechanisms of MT, which affects directly or indirectly salt resistance in many plants. However, MT application had no effects on the growth of wheat and maize under normal conditions (Ke et al., 2018; Ren et al., 2020). It has not fully elucidated a universal pathway on whether or how MT launches crosstalk with vital regulators under stress.

Different concentrations of MT may have differential effects in mediating abiotic stress defense. MT at 200 µmol L⁻¹ can effectively inhibit ROS production, improve the biomass and chlorophyll content, and regulate the photosynthetic characteristics of cotton seedlings under salt stress (Jiang et al., 2021). MT at 300 µmol L⁻¹ alleviates the negative effect of water stress on wheat germination and promotes morphological development including radicle length, radicle number, and plumule length of germinated seeds (Li et al., 2020). About 800 µmol L⁻¹ MT used for priming seed significantly improves germination energy, germination percentage, proline content, and total phenolic content of maize (Jiang X. et al., 2016). Application of exogenous MT (100 µmol L⁻¹) alleviates ROS burst and protects the photosynthetic activity in maize seedlings under salt stress (Chen et al., 2018). For some other reports, MT at a concentration of 100 µmol L⁻¹ inhibits seed germination and seedling growth and enhances the toxic effect of copper on seedlings, but 1 or 10 µmol L⁻¹ MT can eliminate the inhibitory effect of copper on the fresh weight of red cabbage seedlings (Posmyk et al., 2008). Pretreatment with 1 µmol L⁻¹ MT was also found to partially mitigate the inhibition of shoot dry weight induced by salt stress (Ke et al., 2018). Li X. et al. (2017) showed that 10–500 µmol L⁻¹ MT could help to recover seed germination potential, germination index, and vigor index on two varieties of rice treated with 120 mM NaCl to control levels. It seems that MT can regulate and enhance stress resistance by multiple mechanisms in a concentration-dependent manner.

In wheat production, different varieties show different resistances, adaptive areas, and production yields. So far, the roles of MT in mediating salt stresses on physiological regulation still need to be explored in different wheat varieties. In this study, seed germination and seedling growth of winter wheat experiments were conducted in a hydroponic solution with NaCl (100 mM) or without NaCl (100 mM), and the addition of MT. The purposes of this study were to (1) verify that MT enhances seed germination rate and antioxidant activities of seedling, (2) elucidate the regulatory effect of MT on ion exchange in the organelles, and (3) compare the regulatory effects of MT on the different varieties of wheat under salt stress. The results of this experiment provide theoretical support to promote the application of exogenous MT on saline-alkali wheat to increase arable land reserve and improve food security.
MATERIALS AND METHODS

Plant Material
The experiment was conducted in 2021 at the Hebei Agricultural University (38° 85' N, 115° 30' E), Hebei Province, China. Three commercial wheat (T. aestivum L.) cultivars, 'Jimai 22' (JM22), 'Shimai 15' (SM15), and 'Zhoumai 18' (ZM18), were used in this study. All varieties are high-yielding and extensively cultivated in Huang-Huai-Hai region. ZM18 is suitable for planting in south section of the Huanghuai winter wheat area, and JM22 is suitable for planting in north section of the Huanghuai winter wheat area. SM15 has comprehensive resistance and good adaptability.

Experimental Design
This experiment was performed in a growth chamber.

Seed Germination
In total, 450 seeds of each wheat variety were surface-sterilized with 75% ethanol for five minutes. After that, all surface-sterilized seeds were washed thoroughly with distilled water and were transferred into a glass plate for germination. Totally, two layers of filter paper saturated with three different treatment solutions: CK (distilled water), NaCl (100 mM), and NaCl-MT (100 mM NaCl + 300 µM MT) were placed in each plate. The concentrations of NaCl and MT solutions were selected referring to Li X. et al. (2017), Ke et al. (2018), and our former study (Li et al., 2020). The germination conditions were set at a light/dark shift (12/12h). The day/night temperature was 20/15°C. Light intensity was 600 µmol/m²/s. Relative humidity was 60%. The number of seed germinated was recorded daily till the 10th day. Water and other solutions were added into the germination plates to keep suitable humidity.

Hydroponic Experiments
At first, some other healthy seeds of three wheat varieties were germinated in a hole tray with moist vermiculite. The indoor environmental conditions were the same as those described above for the germination plate experiment. When the second leaf appeared, seedlings were removed out of hole tray gently and washed off vermiculite attached to the roots in tap water several times. Then, seedlings were transferred into Hoagland nutrient solution in plastic boxes covered with black film to avoid light. After about 1 week of recovery growth, seedlings were treated with fresh Hoagland nutrient solution as salt stress, 100 mM NaCl solution as salt stress, and 100 mM NaCl + 300 µM MT. Each treatment was repeated three times. After 24 h, leaf and root samples were collected and quickly frozen using liquid nitrogen and then stored in a −80°C freezer until the measurements were taken.

Measurement of Morphological Parameters
Germinated seeds were recorded daily when radicle length exceeded 2 mm according to Liu et al. (2016). Germ and radicle length and radicle number of 10 randomly selected grains from each treatment were recorded with a ruler for 24 h after treatment. Shoot and root length of five randomly selected seedlings from each replication were measured 24 h after treatment. Seedlings of each treatment with triplicates were dissected into roots, stems, and leaf and their fresh weights were recorded immediately. Then, the weighed fresh samples were kept in the paper bags to be dried in an oven at 105°C for 30 min and then at 75°C until the consistent weight was obtained.

\( \text{Na}^+, \text{K}^+, \text{and Ca}^{2+} \) Contents
\( \text{Na}^+, \text{K}^+, \text{and Ca}^{2+} \) contents of the whole-plant tissues were determined using a previous method with minor modifications (Chen et al., 2017). After the measurement of dry weight, samples were digested in a mixed solution of perchloric acid or concentrated nitric acid (volume ratio, \( \text{HClO}_4:\text{HNO}_3 = 1:5 \), v:v) in a glass test tube. More solution has replenished the solution to 12 mL. The ion content of a solution in a glass test tube was extracted in boiling water bath for 5 h and measured using an atomic absorption spectrophotometer (ZA-3000; Hitachi Instruments, Tokyo, Japan).

Hydrogen Peroxide Content and Detection of \( \text{H}_2\text{O}_2 \) and \( \text{O}_2^- \)
Hydrogen peroxide (\( \text{H}_2\text{O}_2 \)) content in the leaf and root samples was determined using a \( \text{H}_2\text{O}_2 \) assay kit (A064, Nanjing Jiancheng Bioengineering Institute, Nanjing, China) according to the manufacturer's user manual. \( \text{H}_2\text{O}_2 \) accumulation in root samples was visualized according to what Liu (2020) described. About 1.5 cm root tip was soaked and stained in 1% (w/v) 3,3′-diaminobenzidine staining (DAB, dissolved with 10 mM MES buffer on pH value = 6.5) and incubated in dark at 25°C for 8 h. Then, the samples were washed several times and observed using an optical microscope (Leica, Wetzlar, Germany).

Superoxide anion (\( \text{O}_2^- \)) accumulation in root samples (about 1.5 cm) was also visualized according to Liu’s (2020) method. Root samples were stained in 100 µM nitro blue tetrazolium chloride (NBT) dissolved with 50 mM phosphate buffer (pH value = 6.4) for 15 min and observed using an optical microscope (Leica, Wetzlar, Germany).

Antioxidant Enzymes, Soluble Sugar, Soluble Protein, and Malondialdehyde Content
Fresh leaf sample (0.5 g each) was triturated in 50 mM phosphate buffer (pH = 6.4). Homogenate solution was centrifuged at 20,000 g at 4°C for 15 min. Superoxide dismutase (SOD), peroxidase (POD), catalase (CAT) activity, soluble sugar, soluble protein, and malondialdehyde (MDA) contents were determined using assay kits (A064, Nanjing Jiancheng Bioengineering Institute, Nanjing, China) according to the manufacturer's instructions. Soluble sugar and protein contents were determined using a soluble sugar assay kit (A145-1-1, Nanjing Jiancheng Bioengineering Institute) and a BCA assay kit (BCP-1-W, Suzhou Comin Biotechnology Co., Ltd.) according to the manufacturer's user manual.
Extraction and Measurement of Amino Acid Content

Sample Preparation

The leaf and root samples (0.20 g) were accurately weighed and placed in the sample bottle (12 mL), into which 10 mL 6 mol/L hydrochloric acid (containing 0.1% phenol) was added. The mixed solution was homogenized by ultrasound. The samples were hydrolyzed in the oven at 110°C for 24 h. After cooling, the filtrate was mixed and filtered by a 0.45 µm water membrane. One milliliter of the filtrate was put into a rotary evaporator and dried at 80°C. Finally, 2.00 mL 0.1 mol/L hydrochloric acid was added to evaporated samples in the rotary evaporator and then mixed evenly by the whirlpool and transferred to the sample bottle for further use.

Standard Curve Development

Seventeen amino acid standards including aspartic acid (Asp), glutamic (Glu), histidine (His), serine (Ser), arginine (Arg), glycine (Gly), threonine (Thr), proline (Pro), alanine (Ala), valine (Val), methionine (Met), cysteine (Cys), isoleucine (Ile), leucine (Leu), phenylalanine (Phe), lysine (Lys), and tyrosine (Tyr) were calculated and accurately weighed in a volumetric flask. The mixed standard was diluted using 0.1 mol/L hydrochloric acid to obtain a final concentration of 500 mg/L of each amino acid. An appropriate amount of 500 mg/L mixed standard was diluted to make 10, 25, 50, 100, 150, 200, 300, and 400 mg/L mixed standard solution. The gradient elution procedures are shown in Table 1.

Aminoacyl Derivatization

Three solutions including 100 µL mixed standards, 200 µL buffer solution (0.5 mol/L sodium bicarbonate solution, 0.5 mol/L sodium carbonate solution, pH value = 9.0), and 100 µL 2,4-dinitrochlorobenzene (100 mg/mL) were swirled mixed and then reacted at 90°C for 90 min in dark. Thereafter, 50 µL of 10% acetic acid solution and 550 µL ultrapure water were added to 1.00 mL. The mixed solution was swirled and mixed and then filtered with 0.45-µm organic film and filter liquor prepared for measuring.

Chromatographic Analysis

An high-performance liquid chromatography (HPLC, Agilent 1200, United States) equipped with a C18 column (4.6 mm × 250 mm, 5 µm; Kromat Universil, catalog no. 35D5) and a diode array detector was used. The column temperature was 40°C. Mobile phase A was pure acetonitrile, and mobile phase B was acetic acid – sodium acetate buffer solution (2.50 g/L sodium acetate, 1.17 mL/L glacial acetic acid, 1.50 mL/L triethylamine, pH value = 5.25 ± 0.05). The flow rate was 1 mL/min. The sample injection amount was 10 µL with 10-to 15-min intervals. The detection wavelength was 360 nm.

Statistical Analyses

All data were processed using analysis of variance (ANOVA) in triplicate in Excel 2003 and IBM SPSS Statistics 17.0 (IBM Corp., Armonk, NY, United States). Duncan's new multiple range (DMR) test at a 5% probability level was used to test the differences among the means. Significant differences were labeled based on DMR.

RESULTS

Exogenous Melatonin Promotes Wheat Germination Trait

The effect of MT on wheat germination rate under salt stress is shown in Figure 1. The germination rate of CK reached the highest value on the third day and then showed a stable performance. The germination rate of wheat treated with NaCl increased rapidly in the first 3 days and then showed differences:

![Germination rate comparison](image-url)
TABLE 2 | Effects of MT on radicle number, root length, and germ length under salt stress.

| Varieties | Treatments | Germ length (cm) | Root length (cm) | Radicle number |
|-----------|------------|-----------------|-----------------|---------------|
| Jimai 22  | CK         | 15.17a          | 22.52a          | 5.47c         |
|           | NaCl       | 11.53b          | 13.34c          | 6.00c         |
|           | NaCl-MT    | 10.47b          | 7.05d           | 7.93a         |
| Shimai 15 | CK         | 13.29a          | 21.77a          | 5.00b         |
|           | NaCl       | 9.5733b         | 12.02c          | 5.07b         |
|           | NaCl-MT    | 7.255c          | 5.38d           | 6.55a         |
| Zhoumai 18| CK         | 13.94a          | 18.13a          | 5.00d         |
|           | NaCl       | 11.60c          | 12.07bc         | 5.27cd        |
|           | NaCl-MT    | 8.69d           | 5.39d           | 7.33a         |

For each trait, bars with the same letter are not significantly different according to Duncan’s test at a $p < 0.05$ threshold.

JM22 increased steadily, and SM15 and ZM 18 were basically stable. The germination rate of wheat decreased significantly under salt stress, especially for Jimai22, which was 11.09% lower than CK ($p < 0.05$), but there was no significant difference between Shimai 15 and Zhoumai18.

Exogenous Melatonin Inhibits Root Length and Germ Length and Promotes Radicle Number

Radicle number, root length, and germ length were also affected by salt stress significantly (Table 2). NaCl stress significantly decreased the length of germ and root of JM22, SM15, and ZM18 by 23.99 and 40.76, 27.97 and 44.79, and 16.79 and 33.43%, respectively. The radicle number did not change significantly under the NaCl treatment than that under CK. MT application further significantly decreased the length of germ and root of JM22, SM15, and ZM18 by 30.98 and 68.69, 45.41 and 75.29, and 37.66 and 70.27%, respectively. Meantime, the radicle number was significantly increased under NaCl-MT than the other two treatments.

Melatonin Application Relieve Inhibited Effect of NaCl on Shoot and Root Length

Figure 2 shows that shoot and root lengths of JM22 and ZM18 were both decreased significantly under NaCl stress compared with CK. MT application increased those indexes to some degrees, in which the shoot length of ZM18 and root length of JM22 were significantly increased under NaCl treatment than those under the NaCl-MT. Additionally, shoot and root lengths of ZM18 did not change significantly under NaCl and NaCl-MT treatments. These results indicated that SM15 showed higher salt resistance compared with the other two varieties.

Melatonin Application Increased Water Content in Plant Under NaCl Stress

Table 3 shows that fresh weight, dry weight, and water content in root, stem, and leaf of JM22 and ZM18 significantly decreased under NaCl stress. MT application had increased those indexes to the level of CK. However, for SM15, the fresh weight, dry weight, and water content in plants did not change significantly under the NaCl and NaCl-MT treatments compared with CK. These results also indicated that SM15 contained a good ability on salt resistance.

Melatonin Application Regulates Na\(^+\), K\(^+\), Ca\(^{2+}\) Compartmentation, and [K\(^+\)]/[Na\(^+\)] at Root and Leaf of Plant

Table 4 shows that K\(^+\), Ca\(^{2+}\), and Na\(^+\) contents in different organelles and varieties followed different changing patterns. Under NaCl stress, compared with CK, Na\(^+\) content significantly increased in root, stem, and leaf of three wheat varieties; K\(^+\) and Ca\(^{2+}\) contents were both significantly decreased in root and stem (Ca\(^{2+}\) content in the stem of SM15 and ZM18 exception); when K\(^+\) content increased, Ca\(^{2+}\) contents decreased significantly in the leaf of JM 22, but Ca\(^{2+}\) increased in the leaf of SM15; both K\(^+\) and Ca\(^{2+}\) in leaf were not obviously changed in ZM18. NaCl-MT treatment significantly decreased Na\(^+\) contents in root compared with that treated by NaCl alone, but increased in stem and leaf. This suggested that MT took effect first in the root, a higher Na\(^+\) concentration formed in stem and leaf. Also, MT application further decreased K\(^+\) content in stem and leaf and increased significantly Ca\(^{2+}\) content in the leaf of JM22; it decreased K\(^+\) content in the leaf of SM15 and decreased significantly K\(^+\) content in stem and increased Ca\(^{2+}\) in root and leaf of ZM18. [K\(^+\)]/[Na\(^+\)] ratio in root and stem + leaf was significantly decreased under NaCl stress compared with CK. MT application could
Melatonin Application Enhance Activity of Antioxidant Enzymes

Figure 3 shows that NaCl stress-induced increase of activity for SOD, POD, and CAT in different degrees compared with CK. MT application could enhance further the activity of SOD and CAT in the leaves of three wheat varieties under NaCl treatments; POD activity was recovered to the level of CK. It is noted that SOD activity of JM22 was significantly decreased under NaCl stress, which was possibly explained by the fact that JM22 was salt-sensitive and biochemical characteristics experienced serious damage.

Melatonin Application Decreased Malondialdehyde Content and Increased Proline Content

As shown in Figure 4, MDA content and proline content in the leaves of three wheat varieties were both increased under NaCl stress. MT application decreased MDA content under NaCl condition, suggesting that the cell membrane structure had been improved significantly. Additionally, MT further increased the proline content in the leaf of SM15 significantly under NaCl + MT compared with that under the NaCl treatment alone.

Melatonin Application Increased the Contents of Soluble Protein and Soluble Sugar

As shown in Figure 5, the contents of soluble protein and soluble sugar were decreased by different degrees under NaCl stress compared with CK. MT application significantly increased the contents of soluble protein and soluble sugar, which helped plants keep good water status under salt stress. Soluble protein content in the leaf of ZM18 did not change significantly under different treatments.

Melatonin Application Increased the Root Vigor Under NaCl Stress

Figure 6 shows that the root vigor of JM22 and ZM18 decreased significantly under NaCl stress compared with CK. However, the root vigor of SM15 was increased significantly under NaCl conditions, suggesting higher salt resistance for this variety. MT application had increased the root vigor of JM22 and ZM18 significantly to alleviate the salt damage.
Figure 7A shows that 

H$_2$O$_2$ and O$_2^-$ Accumulation in Root

Figure 7A shows that $H_2O_2$ contents in root of JM22, SM15, and ZM18 were significantly higher under NaCl stress than that under CK by 23.74, 33.89, and 20.24%, respectively. MT application reduced $H_2O_2$ contents in the roots of three varieties under NaCl stress to the level of CK. $H_2O_2$ and $O_2^-$ accumulation in roots of three varieties with three treatments were visualized by DAB and NBT staining. As expected, the detached roots of MT-treated wheat seedlings displayed less $H_2O_2$ and $O_2^-$ accumulation than those of NaCl-treated seedlings (Figure 7B). However, $H_2O_2$ contents in the leaf of JM22, SM15, and ZM18 were decreased significantly compared with CK. MT application maintained a low level of $H_2O_2$ contents in the leaves of three varieties.

**Figure 3** | Effects of MT on superoxide dismutase (SOD), peroxidase (POD), and catalase (CAT) under salt stress. For each trait, bars with the same letter are not significantly different according to Duncan’s test at a $p < 0.05$ threshold.

**FIGURE 3** | Effects of MT on superoxide dismutase (SOD), peroxidase (POD), and catalase (CAT) under salt stress.

**Melatonin Application Decreased $H_2O_2$ Content in Root and Displayed Less $H_2O_2$ and $O_2^-$ Accumulation in Root**

**Melatonin on Cluster Heatmap of Amino Acid Content of Wheat Seedlings Under Salt Stress**

Figure 8 shows the variations in amino acid content in plants of three wheat varieties under different treatments.

**DISCUSSION**

Salinity, one of the major abiotic stresses, limits the growth and productivity of many field crops. For increasingly extreme climate events, it is becoming the most important scientific research and agricultural practices in improving the salt tolerance of crops and exploiting arable areas of saline soils (Zhang et al., 2011; Himabindu et al., 2016). MT plays significant roles in antisenescence and antistress (Arnao and Hernández-Ruiz, 2014). It is well documented that exogenous MT can also improve the salinity resistance of wheat seedlings (Ke et al., 2018; Liu et al., 2020). During seed germination, exogenous MT (20 and 1 µM) pretreatment enhanced the germination rate of cotton, wheat, cucumber, and so on (Zhang N. et al., 2014; Ke et al., 2018; Chen et al., 2021). In our result, however, the germination rate showed somewhat improvement on ZM18 and SM15, no improvement on JM22, and furtherly inhibited the length of radical and germ under salt stress (%Table 2 and Figure 9). On the one hand, different MT concentrations for different crops might be partly reasons. However, concentration was not the main and only reason because MT (0–500 µM) pretreatment recovered root vigor and growth of two maize varieties and decreased relative electrolytic leakage in roots and leaves during the period of seed germination and seedling cultivation under NaCl stress (Li X. et al., 2017). This is likely for different treatment modes, plant pretreated with MT often shows better stress resistance even under lower concentrations, but MT was added directly to the culture solution especially in high concentrations, the seed germination was inhibited, and the growth of radical and radicle appeared retardation because plants would trigger a defensive response as external stress elements. On the other hand, the radicle number increased significantly under salt and MT condition, which suggested that MT promoted lateral root produced and growth of plants under NaCl stress. This result was consistent with the newest report that MT promotes lateral root under salt stress (Javeed et al., 2021).

Exogenous MT partially mitigated the salt stress-induced inhibition of whole-plant growth by reducing the accumulation...
of H$_2$O$_2$ in wheat leaf and increasing endogenous MT content and polyamine contents (Ke et al., 2018). Rice and cotton plants pretreated with MT also reduced H$_2$O$_2$ contents in leaf and roots, which accumulate high H$_2$O$_2$ concentration induced by salt stress (Li X. et al., 2017; Jiang et al., 2021). In our research, the changing rule of H$_2$O$_2$ content in root was also decreased with salt and MT treatments compared with NaCl alone treatment. However, H$_2$O$_2$ content in wheat leaf was significantly decreased under salt stress alone compared with CK, and MT application did not significantly recover H$_2$O$_2$ content to the level of control (Figure 7). The possible reason was that roots receive stress signals to turn on the defense system including recognizing and appointing exogenous MT as a stress regulator in a short time; for leaf, there was delayed effects for that MT has been reported as a potent long-distance signal to translocate stress message from roots to leaf (Li H. et al., 2017). Whether seeds of rice and wheat were pretreated with MT for 3–7 days, or the cotton seedling was treated with foliar spray MT every 24 h for 12 days, plants have been finished antistress training (Li X. et al., 2017; Ke et al., 2018; Jiang et al., 2021). Besides playing a vital signaling role during stress responses, H$_2$O$_2$ was induced by respiratory burst oxidase homologs (RBOH) to convert eventually into other ROS (OH) by the Fenton reaction and Haber–Weiss reaction (Liu et al., 2020; Michard and Simon, 2020). Additionally, MT treatment enhanced significantly the activities of antioxidant enzymes (especially SOD and CAT) and decreased MDA content in leaves resulted in a decrease in H$_2$O$_2$ level, which was conformed to this conclusion of Li X. et al. (2017). Not only does MT act as an endogenous antioxidant that enhanced the antioxidant capacity, but also MT was applied exogenously to modulate subcellular antioxidant systems in barley (Li et al., 2016). The possible response mechanism is still unknown comprehensively considering systematic reaction and crosstalk between organs.

Another effective strategy to resist salt stress employed by plants is to keep ion homeostasis and relieve ionic toxicity. One important approach that increases salt tolerance is to maintain a high level of [K$^+$]/[Na$^+$] ratio in cells (Flowers and Läuchli, 1983). Many of glycophytes are subjected to NaCl stress which showed that Na$^+$ in roots and leaves significantly increased, and K$^+$ content clearly decreased compared with those of the control (Li X. et al., 2017; Ren et al., 2020). Na$^+$ is sensed possibly by Na$^+$ sensor glucuronosyltransferase, activates Ca$^{2+}$ channels, and increases Ca$^{2+}$ influx into the cytosol (Jiang et al., 2019). In our study, MT application significantly decreased Na$^+$ contents, increased [K$^+$]/[Na$^+$] ratio in root induced by NaCl stress alone, but further increased Na$^+$ content, and decreased [K$^+$]/[Na$^+$] in stem and leaf. This suggested that MT may take ameliorating effect first on the root, and meantime, a higher Na ion concentration formed along with
nutrient solution flows and circulates in stem and leaf. It is known that the ion transport system including ionic equilibrium of [K$^+$] and [Na$^+$] and [K$^+$]/[Na$^+$] is often considered in a signaling network involving H$_2$O$_2$ and Ca$^{2+}$. Kaya et al. (2019) reported that MT treatments increase plant growth attributes to increased Ca$^{2+}$ and K$^+$ in the leaves and reduced MDA, H$_2$O$_2$ in Cd-stressed wheat plants. MT also helps cold-stress plants that possessed higher Ca$^{2+}$-ATPase, which are important for the ATP formation (Sun et al., 2018). In our results, SM15 also showed a higher Ca$^{2+}$ keeping ability in stem and leaf under salt stress alone; ZM 18 was more easily adjusted by MT to increase significantly Ca$^{2+}$ content in root and leaf under salt stress. These results conform to that the increase in Ca$^{2+}$ content in cytosol that triggers RBOH activity directly induces the formation of H$_2$O$_2$ that is eventually converted into other ROS (Michard and Simon, 2020), which is also interpreted as why H$_2$O$_2$ content decreased in leaves under salt treatment (Figure 7).

Salt stress promoted storage protein degradative, and thus, amino acid content changed accordingly (Zhang et al., 2017). Lysine (Lys) content is decreased with MT and transforms into other substances to raise the level of stress tolerance in wheat seeds during germination under drought stress (Li et al., 2020). MT also accelerates the metabolic flow from the precursor amino acids arginine and methionine to polyamines and decreases the degradation of salt-induced polyamines (Ke et al., 2018). From our study, the consistent results showed that salt stress increased Arg and Met contents, which decreased with MT application to some degrees. Additionally, Cys and Met contents in root of JM 22 and Cys content in the root of SM15 and ZM18 decreased under salt stress, but increased when MT application. This suggested that Cys is likely to participate in the H$_2$S-Cys cycle, which enhances its roles in the regulation of the antioxidant system (Huang et al., 2021). In our study, Lys and Tyr contents in leaf both increased under salt stress and further increased with MT application;
FIGURE 8 | The principal component analysis (PCA) (A) and cluster heatmap (B) of amino acid content of wheat seedlings under different treatments. Asp, aspartic acid; Glu, glutamic; His, histidine; Ser, serine; Arg, arginine; Gly, glycine; Thr, threonine; Pro, proline; Ala, alanine; Val, valine; Met, methionine; Cys, cysteine; Ile, isoleucine; Leu, leucine; Phe, phenylalanine; Lys, lysine; Tyr, tyrosine.

FIGURE 9 | Schematic diagram of seed germination and seedling growth with MT under salt stress.
His content in leaf increased significantly under both NaCl and NaCl-MT treatments. Other amino acids increased under NaCl and decreased under NaCl-MT without recovering to the level of control. It is known that MT often helps to induce amino acid accumulation in root and leaf to enhance the cellular osmotic potential (Cui et al., 2018). Lysine has been reported to be transformed into proline, aminobutyric acid, and polyamines during drought resisting processes (Hare and Cress, 1997; Klessig et al., 2000). So, our results also provided the evidence that MT increases the ability for the leaves of different wheat varieties to keep high water status and accumulate high soluble protein, soluble sugar, and proline content under NaCl stress (Table 3 and Figures 4, 5). On the other hand, lysine resides on histone terminus are deacetylated by histone deacetylase 14, which is involved in the biosynthesis of MT in Arabidopsis thaliana (Zhao et al., 2018). Both Lys and His contents increased in leaf under NaCl and NaCl + MT which indicated a complex biochemical process, during which the MT content changing the interaction between MT and amino acid, and involved mechanism would be the next important research point.

It is noted that MT application methods (pretreated coating or soaking, leaf pray, root or rhizospheric application, mixed application with other growth regulators) and varied experimental elements (MT concentration, plant species, varieties, and adversity types) appear to have different regulatory effects during stress-resistance process (Kostopoulou et al., 2015; Jiang C. et al., 2016; Li X. et al., 2017). In our research, different wheat varieties showed different salt-sensitivities. JM22, a widely adaptive super-high-yield variety, was not salt-resistant and shows an obvious decrease on germination rate and activity of SOD and CAT under NaCl stress. SM15 and ZM18 showed higher salt-resistant, considering that SOD activity of SM15 did not decrease but increased significantly and higher germination rate and proline accumulation under NaCl stress alone. MT at a concentration of 300 µmol L\(^{-1}\) played roles as regulated antioxidize to keep the physiological equilibrium of wheat seedling and to promote germination and lateral roots, but it took an inhibitory effect on the length of radicle and germ. The optimal concentration of MT on wheat seed germination and growth of wheat seedling and the dominant varieties of wheat for coping with salt stress should be further researched for meeting future production reality needs.

### DATA AVAILABILITY STATEMENT

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

### AUTHOR CONTRIBUTIONS

DL and RL initiated and designed the research. ZZ, HL, XF, and SZ performed the experiments and collected the data. XZ, NY, and JS wrote the code and tested the methods. LL, ZZ, and DL analyzed the data and wrote the manuscript. All authors contributed to the article and approved the submitted version.

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### SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fpls.2021.787062/full#supplementary-material
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