Mechanisms of Salt Tolerance and Molecular Breeding of Salt-Tolerant Ornamental Plants

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As the area of salinized soils increases, and freshwater becomes more scarcer worldwide, an urgent measure for agricultural production is to use salinized land and conserve freshwater resources. Ornamental flowering plants, such as carnations, roses, chrysanthemums, and gerberas, are found around the world and have high economic, ornamental, ecological, and edible value. It is therefore prudent to improve the salt tolerance of these important horticultural crops. Here, we summarize the salt-adaptive mechanisms, genes, and molecular breeding of ornamental flowering crops. We also review the genome editing technologies that provide us with the means to obtain novel varieties with high salinity tolerance and improved utility value, and discuss future directions of research into ornamental plants like salt exclusion mechanism. We considered that the salt exclusion mechanism in ornamental flowering plants, the acquisition of flowers with high quality and novel color under salinity condition through gene editing techniques should be focused on for the future research.

Keywords: ornamental plants, salinized soil, salt tolerance, breeding, value

INTRODUCTION

Degradation of soils caused by salinization, a major abiotic stress factor, is an increasing limitation of arable land. Almost 10% of all soils and 50% of the irrigated land worldwide are affected by salinity (Guo et al., 2015, 2018; Wang et al., 2015, 2018; Song et al., 2016, 2017). Salinized soils limit the growth, development, and survival of plants that grow in such environments. Salinity poses a serious threat to food production and security (Munns and Tester, 2008; Abdullakasim et al., 2018). As the human population grows and urbanization increases, the area of land suitable for cultivation is decreasing. Improving living standards also mean increased demand for habitable land. Planting salt-tolerant flowering plants has become a feasible and sustainable strategy by which to use saline land without incurring competition for soils for food cultivation. Therefore, there is much interest in breeding high-value salt-tolerant ornamental flowering varieties, and to expand the usage of underutilized ornamental species for exploitation of saline soils.

Excess salinity in soils directly reduces the water potential (become more negative) around the roots, making it difficult for root cells to extract water and leading to water deficit (Chinnusamy et al., 2005; Ulczycka-Walorska et al., 2020; Wani et al., 2020; Zheng et al., 2021).
Thus, to survive in such environments, plants must reduce the water potential of their own cells (Shabala et al., 2015). When salt ions accumulate in plant cells, ionic toxicity occurs. Subsequently, secondary damage such as oxidative stress and nutrient deficiencies can occur (Breš et al., 2016; Van Zelm et al., 2020). Some salt-sensitive plants may display stunted growth or even die because of damage caused by salinity. However, salt-tolerant plants can initiate various protective mechanisms that allow them to grow in saline environments; for example, changes in gene expression and regulation allow these plants to adapt their morphology, physiology, and biochemistry in response to salinity (Bai et al., 2018; Zandalinas et al., 2018; Guo et al., 2019a; Qi et al., 2020; Zhao et al., 2020). Additionally, the damage to plants caused by salinity depends on the species and environmental factors.

Flowering plants have many important roles for humans. They can be used as ornamental growing plants, as cut flowers, for environmental greening, as medicine, as fruits and vegetables. In addition, the flowering time of many plants can be changed to meet the option or the market needs. Recent reviews have explored the biodiversity of edible flowers (Boutigny et al., 2020), flower color regulation (Zhao and Tao, 2015), and breeding of mutant ornamental flowering plants (Anne and Lim, 2020). However, a review of the mechanisms of salt tolerance in ornamental flowering plants and of strategies to produce salt-tolerant ornamental plants or breed new, salt-tolerant varieties is lacking. Salinity is particularly responsible for degrading the visual quality of ornamental flowers (Jaleel et al., 2008; Yang et al., 2019). Therefore, here we review the salt-tolerant mechanisms of ornamental flowering plants, and the selection and breeding of new salt-tolerant ornamental flowering varieties.

**SALT TOLERANCE MECHANISMS OF ORNAMENTAL PLANTS**

The level of salt tolerance of ornamental plants depends on the species, their development, the level of salt stress, and environmental factors. If the ornamental plant could grow and survive at the salt level at or over 200 mM NaCl that could be considered as a halophyte, thus the ornamental plants can be divided in halophytic and non-halophytic ones according to their salt tolerant ability. For example, the salinity threshold was 400 mM NaCl of the halophytic Limonium sinuatum under laboratory conditions (Mi et al., 2021). Many research findings have indicated that the development of salt-tolerant ornamental flowers is a viable approach. Although the opening time of the first flower was delayed, marigold (Tagetes erecta) can be planted in saline fields (García-Caparrós et al., 2016; Hao et al., 2017). Some ornamental species, such as fuchsia (Fuchsia hybrida), coleus (Solenostemon scutellarioides), and begonia (Begonia hiemalis; Villarino and Mattson, 2011), can tolerate a certain concentration of NaCl (7.0–9.8 dS m⁻²) without demonstrable growth inhibition. No negative impact was detected in lily (Lilium × elegans) seedlings treated with a moderate concentration of salt (3 dS m⁻²). Furthermore, the effect of salinity on growth could be alleviated by applying K⁺ to the grown medium (Ayad et al., 2019).

Although there has been some research into the salt tolerance of ornamental plants, the underlying mechanisms are complex, and vary between ornamental plants. Therefore, detailed studies still need to be conducted to provide basis for future research and utilization of ornamental plants. The salt-tolerant mechanisms of ornamental plants are summarized in Figure 1, and the detailed experimental condition of plant species is tabulated in Table 1. These mechanisms include the regulation of osmotic balance under the osmotic stress caused by high salinity; adjustment of ionic balance to avoid ionic toxicity; active oxygen scavenging to reduce oxidative damage; the exclusion of salt from the roots by enhancing apoplastic barriers (like Chrysanthemum), salt release via Na⁺/H⁺ antiporters (like Salicornia), and salt secretion by salt glands or salt bladders (like Atriplex); photosynthetic regulation to maintain high photosynthetic efficiency (like Aster); limiting water loss by closing stomata, and thickening leaf surface wax (like Dianthus).

**Adaptation to Salinity by Osmotic Adjustment and Ionic Homeostasis**

Like many other plants, the first threat faced by ornamental plants in a saline environment is the lowered external water potential caused by the presence of salt ions. This prevents plant roots from taking up water from the soil, and can even cause water to drain out of the cells (Feng et al., 2014a), thus reducing plants’ ability to grow and survive. Accumulation of organic and inorganic solutes in the root cell cytoplasm (Feng et al., 2015) reduces the water potential of the cells and ensures water uptake under salt stress conditions (Han et al., 2012; Shao et al., 2014). In the process, some small organic molecules (compatible solutes), such as proline, will synthesize and accumulate in the cytosol of plant cells as the osmotic adjustment substances to cope with the osmotic stress (El-Shawa et al., 2020). These are also used to protect the structure of cells and macromolecular substances. In some cases, exogenous application of compatible solutes can increase plant salt tolerance. For example, increased antioxidant capacity was observed in plants to which exogenous glycine betaine (GB) was applied, compared with those without (Roychoudhury and Banerjee, 2016). At the same time, inorganic ion accumulation is also the way in which most salt-tolerant plants, such as halophytes, to reduce the osmotic potential. In this process, ions such as Na⁺ and Cl⁻ mainly accumulate in the vacuole so they can be used for osmotic adjustment of the plant cell (Chen and Jiang, 2010). Accumulation of inorganic ions as a strategy that consumes less energy than synthesize organic substances.

In addition, more Na⁺ and Cl⁻ accumulated in the roots than in the shoots of Dendrobium orchid under saline conditions. This high Na⁺/K⁺ ratio in the roots contributed to osmotic adjustment (Abdullakasim et al., 2018). More proline accumulated in seedlings of sweet osmanthus (Osmanthus fragrans (Thunb.) Lour.), germinated from gamma-irradiated seeds than those of controls, thus the salt tolerance of the seedlings was increased by improving osmotic adjustment (Geng et al., 2019). Salinity
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Reduced the leaf area and flower weight of marigold (Calendula officinalis L.), but increased the proline content in these plants. This increased proline content conferred salt tolerance to marigolds in saline conditions below 150 mM NaCl (Adamipour et al., 2019). Despite decreased seedling biomass, proline accumulated and was the main osmolyte responsible for the osmotic adjustment of ornamental plants in response to saline conditions (El-Shawa et al., 2020). In the ornamental crop Hyacinthus orientalis L., proline in the leaves increased under saline conditions of 6.1–8.6 g L⁻¹. The ornamental qualities of these flowers also increased, indicating promise that this crop could be cultured in saline soils with concentrations of sodium chloride as was used in the laboratory culture (Ulczycka-Walorska et al., 2020). In the ornamental crop Hyacinthus orientalis L., proline in the leaves increased under saline conditions of 6.1–8.6 g L⁻¹. The ornamental qualities of these flowers also increased, indicating promise that this crop could be cultured in saline soils with concentrations of sodium chloride as was used in the laboratory culture (Ulczycka-Walorska et al., 2020).

Ionic homeostasis in plant cells is the basis for development and metabolism, especially the higher K⁺ and lower Na⁺ concentration in the cytosol of salt-tolerant plant cells (Blumwald, 2000). In salt sensitive plants under salt stress, ionic disturbance is very important, while salt-tolerant plants have a greater ability to maintain ionic balance (Guo et al., 2019a). With a similar charge and physiological characteristics, Na⁺ might compete with K⁺ for active sites of enzyme transporters. Thus, competition between the two ions leads to disruption of enzymatic function and biosynthesis (Zhu, 2003). For plants to grow and develop well under saline conditions, maintenance of a proper ratio of cytosolic K⁺/Na⁺ is required. This, along with the homeostasis of other ions, such as Ca²⁺, Mg²⁺ and Fe²⁺, is also a reliable indicator of the level of plants' salt tolerance (Munns and Tester, 2008).

In salt-tolerant plants, ionic homeostasis might be maintained by excluding salt ions in soil through the roots by Salt Overly Sensitive 1 (SOS1) and apoplastic barriers (Hose et al., 2001; Meyer et al., 2009; Krishnamurthy et al., 2011; Song et al., 2012; Gao et al., 2016); thus fewer ions might be translocated to, or accumulate in, the aboveground parts of the plant, such as leaf cells, by compartmentalizing salt ions into the vacuole and reducing the ionic concentration in cytoplasm (Qiu et al., 2007). Therefore, restraining ionic transport to the shoots and

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**FIGURE 1** | Graphical summary of the adaptation of ornamental plants to salinized environments. (A) Adaptation through ionic homeostasis and osmotic adjustment; (B) Adaptation through ROS scavenging; (C) Adaptation through salt exclusion, removing and pumping out of cells in roots; (D) Adaptation through salt secretion, leaf succulence, photosynthesis protection, and reduction of water loss in shoots. SOS1: Salt Overly Sensitive 1, e.g., CcSOS1 gene from Chrysanthemum cismus; NHK1: vacuolar Na⁺/H⁺ antiporter, e.g., IbNHX2 gene from Ipomoea batatas; HKT1: high-affinity K⁺ transporters, e.g., AhHKT2:1 gene from Aeluropus lagopoides; WRKY: e.g., DgWRKY4 genes of from Dendronthema grandiform; NAC: e.g., DgNAC1 gene from Dendronthema grandiform; ROS: reactive oxygen species; SOD: superoxide dismutase; CAT: catalase; APX: ascorbate peroxidase; POD: peroxidase; GR: glutathione reductase; RC: reaction center. Symbol size represents relative content of ions and organic matter in plant cells.
TABLE 1 | The detailed experimental conditions of the ornamental flowering plants exposed to salt stress.

| Species                          | Duration and extent of NaCl stress | Main phenotype                                                                 | References                                      |
|---------------------------------|-----------------------------------|--------------------------------------------------------------------------------|------------------------------------------------|
| Gerbera jamesonii               | 30 mM/57 days                     | Mitigated phenotypes in leaves and flowers                                    | Gómez Bellot et al., 2018                      |
| Narcissus sp. (L.)              | 150 mM/5–6 weeks                  | Maintained a steady K+/Na+ ratio                                              | Veatch-Blohm et al., 2014                      |
| Lobelia erinus                  | 30–50 mM/60 days                  | Reduced growth and biomass with no effect on flower production                | Escalona et al., 2013                          |
| Viola x wittrockiana Gams.      | 5–7 g L⁻¹/8 weeks                 | No effect on flower production with reduced growth of plant                   | Pušić et al., 2019                             |
| Catharanthus roseus             | 80 mM/30 days                     | Inhibited growth and the activities of antioxidant enzymes                   | Jakeel et al., 2008                            |
| Limonium                        | More than 200 mM/60 days          | Grown well with salt excreting in leaves                                      | Soud et al., 2016; Mi et al., 2021; Duarte et al., 2017 |
| Aster tripolium L.              | 250 mM/2 weeks                    | High light harvesting efficiencies and low energy dissipation rates           |                                               |
| Callistemon citrinus            | 4 dS m⁻¹/56 weeks                 | Greater Na⁺ storage                                                          | Álvarez and Sanchez-Blanco, 2014               |
| Portulaca                       | 400 mM/5 weeks                    | Leaf succulence with high ionic accumulation in the vacuoles                 | Borsai et al., 2020                            |
| Iris germanica                  | 140 mM/28 days                    | Lower Na⁺ content in rhizomes than in leaves or roots under saline conditions  | Zhao et al., 2021                              |
| Tagetes erecta                  | 4.5 or 7.5 dS m⁻¹/60 days         | Delayed the opening time of the first flower                                 | García-Caparrós et al., 2016; Hao et al., 2017 |
| Dianthus caryophyllus           | 200 mM/15 days                    | Reduced stomatal conductance and increased epicuticular wax                  | Kwon et al., 2019                              |
| Chrysanthemum morifolium        | 0.4–2.0%/1 week                   | Increased photosynthetic pigments with a better seedling growth              | Vanlairuati et al., 2019                      |
| Fuchsia hybrida                 | 7.0–9.8 dS m⁻¹/5 weeks            | With no demonstrable growth inhibition                                        | Villarino and Mattson, 2011                   |
| Solenostemon scutellarioides    | 7.0–9.8 dS m⁻¹/5 weeks            | With no demonstrable growth inhibition                                        | Villarino and Mattson, 2011                   |
| Begonia hiemalis                | 7.0–9.8 dS m⁻¹/5 weeks            | With no demonstrable growth inhibition                                        | Villarino and Mattson, 2011                   |
| Hyacinthus orientalis L.        | 6.1–8.6 dS m⁻¹/12 weeks           | Accumulated more proline                                                     | Ułczycka-Walorska et al., 2020                |
| Pelargonium hortorum L.H. Bailey| Lower than 1.976 g L⁻¹/77 days    | Accumulated more proline and anthocyanin                                     | Břeš et al., 2016                              |
| Osmanthus fragrans (Thunb.) Lour.| 40–120 mM/3 days                  | Accumulated more proline                                                     | Geng et al., 2019                              |
| Nerium oleander L.              | 80 mM/15–30 days                  | Increased activities of SOD and CAT                                          | Kumar et al., 2017                            |
| Lavandula angustifolia Mill.    | 300 mM/30 days                    | Increased activities of SOD and CAT                                          | Szekely-Varga et al., 2020                    |
| Parthenium hysterophorus        | 160 mM/10 days                    | Increased activities of SOD, GR, CAT and contents of proline, Ascorbate, and glutathione | Ahmad et al., 2017                            |
| Eugenia myrtifolia L.           | 88 mM/30 days                     | Increased activities of SOD                                                   | Acosta-Motos et al., 2015                     |
| Tagetes patula L.               | 100 mM/10 days                    | Increased activities of antioxidant, and contents of polyphenol, carotenoid   | Chryssargyris et al., 2018                     |
| Echinacea purpurea              | 50–100 mM/2 weeks                 | Increased activities of SOD and APX                                           | Sabra et al., 2012                            |
| Amaranthus tricolor             | 50–100 mM/24 days                 | Increased activities of SOD, APX and ascorbate, carotenoids                  | Sarker and Oba, 2020                          |
| Portulaca grandiflora           | 400 mM/5 weeks                    | Enhanced salt tolerance                                                      | Borsai et al., 2020                            |
| Iris halophila (Pall.)          | 400 mM/5 weeks                    | Enhanced salt tolerance                                                      | Borsai et al., 2020                            |
| Eragrostis spectabilis (Pursh) Steud.| 5.0 or 10.0 dS m⁻¹/65 days      | High visual quality was maintained                                            | Wang et al., 2019c                             |
| Viburnum lucidum L.             | 200 mM/103 days                   | Higher chlorophyll and K⁺ content when applying of exogenous GB or proline   | Citrillo et al., 2016                          |
| Callistemon citrinus Stapf      | 200 mM/103 days                   | Higher chlorophyll and K⁺ content when applying of exogenous GB or proline   | Citrillo et al., 2016                          |
| Aloe vera (L.)                  | 2.0–7.5 dS m⁻¹/60 days            | Higher ions in roots than in shoots                                           | García-Caparrós et al., 2016                   |
| Kalanchoe blossfeldiana Poelln.| 2.0–7.5 dS m⁻¹/60 days            | Reduced Na⁺ when older leaves are shed                                        | García-Caparrós et al., 2016                   |
| Gazania splendens L. sp.        | 2.0–7.5 dS m⁻¹/60 days            | Increased the succulence index with more Na⁺ and Cl⁻ in roots                 | García-Caparrós et al., 2016                   |
| Viburnum lucidum                | 10–70 mM/120 days                 | Higher ions in roots than in shoots                                           | Cassaniti et al., 2009                        |
| Alternanthera bertzickiana (Regel) G. Nicholson | 40 dS m⁻¹/20 days | Higher Na⁺ content in roots than shoots                                      | Ali et al., 2012                               |
| Callistemon laevis              | 4 dS m⁻¹/56 weeks                 | Higher Na⁺ content in roots than stems                                        | Álvarez and Sanchez-Blanco, 2014               |
| Dendrobium orchid               | 2–15 dS m⁻¹/30 days               | Higher Na⁺ and Cl⁻ content in roots than shoots                               | Abdullahakasim et al., 2018                    |

compartmentalizing toxic ions in the vacuole are the important pathways to enhance plants’ salt tolerance in saline environments (Hasegawa, 2013). In the ornamental plant *Eugenia myrtifolia* L., more salt ions accumulated in roots than in leaves to perform normal physiological metabolism under salinity conditions (Acosta-Motos et al., 2017). In salt-tolerant chrysanthemum varieties, growth increased under saline conditions, together with increased K⁺, Ca²⁺ and Mg²⁺ content,
and more efficient usage of N and P (Rahi and Singh, 2011). Results in marigold (Tagetes patula) indicated that salt tolerance could be improved by enhanced potassium application (Aboutalebi Jahromi and Hosseini Farahi, 2016).

For the ornamental plant gerbera (Gerbera jamesonii Bolus ex Hook, f., cv. "Forsa"), the presence of NaCl (such as 30 mM) could mitigate the phenotypes in leaves and flowers affected by boron (Gómez Bellot et al., 2018). A steady K+/Na+ ratio was maintained despite high Na+ accumulation in the leaves of daffodils [Narcissus sp. (L.) Amaryllidaceae] when treated with different salt concentrations—even under 150 mM NaCl conditions (Veatch-Blohm et al., 2014). While for the halophytic ornamental plant Lobularia maritima L., which growth was unaffected by 100 mM NaCl with the relative ionic balance in roots and leaves (Hsouna et al., 2020). Another halophytic ornamental plant Sesuvium portulacastrum displayed an optimal growth treated with moderate salinity with accumulating of large amounts of true halophyte and shows an optimal development under moderate salinity with a large amount of salt ions accumulating in the leaves (Nikalje et al., 2018; Ding et al., 2022). In Callistemon laevis treated with NaCl, little Na+ concentration was detected in the leaves, and more Na+ accumulated in the roots and stems, indicating that more salt ions were prevented from reaching the aboveground parts of the plant (Álvarez and Sanchez-Blanco, 2014). In ornamental grasses, such as Eragrostis spectabilis (Pursh) Steud., and Panicum virgatum L. “Northwind,” high visual quality was maintained when treated with NaCl, despite high concentrations of salt ions (Na+ and Cl−) accumulating in leaves (Wang et al., 2019c).

In Lobelia erinus, plant growth and biomass were reduced with increased salinity, while salinity (30 or 50 mM NaCl) had no effect on flower production, and no toxic symptoms were observed on leaves, despite high accumulation of Na+ and Cl− in leaf tissues. This indicates that L. erinus could be used for urban landscaping (Escalona et al., 2013). A similar result was observed in pansy (Viola x wittrockiana Gms.), which displayed salt tolerance at levels of 5 or 7 g L−1 NaCl (Pušić et al., 2019). Under salt stress, uptake of Ca2+, Mg2+, and Na+ in the shoots of marigold (Tagetes erecta L.) increased, indicating an increase in salt tolerance (Koksal et al., 2016). When Callistemon citrinus plants were subjected to salt stress, greater Na+ storage was detected, which played a role in maintaining plant quality, and indicating that C. citrinus could be cultured with 4 dS m−1 saline water (Álvarez and Sanchez-Blanco, 2014).

It is well known that high salinity decreases K+ content in plants. A saline environment reduced the growth of marigold, with decreased K+ content, while the Mg2+ content increased in the leaves under such conditions (Koksal et al., 2016). However, the K+ content, as well as the carotenoid content, in pelargonium (Pelargonium hortorum L.H. Bailey) leaves was not affected when plants were grown at salinities lower than 1.976 g L−1. Meanwhile, the increased anthocyanin and proline content in leaves might be important in ameliorating the adverse effects caused by salt stress (Brés et al., 2016).

Salt-tolerant species can maintain ion homeostasis during growth, and this is inseparable from the roles of membrane-bound ion transporters, such as Na+/H+ antiporters. Under salt stress conditions, the plasma membrane-located Na+/H+ antiporter SOS1, and the NHX antiporters on tonoplastic membranes are vital in maintaining the cellular ion homeostasis by transporting excess Na+ out of the cells and into the vacuole (Munns and Tester, 2008; Yue et al., 2012). In the working model of the SOS system pathway, cellular Ca2+ signals were identified as being involved. Firstly, salt stress induced an increased concentration of Ca2+ in the cytosol, which is sensed by, and causes the activation of SOS3. SOS2 is then activated by activated SOS3, forming an SOS2–SOS3 complex. Thirdly, SOS1 is activated, which mediates the transport of excess Na+ out of the cells and maintains a relatively low Na+ concentration in the cytoplasm (Tang et al., 2015). A significant reduction in Na+ content, and a favorable K+/Na+ ratio, was detected in transgenic plants—especially in the youngest leaves—in Chrysanthemum crassum overexpressing the CaSOS1 gene. Thus, the transgenic plants demonstrated higher salt tolerance, up to 200 mM NaCl (An et al., 2014).

NHX family transporters are localized to the tonoplast, they are widely considered as the players to sequestrate sodium (Na+) into vacuoles and to avert the cytoplasmic ion accumulation when plants exposed to salinity. In Arabidopsis, the expression levels of AtNHX1 and AtNHX2 genes were upregulated under osmotic stress or by abscisic acid (ABA; Jiang et al., 2011). Overexpression of HtNHX1 and HtNHX2 transporters from Helianthus tuberosus improved the salt tolerance of rice (Zeng et al., 2018). Salt or drought tolerance was enhanced in transgenic sweet potato [Ipomoea batatas (L.) Lam.] with overexpression of the IbhNHX2 gene (Wang et al., 2016).

In view of this research on the ornamental flowering plants, salt-tolerant varieties seem to prefer to accumulate inorganic ions in the vacuole to lower cell water potential, and simultaneously synthesize some compatible solutes in the cytosol to balance the decline of vacuolar water potential. This strategy thereby ensures the growth of plants and their increasing salt tolerance. A reconstruction of osmotic and ion homeostasis pathways in a plant cell under saline stress is shown in Figure 1A.

**Adaptation to Salinity by Maintaining the Balance of Reactive Oxygen Species**

Reactive oxygen species (ROS) are free radical molecules that are inevitably produced during plant metabolism. They have important roles in plant growth, development, and adaptation to stressors. Under normal conditions, ROS production and scavenging is kept in balance. However, when plants are subjected to adversity, such as salt or drought stress, more ROS are produced, and they will accumulate if production exceeds scavenging. Excess accumulation of ROS damages structural and metabolic plant processes (Pang et al., 2011). In salt-tolerant species, antioxidant capacity is enhanced to scavenge accumulated ROS, thus restoring the balance of ROS. Usually, ROS scavenging is implemented in two main ways: non-enzymatic and enzymatic mechanisms (Bowler et al., 1992; Mittler, 2002). Superoxide dismutase (SOD) and catalase (CAT) are usually considered the two most important antioxidant enzymes, but
others, such as ascorbate peroxidase (APX), peroxidase (POD), and glutathione reductase (GR), are all employed to scavenge ROS in enzymatic reaction systems (Kozi, 1999). Furthermore, they work together to regulate the balance of ROS in plants.

When ROS levels are in equilibrium, membrane stability is maintained and normal metabolism occurs in plants (Li et al., 2012; Luo et al., 2013). Tolerance to other stresses, such as drought or heat, is also enhanced (Reddy et al., 2004). During ROS scavenging, enzymatic and non-enzymatic activity are strongly related to plants’ ROS scavenging ability. Overexpression of genes related to antioxidant enzymes could improve ROS scavenging, thus enhancing stress tolerance (Caverzan et al., 2016). In the ornamental plant Lavandula angustifoli Mill., antioxidant enzymes, including SOD and CAT, were activated under salt stress (even at 300 mM NaCl), with enhanced accumulation of flavonoids and phenolic substances, thus improving plants’ oxidation tolerance (Szekely-Varga et al., 2020). In salt-treated Echinacea species, E. purpurea displayed higher Na⁺ exclusion ability, and enhanced antioxidant activity of APX and SOD, but CAT activity was reduced and there was no change in GR activity (Sabra et al., 2012). The highly salt-tolerant ornamental plant, Parthenium hysterophorus could grow in a concentration as high as 160 mM NaCl. It also displayed increased proline, ascorbate, and glutathione contents, and elevated activity of enzymatic antioxidants, such as SOD, GR, and CAT (Ahmad et al., 2017). With the high salt tolerance (800 mM NaCl) of the ornamental plant Nerium oleander L., the activities of SOD and CAT were increased in the leaves and roots, even though more salt ions accumulated in these organs (Kumar et al., 2017). In tagetes (Tagetes patula L.), polyphenol and carotenoid contents, and antioxidant activities increased under 100 mM NaCl, and there was high accumulation of N, P, Zn, and Na in the flowers (Chrysargyris et al., 2018). In Eugenia myrtifolia L., SOD content and activity in the leaves increased despite reduced APX content and activity under saline conditions, and lower Na⁺ in the leaves than roots ensured normal photosynthesis (Acosta-Motos et al., 2015). The antioxidant ability of plants could also be enhanced by gene mutation. The activities of SOD, CAT, and POD were significantly increased in seedlings of sweet osmanthus germinated from gamma-irradiated seeds compared with controls; seedlings’ salt tolerance was improved by regulating ROS balance (Geng et al., 2019). When treated with 80 mM NaCl, growth and the activities of antioxidant enzymes of Catharanthus roseus plants were inhibited. However, the salt tolerance of the seedlings was enhanced, with increased plant weight, SOD, and POD activities when plants were grown with NaCl and propiconazole, except leaf area and stem length decreased (Jaleel et al., 2008).

Salt tolerance is different even in different varieties of the same species. In Amaranthus tricolor, ascorbate, carotenoids (non-enzymatic antioxidant), and SOD and APX (antioxidant enzymes) activities were increased in the salt-tolerant varieties compared with the salt-sensitive ones, indicating that increased ROS quenching ability conferred higher tolerance (Sarker and Oba, 2020). The activities of APX, SOD, POD and CAT in marigold were enhanced when plants were grown under conditions below 150 mM NaCl, then decreased at 200 mM NaCl (Adamipour et al., 2019), indicating that marigold could be developed for cultivation on saline lands.

In saline environments, large amounts of ROS accumulate in plants, resulting in ROS imbalance. Salt-tolerant ornamental varieties usually show improved ROS scavenging ability, both from enzymatic and non-enzymatic reaction processes, so this could be considered an indicator of high salt tolerance in ornamental flowering plants (Figure 1B).

**Adaptation to Salinity by Root Salt Exclusion**

Another measure that plants employ to reduce cellular ion content is to reduce the uptake of salt ions in the roots, or to excrete some absorbed salt ions out of the plant (Figure 1C). Thus, a relatively higher Na⁺ content will be detected in roots than in shoots (Abdullakasim et al., 2018). The same was found under cadmium stress (Qi et al., 2020). Salt tolerance was conferred by a special structure, such as thickened apoplastic barriers (Krishnamurthy et al., 2020). In some ornamental plants, such as Escallonia rubra, Cestrum fasciculatum, Viburnum lucidum (Cassaniti et al., 2009), and Aloe vera (L.; García-Caparrós et al., 2016), the mechanism employed to respond to salinity correlated with higher accumulation of ions in roots than in shoots. Na⁺ content in rhizomes of Iris germanica was usually lower than in leaves or roots under saline conditions, which may indicate a salt stress adaptation mechanism (Zhao et al., 2021). Perhaps, in such plants under saline conditions, the greater accumulation of salt ions in roots is associated with structural changes of roots related to the apoplastic exodermal barriers, such as the thickened endothelium, preformed suberin lamellae and Casparian bands (Andersen et al., 2015). In Opisthopappus taihangensis (Ling) Shih and Iris germanica (Meyer et al., 2009; Yang et al., 2020), salt ion uptake was reduced in root cells. Additionally, the absorbed ions in root cells could also be excreted out of the roots cells through antiporters such as SOS1 (Song et al., 2012; Gao et al., 2016; Mohammadi et al., 2019), resulting in increased ion accumulation in roots and reduced salt ion transport to shoots. However, the relationship between salt ion accumulation and apoplastic barriers remains to be investigated in ornamental flowering plants.

As well as decreasing salt ion uptake through structural features, absorbed salt ions can also be removed from xylem. The HKT1 (high-affinity K⁺ transporters) gene plays vital roles in this process (Ali et al., 2016), and the HKT1 transporter can also restrict Na⁺ entry into plant roots (Rus et al., 2001). With the function of HKT1 localized in xylem parenchyma cells, Na⁺ flux to the shoot tip was actively reduced and excess Na⁺ accumulated in the root zone of plants. In Arabidopsis, AHKT1 was proposed to recirculate Na⁺ in the phloem, reducing the allocation of Na⁺ in the shoot, and correspondingly increasing Na⁺ content in roots (Davenport et al., 2007).

Furthermore, absorbed Na⁺ can also be excreted from root cells through transporters such as SOS1, which are localized in the plasma membrane of root cells (Shi et al., 2000; Oh et al., 2009), like the CcSOS1 transporter in chrysanthemum (Song et al., 2012; Gao et al., 2016). Perhaps, in ornamental
flowering plants, the SOS1 transporters also play a vital role in pumping Na⁺ out of root cells.

Finally, absorbed salt ions (especially Na⁺) can be partitioned into the vacuole via transporters such as NHX1, which is localized in the tonoplast. On the one hand, vacuoles act as the ultimate Na⁺ sinks to reduce Na⁺ accumulation in cytoplasm. On the other hand, the water potential of root cells is also reduced to ensure water uptake in saline environments (Liang et al., 2015; Guo et al., 2020).

**Adaptation to Salinity by Shoots**

Under saline conditions, the response mechanisms triggered are different in different ornamental species. In some ornamental species with salt secreting structures like salt glands or salt bladders, especially Limonium species (Feng et al., 2014b; Leng et al., 2018; Lu et al., 2020; Mi et al., 2021), excess salt ions transported to aboveground parts of the plant are also excreted outside of the plant body through salt glands (García-Caparrós et al., 2016). Limonium species have remarkable salt tolerance thanks to their typical salt excreting salt glands in the leaves; they can grow and develop well in environments containing NaCl concentrations of more than 200 mM NaCl (Soudid et al., 2016; Mi et al., 2021). Besides excreting excess salt ions out of the leaves, some of the absorbed salt ions are regionalized; they accumulate in vacuoles and regulate the osmotic potential of leaf cells (González-Orenga et al., 2021). Therefore, more ornamental flowering plants that possess these leaf structures, and which also have high value, should be exploited further.

Some ornamental plants, such as Portulaca (Borsai et al., 2020), as well as Aloe vera (L.), exhibit leaf succulence when they grow in saline or drought environments (García-Caparrós et al., 2016). In response to salt stress, large amounts of ions were accumulated in the vacuoles of leaf cells in such plants, and the leaf succulence was subsequently displayed higher ability to cope with the salt stress. And the specific anatomical adaptation to salinity in halophytic ornamental flowering plants is summarized in Supplementary Figure S1.

To respond to saline conditions, ornamental plants have evolved many other adaptations; for example, altering leaf function, reducing chlorophyll content, and reducing stomatal size and aperture (García-Caparrós and Lao, 2018). In carnation (Dianthus caryophyllus), salt tolerance was mainly associated with the reduction of water loss by reducing stomatal conductance and increasing epicuticular wax (Kwon et al., 2019). Chrysanthemum morifolium treated with different levels of salinity displayed salt tolerance, with increased photosynthetic pigments, and a better seedling growth (Vanalruati et al., 2019). Photosynthetic pigments are vital in photosynthesis. Under stress, inhibited photosynthetic efficiency in plants was related to reduce chlorophyll content and damaged photosynthetic systems (Wu et al., 2016). Some plants showed higher tolerance to salinity in photosynthetic processes. For example, low energy dissipation rates and high light harvesting efficiencies were detected in Aster tripolium L. plants exposed to even 250 mM NaCl, with increased density of reaction centers (RC), enhanced electron transport ability, and lower energy dissipation rates (Duarte et al., 2017). In the ornamental plant Eugenia myrtifolia L., when treated with salinity, the normal photosynthetic system was protected by limiting salt ion accumulation in leaves, as well as by increasing chlorophyll content (Acosta-Motos et al., 2017). The salt tolerance of Aloe vera associated with high efficiency of photosystem II and ROS scavenging was enhanced by exogenous application of proline (Nakhaie et al., 2020). With the application of exogenous GB or proline, higher chlorophyll and K⁺ content was detected in Viburnum lucidum L. and Callistemon citrinus Stapf (Cirillo et al., 2016), as well as calendula (Calendula officinalis L.) plants (Soroori et al., 2021). Proline content and the stability of the cellular membrane were increased by exogenous application of ascorbic acid on calendula plants under salinity stress (Azizi et al., 2021).

The salt tolerance of ornamental flowering plants could also be enhanced, and the salt content reduced, by shedding old leaves. For example, older leaves are shed to reduce Na⁺ content in Kalanchoe blossfeldiana Poelln. In Gazania splendens Lem sp., multiple strategies are employed, such as accumulating the majority of Na⁺ and Cl⁻ in the roots, secreting salt ions out of leaves through salt glands, increasing the succulence index of living leaf tissue, and shedding old leaves (García-Caparrós et al., 2016). Thus, G. splendens is suitable for planting in saline environments. The adaptive strategies of shoots are summarized in Figure 1D.

**MOLECULAR BREEDING FOR IMPROVEMENT OF SALT TOLERANCE IN ORNAMENTAL PLANTS**

Salinity is one of the most important adverse factors limiting agricultural production. Cultivating salt-tolerant ornamental crops or flowers is considered to be an effective way of utilizing saline soils. Therefore, selecting and breeding salt-tolerant ornamental flowering crops is a first and important step for the sustainable development of saline land. Many salt-tolerant plants can survive in heavily saline soils, and current research demonstrates the benefit of selecting and breeding species that are both ornamental and highly salt tolerant. And new varieties with higher salt tolerance could be obtained through gene mutation, such as via gamma irradiation. Seedlings with enhanced salt tolerance, enhanced osmotic regulation ability, and greater ROS scavenging capacity were obtained via gamma irradiation in sweet osmanthus (Geng et al., 2019).

To survive in a saline environment, plants—on the one hand—reduce the translocation of toxic ions to aboveground parts of the plant by draining salt ions to the root extracellular spaces. On the other hand, they accumulate excess salt ions into the vacuoles. Thus, the cytoplasmic ion content is maintained in controlled levels, the osmotic potential of plant cells is reduced, and normal water uptake and physiological metabolism can occur (Chen et al., 2010; Song and Wang, 2015). In this process, many genes are upregulated to ensure recruitment of proteins that participate in ion uptake, translocation, and redistribution. This, in turn, ensures that salt ion concentrations in plant cells are reduced to a minimum (Yang et al., 2017). If plants can
successfully redistribute ions, the plant will survive; if the plant cannot successfully redistribute salt ions well, then plant growth will be inhibited, or the plant might even die, once the salt content exceeds the plant's level of tolerance. Therefore, the ability of a plant to retain K⁺ and balance Na⁺ in the cytoplasm is one of the most important indicators of salt tolerance in plants. Research finds that the ability to retain K⁺ is particularly critical (Reddy et al., 2004; Caverzan et al., 2016; Wang et al., 2019b,c). Maintenance of intracellular ionic homeostasis in Arabidopsis thaliana depends primarily on AtNHX1 (an Na⁺/H⁺ antiporter gene located in the tonoplast) and AtSOS1 (an Na⁺/H⁺ antiporter gene located in plasma membrane; Zhu, 2003). Furthermore, the mechanism responsible for regulating salt ions in most plants is the same as in A. thaliana. Interestingly, species with high salt tolerance, such as halophytes, can better adjust or compartmentalize ions than Arabidopsis (Song, 2009; Guo et al., 2018, 2019a). Perhaps, higher expression levels of related genes are activated in such plants. As in Arabidopsis, increased Na⁺ efflux and reduced intracellular Na⁺ content was detected in transgenic plants overexpressing AtSOS1 and AtHKT1 genes (Wang et al., 2014, 2019a). In the important ornamental plant Chrysanthemum crismum, salt tolerance in CcSOS1 over-expressing plants was enhanced, with a significant reduction in Na⁺ content and a favorable K⁺/Na⁺ ratio. This plant could grow under 200 mM NaCl conditions (An et al., 2014). Overexpression of CmPIP1 and CmPIP2 genes from Chrysanthemum morifolium in transgenic chrysanthemum plants also resulted in salt tolerance (Zhang et al., 2019). And the salt tolerance was improved by overexpressing LiNHX from Iris lactea (Guo et al., 2020). In the halophyte Aeluropus lagopoides, the promoter of the gene encoding the high-affinity potassium transporter AtHKT2;1 was involved in the response to salt stress in plants (Dave et al., 2021).

As well as the functional genes involved in ion transport during salt tolerance in ornamental plants, transcription factor genes are also involved in salt tolerance; this lays the foundation for further research into salt-tolerant mechanisms in ornamental plants. Novel varieties with high salt tolerance could be obtained using transgenic transcription factor genes. For example, the salt tolerance of transgenic chrysanthemum was enhanced by overexpressing the DgNAC1 or DgWRKY4 transcription factor genes from chrysanthemum (Dendronthema grandiform; Wang et al., 2017a,b). Salinity tolerance of Arabidopsis was conferred by overexpressing the PSK1 gene (a homologous gene of S-phase kinase-associated protein 1-like) from Paeonia suffruticosa, indicating that this gene is important for salt tolerance, together with the function of flowering in peony (Hao et al., 2017). The salt tolerance of chrysanthemum (Dendronthema grandiform) transgenic plants was improved with overexpression of the transcription factor gene DgNAC1 (a salt responsive gene); higher activities of SOD, POD, and CAT were detected in the transgenic plants compared with wild type (Wang et al., 2017b). Thus, this could provide evidence for the molecular modification of salt tolerance in ornamental plants. Genes known to confer salt tolerance to ornamental flowering plants are summarized in Table 2.

TABLE 2 | Genes involved in salt tolerance of salt-tolerant ornamental flowering plants.

| Gene name | Species | Accession number | Probable function | References |
|-----------|---------|------------------|-------------------|------------|
| LfSOS1    | Leptochloa fusca | KC525946         | Encoding a cytomembrane Na⁺/H⁺ antiporter that transport Na⁺ out of the plant cells | Mohammadi et al., 2019 |
| CcSOS1    | Chrysanthemum crismum | AB439132      | Encoding a cytomembrane Na⁺/H⁺ antiporter that transport Na⁺ out of the plant cells | An et al., 2014 |
| CmSOS1    | Chrysanthemum morifolium | KP896477     | Encoding a cytomembrane Na⁺/H⁺ antiporter that transport Na⁺ out of the plant cells | Gao et al., 2016 |
| AjSOS1    | Artemisia japonica | KP896475        | Encoding a cytomembrane Na⁺/H⁺ antiporter that transport Na⁺ out of the plant cells | Gao et al., 2016 |
| CcSOS1    | Cossostephium chinense | KP896476     | Encoding a cytomembrane Na⁺/H⁺ antiporter that transport Na⁺ out of the plant cells | Gao et al., 2016 |
| LfNHX1    | Leptochloa fusca | JF833902         | Encoding a tonoplast Na⁺/H⁺ antiporter that transport Na⁺ into vacuole | Mohammadi et al., 2019 |
| HtNHX1    | Helianthus tuberosus | EF159151      | Encoding a tonoplast Na⁺/H⁺ antiporter that transport Na⁺ into vacuole | Zeng et al., 2018 |
| HtNHX2    | Helianthus tuberosus | DQ343304      | Encoding a tonoplast Na⁺/H⁺ antiporter that transport Na⁺ into vacuole | Zeng et al., 2018 |
| IbNHX2    | Ipomea batatas | JN888961        | Encoding a tonoplast Na⁺/H⁺ antiporter that transport Na⁺ into vacuole | Wang et al., 2016 |
| lInHX     | Iris lactea | AY730277        | Encoding a tonoplast Na⁺/H⁺ antiporter that transport Na⁺ into vacuole | Guo et al., 2020 |
| AtHKT2;1  | Aeluropus lagopoides | MW535306      | Encoding a transporter recirculating Na⁺ from shoot to root | Dave et al., 2021 |
| PaPSK1    | Paeonia suffruticosa | FE529999      | Encoding a kinase-associated protein involving in response to salt tolerance | Hao et al., 2017 |
| DgNAC1    | Dendronthema grandiform | HX317452     | NAC transcription factor gene that response to salinity | Wang et al., 2017a |
| DgWRKY4   | Dendronthema grandiform | KC615358     | WRKY transcription factor gene that response to salinity | Wang et al., 2017b |
| CmPIP1    | Chrysanthemum morifolium | KJ489416     | Encoding aquaporin involving in salt tolerance | Zhang et al., 2019 |
| CmPIP2    | Chrysanthemum morifolium | KJ756774     | Encoding aquaporin involving in salt tolerance | Zhang et al., 2019 |
The ornamental flower *Alternanthera bettzickiana* (Regel) G. Nicholson grew under saline conditions, with less Na⁺ content in shoots than roots, even at 40 dS m⁻¹, so was considered to be a potential halophyte (Ali et al., 2012). Plants with high salt tolerance, such as *Portulaca grandiflora* and *Iris halophila* (Pall.) plants, which exhibit high salt tolerance during the growth period (Borsai et al., 2020), could be used as greening species on saline soils. Selection and utilization of such species could enable further study of salt tolerance mechanisms at the molecular level (Liu et al., 2018). Systematic genomic analysis to identify salt tolerance-related genes or transcription factors will be of great significance in plant breeding (Wang et al., 2021). Many transcription factor genes and functional genes might serve as candidate genes for breeding salt-tolerant ornamental flowering plants.

**ACQUISITION OF NOVEL VARIETIES BY GENETIC MODIFICATION**

The advent of genome editing technology, especially the CRISPR editing system, has provided an efficient method to obtain improved flower varieties. The genome sequences of some ornamental crops are available, such as *Nankingense*, *Chrysanthemum*, *Rosa chinensis*, *Ipomoea nil*, *Petunia hybrid*, and *Dendrobium officinale*. These genomes will provide the basis for obtaining and developing new varieties of ornamental crops, such as early flowering varieties of orchid and chrysanthemum (Ahmad et al., 2021). Using CRISPR/Cas9 genome editing technology, more flower colors were obtained in *Lilium* (Yan et al., 2019). In *Toreinia fournieri* L., CRISPR/Cas9 modification of a gene related to flavonoid synthesis resulted in flowers with a pale purple and almost white color (Nishihara et al., 2018). Flower longevity was extended in *Petunia hybrida* by knocking out the gene for 1-amino-cyclopropane-1-carboxylate oxidase 1 (ACO), which reduced the production of ethylene (Xu et al., 2020). Altered flower color (influenced by biosynthesis of anthocyanin) was also obtained in *Ipomoea nil* (Japanese morning glory) using CRISPR/Cas9 (Watanabe et al., 2017).

Further work based on the genome sequencing of ornamental plants (Wani et al., 2020) will lay the foundation for subsequent gene editing to obtain highly salt-tolerant ornamental flowers for greening saline–alkaline areas. Transcriptome analysis of ornamental plants will also reveal previously unknown genes related to flower color or quality, and genes related to the response to a saline environment (Guo et al., 2019b). Current methods of obtaining novel varieties are shown in Figure 2.

![Figure 2](image-url)
CONCLUSION AND PERSPECTIVES

Saline soils are widely distributed around the world, and these are considered to be a valuable land resource for development. Protection by vegetation and “greening” is the most effective and feasible biological methods for improving saline lands. Photos of some ornamental plants with beautiful flowers are shown in Figure 3. Planting salt-tolerant ornamental flowering
crops is a feasible and sustainable greening strategy. Achieving this goal relies on selecting and breeding ornamental flowering crop varieties with both high salt stress tolerance and high value.

Like most plants in a salinized environment, the responses or adaptive strategies of ornamental flowering crops to salt aim to mitigate the effects of osmotic, ionic, and oxidative stresses, as well as nutritional imbalances induced by salinity. Under saline conditions, ornamental plants accumulate osmolytes like proline, redistribute ions, enhance their ROS scavenging ability, reduce the salt ion uptake by roots, and secrete salt ions from leaves.

With the help of genetic engineering approaches such as genetic modification and gene editing, future studies should focus on understanding the salt tolerance mechanisms of ornamental flowering crops at the molecular level, and on providing the basis for breeding ornamental flowering crops with high salt tolerance, more flower colors, and greater economic value. Other strategies include selection and acceptance of model ornamental flowering plants with high salt tolerance; genome sequencing of model plants; and establishment of highly efficient transgenic systems for model plants. These strategies would form the basis for deciphering the salt-tolerant mechanisms of ornamental plants at the molecular levels; for example, they could provide insight into the role of the apoplastic barrier in ornamental plant roots for saline tolerance. Novel ornamental plant cultivars with high salt tolerance and expanded usages will be obtained using modern biological techniques.

**AUTHOR CONTRIBUTIONS**

JG and BW conceived and designed this study and revised the manuscript. JG, CS, and YifanZ wrote the manuscript.

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