RESEARCH ARTICLE



Genome-wide identification and expression analysis of $Na^+/$ H^+ antiporter (NHX) genes in tomato under salt stress

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Abstract

Plant Na^+/H^+ antiporter (NHX) genes enhance salt tolerance by preventing excessive Na⁺ accumulation in the cytosol through partitioning of Na⁺ ions into vacuoles or extracellular transport across the plasma membrane. However, there is limited detailed information regarding the salt stress responsive SINHXs in the most recent tomato genome. We investigated the role of this gene family's expression patterns in the open flower tissues under salt shock in Solanum lycopersicum using a genomewide approach. A total of seven putative SINHX genes located on chromosomes 1, 4, 6, and 10 were identified, but no ortholog of the NHX5 gene was identified in the tomato genome. Phylogenetic analysis revealed that these genes are divided into three different groups. SINHX proteins with 10-12 transmembrane domains were hypothetically localized in vacuoles or cell membranes. Promoter analysis revealed that SINHX6 and SINHX8 are involved with the stress-related MeJA hormone in response to salt stress signaling. The structural motif analysis of SINHX1, -2, -3, -4, and -6 proteins showed that they have highly conserved amiloride binding sites. The protein-protein network revealed that SINHX7 and SINHX8 interact physically with Salt Overly Sensitive (SOS) pathway proteins. Transcriptome analysis demonstrated that the SINHX2 and SINHX6 genes were substantially expressed in the open flower tissues. Moreover, quantitative PCR analysis indicated that all SINHX genes, particularly SINHX6 and SINHX8, are significantly upregulated by salt shock in the open flower tissues. Our results provide an updated framework for future genetic research and development of breeding strategies against salt stress in the tomato.

KEYWORDS

gene expression, Na⁺/H⁺ antiporter, NHX, Solanum lycopersicum, salt stress

INTRODUCTION 1

Salt stress is considered one of the most important abiotic stress factors that negatively affects plant growth and development and causes serious yield losses in important crop species (Deinlein et al., 2014; Nataraja & Parvathi, 2016). Global warming has accelerated as well as expanded the land that limits the planting of important crop species due to high levels of salinity, which are thought to affect more than

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6% of existing farmland and about 20% of irrigated land around the world (Munns & Tester, 2008). The adverse effects of salt can vary, depending on the developmental stages of a plant such as germination, early growth, and flowering or seeding stages (Chandna et al., 2013). Salt stress could affect essential biological processes including photosynthesis, water relationship, and nutrient uptake (Parihar et al., 2015). High concentrations of salt in the soil initially cause slower growth by reducing leaf and root growth, and then it becomes toxic to plant cells, leading to physiological disorders and cell death (Munns, 1993; Parihar et al., 2015). Plants under salt stress conditions accumulate high concentrations of Na⁺ and Cl⁻ ions in the chloroplasts, inhibiting photosynthesis and decreasing chlorophyll content (Chutipaijit et al., 2011; Zhang et al., 2005). Over time, increased Na⁺ concentrations in plant tissues become toxic, and this toxicity directly affects intracellular K⁺ homeostasis (Shabala & Cuin, 2008). A critical Na^+/K^+ ratio must be reached in the cell for effectiveness of many cytosolic enzyme activities (Mahaian et al., 2008). With an increase in the amount of Na⁺ in the external environment, the entry of Na⁺ ions into the cell also increases, whereas the uptake of K^+ ions into the cell decreases, and accordingly, the Na⁺/K⁺ balance is disturbed. This is because Na⁺ competes with K^+ for the sites where the K^+ ion binds (Tester & Davenport, 2003). Because K^+ ions play a direct in activating enzymes and to neutralizing negative charges of proteins, cells need high concentrations of K⁺ ions for the stabilization of protein synthesis (Harrewijn, 1979; Marschner, 1995; Pandolfi et al., 2012). Furthermore, some plants show a positive correlation between K⁺ content and yield (Ashraf & McNeilly, 2004; Bandeh-hagh et al., 2008; Valiollah. 2013).

To date, it has been demostrated that many ion transporters play important roles in maintaining the pH and ion homeostasis of plant cells in the presence of salt stress (Hamamoto et al., 2015; Pardo et al., 2006; Ward et al., 2009; Yamaguchi et al., 2013). The Na⁺/H⁺ antiporter (NHX) gene family, a subfamily of ion transporters, plays an important role in salt stress tolerance by controlling cellular pH and Na⁺ ion balance in plants (Brett et al., 2005; Van Zelm et al., 2020). The first plant NHX1 gene was identified in Arabidopsis thaliana (Gaxiola et al., 1999). The NHX gene family in A. thaliana includes eight members, which are divided into three subgroups (Brett et al., 2005). The AtNHX1 (A. thaliana NHX1), -2, -3, and -4 are referred to as Vac class NHXs on the vacuole membranes (Aharon et al., 2003), whereas AtNHX5 and AtNHX6, which are in the Endo class of NHXs, are located on the endosomal region (Bassil et al., 2011). The amiloride binding domain (FFIYLLPPI) is located at the N-terminal of TM3 (transmembrane-3), and this domain, known as an inhibitor of NHX activity, is a characteristic feature of vacuolar class NHX members (Counillon et al., 1993; Yamaguchi et al., 2003). The AtNHX7 and AtNHX8 are found in the plasma membranes (PMs) and are referred to as PM class (Shi et al., 2000). AtNHX7 (AtSOS1) is regulated by Serine/threonine Protein Kinase (SOS2) and Calcineurin B-like Calcium-Binding Protein (SOS3) in the Salt Overly Sensitive (SOS) signaling pathway (Qiu et al., 2002). In response to salt stress, intracellular Ca²⁺ increases, which triggers the activation of the SOS signaling

pathway (Manishankar et al., 2018). SOS3 activates the SOS2 protein kinase, and the SOS2-SOS3 complex then activates SOS1/NHX7 by phosphorylation to efflux Na⁺ ions out of the cell (Halfter et al., 2000; Liu et al., 2000; Shi et al., 2003). Previous reports have documented six NHX genes in Medicago truncatula (Sandhu et al., 2018), nine NHX genes in Capsicum annuum (Luo et al., 2021), five NHX genes in Beta vulgaris (Wu et al., 2019), six NHX genes in Vitis vinifera (Ayadi et al., 2020), 10 NHX genes in Glycine max (Chen et al., 2015), eight NHX genes in Populus trichocarpa (Tian et al., 2017), 10 NHX genes in Punica granatum (Dong et al., 2021), and eight NHX genes in Actinidia chinensis were identified (Liu et al., 2023). Such variation among the NHX gene numbers of plant species indicated that gene duplications or loss events occurred during species evolution, which also provided an opportunity to expand the NHX gene family by generation functionally divert new genes (Huang et al., 2022).

As a member of the Solanaceae family, tomato (Solanum lycopersi*cum*) is diploid with 2n = 24 chromosomes (Díez & Nuez, 2008). It is highly cultivated worldwide and contains invaluable components for human nutrition such as lycopene, β-carotene, and vitamin C (Clinton, 2005). Tomato plants exhibit premature cell senescence. accumulation of Na⁺ in the leaves, and a decrease in photosystem II efficiency when exposed to salt stress (100-mM NaCl) (Ghanem et al., 2012). It has been reported that both leaf area and dry matter content as well as K⁺/Na⁺ ratio of tomato plants decreased along with increased salt stress (Babu et al., 2012). Salt stress during the inflorescence development increases flower abortion of tomato plants and causes a decrease in pollen number and viability (Ghanem et al., 2009). Moreover, salt stress at the flowering stage has a direct adverse effect on yield by reducing the number of tomato fruits (Zhang et al., 2017). Because the importance of the NHX family in regulating salt tolerance is well established, the roles of several NHX members in the tomato under salt stress have been reported (Baghour et al., 2019, 2023; Gálvez et al., 2012; Huertas et al., 2013; Maach et al., 2020, 2021; Olías et al., 2009). Gálvez et al. (2012) reported that LeNHX1, -2, -3, -4 genes are generally induced by salt treatment in root, stem, and leaf tissues of salt-sensitive and salttolerant tomato species, and these isoforms are involved in Na⁺ ion accumulation in the aerial parts of the plant. Overexpression of the LeNHX2 (Huertas et al., 2013) and LeNHX4 (Maach et al., 2020) genes in transgenic tomato plants has been reported to improve salinity tolerance. In the tomato, gene expression analyses of tomato NHX genes under salt stress in previous reports were mainly limited to root, leaf, or stem tissues, and there were also no reports focusing on determining the expression patterns of NHX genes in flower tissues under salt stress. Recently, Hussain et al. (2022) have reported the identification of seven NHX genes in the tomato using a genome-wide approach. Herein, we have discussed and reviewed the findings of Hussain et al. (2022) related to the SINHX genes in the tomato.

In the present study, we identified members of the NHX gene family based on a comprehensive genome-wide approach using the current tomato genome and used bioinformatics tools to reveal phylogenetic relationships, synteny analysis, motif analysis, promoter analyprotein-protein interaction (PPI), and gene sis. structures.

Furthermore, the gene expression patterns of the identified *SINHX* genes were also analyzed by RT-qPCR in the open flower tissues obtained at 0, 6, 12, and 24 h after the plants were treated with 240-mM NaCl shock.

2 | MATERIALS AND METHODS

2.1 | Identification and characterization of the *NHX* genes in tomato genome

To identify putative NHX genes in tomato, the Pfam ID (PF00999) number belonging to the NHX gene family was searched in the Phytozome database (Goodstein et al., 2012). In addition, keywords related to NHX genes, such as sodium/hydrogen exchanger, and Na⁺/H⁺ antiporterl were also searched in the Sol Genomics Network (Fernandez-Pozo et al., 2015). The presence of the NHX domain genes was also confirmed by the Hidden Markov Model (Cook et al., 2018). The peptide sequences of the hypothetical tomato NHX genes were aligned to the sequences of the NHX genes in Arabidopsis, and the percentage of identity between the hypothetical tomato NHX genes and the AtNHX genes was determined. The Arabidopsis Information Resource (TAIR) database was used to retrieve Arabidopsis NHX protein sequences (Berardini et al., 2015). The isoelectric point (pl) and molecular weight (MW) of the SINHX genes were calculated with the ExPASy tool (Gasteiger et al., 2003). Intracellular localization of SINHX genes was predicted with the Plant-mPLoc server (Chou & Shen, 2010). The peptide sequences of the SINHX genes were screened with the TMHMM 2.0 web-based tool to confirm their transmembrane helix domains (Krogh et al., 2001).

2.2 | Multiple sequence alignment and phylogenetic analysis

The amino acid sequences of *NHX* genes of tomato and selected species (A. *thaliana*, G. *max*, M. *truncatula*, and V. *vinifera*) were used for multiple sequence alignment with the MUSCLE algorithm, and a phylogenetic tree was created using MEGA 11 software (Tamura et al., 2021). A thousand replicates were used to determine the bootstrap value. The phylogenetic tree was visualized via iTOL (Letunic & Bork, 2021).

2.3 | Chromosomal localization and synteny analysis

Sol Genomics Network database was used to visualize and to locate *SINHX* genes on the chromosomes of tomato genome (Fernandez-Pozo et al., 2015). Phytozome and Ensembl databases were used to determine the genetic relationship of *S. lycopersicum* with *C. annuum*, *S. lycopersicum*, and *A. thaliana* (Cunningham et al., 2022). Fasta and GFF3 files were downloaded from related databases and analyzed

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2.4 | Conserved motif and gene structure analysis of *SINHX* members

The Multiple Expectation Maximization for Motif Elicitation (MEME) Suite tool was used to identify conserved motifs in the amino acid sequences of the *SINHX* gene family (Bailey et al., 2015). The TBtools software's Gene Structure View was used to show schematic representation of the *SINHX* gene structure.

2.5 | Calculation of *Ka/Ks* and *cis*-regulatory element analysis

The *Ks* (synonymous) and *Ka* (non-synonymous) substitution of each duplicated gene pair were calculated using the Computational Biology Unit database (Siltberg & Liberles, 2002). The upstream 1500-bp region of the *SINHX* genes was selected for analysis of *cis*-regulatory elements on the promoters and predicted using the PlantCARE tool (Lescot et al., 2002).

2.6 | Transcriptome profiling of SINHX genes

The TomExpress RNA-Seq platform provided transcriptome data for the "Microtom" cultivar's roots, flowers, leaves, and flower buds tissues (Zouine et al., 2017). The results were displayed by using TBtools software after the data values were obtained as the normalized mean count per base of each *SINHX* gene.

2.7 | Three-dimensional (3D) structure analysis and PPI of SINHX proteins

The 3D structures of SINHX proteins were created by using the I-TASSER tool (Yang & Zhang, 2015). The outputs of the I-TASSER modeling results were visualized by using the RCSB Protein Data Bank 3D Viewer (Berman et al., 2000). The prediction of the PPI network of SINHX proteins was performed by using the STRING database (Jensen et al., 2009).

2.8 | Growth conditions of plant material and salt stress treatment

Tomato seeds (S. *lycopersicum* L., cv. Microtom) were sown on wet filter paper in plastic dishes and allowed to germinate in the dark at 25° C for 1 week. Germinated seeds were transferred to pots (125×73 mm) containing peat soil and perlite (3:1) in a plant growth

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TABLE 1 The sequences of Forward (F) and Reverse (R) primers used in RT-qPCR reactions.

Accession number	Gene name	Primer sequence	Product size (bp)
Solyc01g067710	SINHX1	F: GGCTTACCTATCTTACATGCTTGC R: AGCTCTCAGTCACATTATGCC	gDNA 244 cDNA 113
Solyc01g098190	SINHX2	F: TCCTCTTCCTCTATGTGGGCA R: AAACAAAAGCAGCTCTCCCCA	gDNA 314 cDNA 135
Solyc10g006080	SINHX3	F: CTCAGTGGGATTTTGACCGTC R: CAATGTCCAACGCATCCATCC	gDNA 522 cDNA 169
Solyc06g008820	SINHX4	F: ACTGATCGTGAAGTTGCTCTC R: TGCCAGGTATAGTGTGACATG	gDNA 209 cDNA 128
Solyc04g056600	SINHX6	F: TCTTGTACGACCTCCACACC R: GGACTGACTGCAAAGCAAGG	gDNA 192 cDNA 107
Solyc01g005020	SINHX7	F: CCTGGCGTGCTTATTTCCAC R: CCCAATTTCTTGCTGGCACC	gDNA 242 cDNA 167
Solyc04g018100	SINHX8	F: CTTTTGCTTGCTGGACCTGG R: ACAGCCACAGGATCAGTAGC	gDNA 209 cDNA 140
Solyc07g025390	EXPRESSED (housekeeping)	F: GCTAAGAACGCTGGACCTAATG R: TGGGTGTGCCTTTCTGAATG	gDNA 291 cDNA 183

chamber with $25 \pm .5^{\circ}$ C temperature, $65 \pm 5\%$ humidity, and 18/6 h (day/night) photoperiod (350μ mol m⁻² s⁻¹). The commercial liquid fertilizer (Black Diamond) was applied with irrigation as needed while no pesticide was used. Because the toxic effects of salt were not observed in the open flower tissues in our preliminary experiments using different concentrations of NaCl (60, 120, and 180 mM), the 240-mM NaCl salt shock was applied to the plants as at the flowering stage. Furthermore, previous reports support the suitability of high salt concentration in the tomato Microtom cultivar (Bacha et al., 2017). Open flower tissues were collected from three biological replicates at 0 (control), 6, 12, and 24 h after salt shock treatment and were immediately stored at -80° C until total RNA extraction.

2.9 | Total RNA isolation and cDNA synthesis

Total RNA was isolated from open flower tissues by using the GF-1 Total RNA Extraction Kit, followed by DNase treatment to eliminate genomic DNA contamination. The quality of the isolated RNA was checked with both gel electrophoresis and a NanoDrop One (Thermo ScientificTM NanoDropTM One Microvolume UV–Vis Spectrophotometers) instrument. The first-strand cDNA was synthesized by using RevertAid First Strand cDNA Synthesis Kit (Thermo ScientificTM).

2.10 | Real time-quantitative PCR (RT-qPCR)

RT-qPCR reactions were carried out using the Ampliqon RealQ Plus SYBR Green/ROX Master Mix Kit. Gene-specific forward and reverse primers for members of *SINHX* gene family were designed using Mac-Vector v18.2 software (MacVector Inc., Cary, NC, USA) (Table 1). The *EXPRESSED* gene was used as housekeeping for normalization in RTqPCR analyses (Choi et al., 2018). The RT-qPCR conditions: initial denaturation at 95°C for 8 min, 40 cycles of 95°C denaturation for 15 s, 55°C annealing for 30 s, and 72°C extension for 30 s. Three biological replicates were used for each sample. The relative expression levels of the *SINHX* genes were calculated by the $2^{-\Delta\Delta Ct}$ method (Livak & Schmittgen, 2001).

2.11 | Statistical analysis

GraphPad Prism 9.3 software (GraphPad Software Inc., San Diego, CA, USA) was employed to perform statistical analysis. The data were statistically analyzed using one-way analysis of variance (ANOVA), and the difference was considered significant when the p value was less than .05.

3 | RESULTS

3.1 | Identification of SINHX genes

The amino acid sequences of *AtNHXs* were used in the alignments to identify the *NHX* gene family in the tomato genome. The results revealed seven *SINHX* genes encoding the *NHX* in the tomato genome (Table 2), namely, *SINHX1* (Solyc01g067710), *SINHX2* (Solyc01g098190), *SINHX3* (Solyc10g006080), *SINHX4* (Solyc06g008820), *SINHX6* (Solyc04g056600), *SINHX7* (Solyc01g005020), and *SINHX8* (Solyc04g018100).

Sequence analyses revealed that genomic sequences of *SINHX* genes ranged from 4232 bp (*SINHX3*) to 15,997 bp (*SINHX6*), the number of exons in the coding sequence (CDS) ranged from 14 to 23, and the total length of CDS ranged from 1578 bp (*SINHX3*) to 3456 bp (*SINHX7*) (Table 2). The size of ORF ranged from 526 aa (*SINHX3*) to 1152 aa (*SINHX7*) (Table 2). In addition, the size of SINHX

TABLE 2 Some characteristics of tomato SINHX genes.



Gene name	Accession number	Genomic sequence (bp)	CDS (bp)	ORF (aa)	тм	Ch	NHX class	Subcellular localization	MW (kDa)	pl	Identity with NHX protein of Arabidopsis (%)
SINHX1	Solyc01g067710	4956	1614	538	10	1	I	Vacuole	59.4	8.55	72.5
SINHX2	Solyc01g098190	5857	1611	537	10	1	I	Vacuole	58.7	7.24	75.5
SINHX3	Solyc10g006080	4232	1578	526	11	10	I	Vacuole	59.1	8.48	69.9
SINHX4	Solyc06g008820	7602	1605	535	10	6	I	Vacuole	59.0	6.60	63.1
SINHX6	Solyc04g056600	15,997	1596	532	12	4	II	Vacuole	58.5	5.42	81.6
SINHX7	Solyc01g005020	13,405	3456	1152	12	1	III	СМ	127.5	5.89	64.4
SINHX8	Solyc04g018100	15,127	2952	983	10	4	Ш	CM-vacuole	108.2	5.75	49.8

Abbreviations: CDS, coding sequences; Chr, chromosomal location; CM, cell membrane; MW: molecular weight; ORF: open reading frame; pl: isoelectric point; TM: transmembrane domain.

FIGURE 1 Phylogenetic relationships among Na⁺/H⁺ antiporter (NHX) proteins of tomato and selected species, Solanum lycopersicum (SI), Arabidopsis thaliana (At), Medicago truncatula (Mt), Glycine max (Gm), and Vitis vinifera (Vv).



proteins ranged from 58.5 (kDa, SINHX6) to 127.5 (kDa, SINHX7), whereas isoelectric points (pl) varied from 5.42 (SINHX6) to 8.55 (SINHX1) (Table 2). SINHX proteins had transmembrane domains ranging from 10 to 12 (Table 2). The amino acid sequences of the *SINHX* genes were aligned with the amino acid sequences of the *AtNHX* genes, and the identity of the genes with the *AtNHX* genes was determined. The highest identity matrix score (81.6%) was determined between *SINHX6* and *AtNHX6* genes, whereas the lowest (49.8%) was between *SINHX8* and *ATNHX8* genes (Table 2).

3.2 | Phylogenetic analysis

To determine the evolutionary relationships of NHX proteins with some of other plant species, SINHXs were compared with A. *thaliana* (At, eight sequences), G. max (Gm, eight sequences), M. *truncatula* (Mt, six sequences), and V. *vinifera* (Vv, six sequences). These plant species were chosen due to their status as model plants and their representation in several plant families allowing the investigation of the evolution and diversity of NHX genes. A total of five species were used for



the analysis, as the results of the preliminary phylogenetic analysis showed that including more plant species in the phylogenetic analysis led to reduced interpretability. Phylogenetic analysis revealed that 35 SINHX proteins were divided into three classes based on their predicted subcellular localization, indicated by different colors using iTOL program (Figure 1). *SINHX1*, -2, -3, and -4 were localized on the vacuole membranes (Vac class/Class I); *SINHX6* was localized on the endosomal region (Endo class/Class II); and *SINHX7* and -8 were localized on the PM (PM class/Class III) (Figure 1).

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3.3 | Chromosomal localization, *Ka/Ks*, and synteny analysis

Location information of *SINHX* genes on the chromosomes was obtained from the tomato genome. Seven putative *SINHX* genes were scattered on 4 chromosomes. *SINHX3* gene was located on chromosome 10; genes *SINHX6* and *SINHX8* were located on chromosome number 4; *SINHX1*, -2, and -7 genes were located on chromosome 1, and *SINHX4* gene was located on chromosome 6 (Figure 2). There were no *SINHX* genes on chromosomes 2, 3, 5, 7, 8, 9, 11, and 12.

The formula $T = (Ks/2\lambda) \times 10^{-6}$ million years ago was used to estimate the date of duplication events ($\lambda = 1.5 \times 10^{-8}$) (Blanc & Wolfe, 2004; Madrid-Espinoza et al., 2019). According to the *Ka/Ks* ratio, the selection type is divided into three different groups as

purifying (Ka/Ks < 1), neutral (Ka/Ks = 1), and positive selection (Ka/Ks > 1) (Table 3) (Lynch & Conery, 2000). The term "tandem duplication" refers to the presence of two or more genes on the same chromosome, whereas the term "segmental duplication" refers to the presence of genes on different chromosomes (Table 3) (Akram et al., 2020).

To better understand the evolutionary relationships of the *SINHX* genes, comparative syntenic schemes were constructed between *S. lycopersicum*, *C. annuum*, and *A. thaliana* genomes. The *SINHX1*, *SINHX2*, *SINHX3*, and *SINHX7* genes showed collinear relationships with *A. thaliana* and *C. annuum* (Figure 3). *SINHX4*, *SINHX6*, and *SINHX8* genes did not show a synteny with the genomes compared (Figure 3).

3.4 | Gene structure and conserved motif analysis

The exons and introns of *SINHXs* genes were analyzed by using TBtools. Vac class *SINHX1*, -2, -3, and -4 genes provided 14 exons and 13 introns (Figure 4). Endo class *SINHX6* had 22 exons and 21 introns, whereas PM class *SINHX7* and *SINHX8* genes showed 23 exons and 22 introns and 22 exons and 21 introns, respectively (Figure 4). Among the *SINHXs* genes, *SINHX6* had the largest genomic sequence length at 15997 bp, whereas *SINHX3* had the shortest with



FIGURE 2 The chromosomal distribution of SINHX gene family members in the tomato genome.

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TABLE 3 *Ka/Ks* ratio, duplication, selection types, and divergence time of *SINHX* genes.

Gene pairs	Ка	Ks	Ka/Ks	Duplicated type	Selection type	Time (million year ago)
SINHX7-SINHX8	.183	.402	.456	Segmental	Purify	3.011
SINHX3-SINHX4	.132	.417	.316	Segmental	Purify	3.126
SINHX1-SINHX2	.068	.407	.166	Tandem	Purify	3.054

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Abbreviations: Ka, the number of nonsynonymous substitutions per non-synonymous site; Ks, the number of synonymous substitutions per synonymous site.



FIGURE 3 Comparative synteny analysis of Na^+/H^+ antiporter (NHX) genes in Solanum lycopersicum, Arabidopsis thaliana, and Capsicum annuum genomes. (a) Syntenic relationship among SINHX genes in S. lycopersicum and A. thaliana. (b) Syntenic relationship among SINHX genes in S. lycopersicum and C. annuum. The gray lines in the background represent the synteny pairs of the whole genome, whereas the red lines indicate the synteny of SINHX gene pairs and their corresponding positions on each chromosome.

4232 bp. The *NHX* domain (PF00999) was also confirmed by using the Hidden Markov Model (HMM).

Parameter settings related to motif analysis: the maximum number of motifs was 16, and the optimum aa width was set to 6–50 in length. The MEME tool was used to investigate the conserved motifs of *SINHX* genes, and the conserved motifs were determined in different colors (Figure 5a). Analysis results revealed that there were 10 conserved motifs in *SINHX* family members, and these conserved motifs ranged from 21 to 50 amino acids in length (Figure 5a). Whereas motif 1 was found in all members of *SINHX*, motifs 2, -3, -4, -5, -6, -7, and -9 were found in the members of *SINHX*1, -2, -3, and -4 (Figure 5a). Motif 10 was also found in members of *SINHX*6, -7, and -8, whereas Motif 8 was found in all members

except in *SINHX7* and *SINHX8* genes (Figure 5a). The amiloride binding site [FFIYLLPPI], which is a characteristic feature of NHX proteins, was fully conserved in *SINHX1*, -2, and -3 genes, whereas it mostly retained in *SINHX4* and *SINHX6* genes. The amiloride binding site was absent in *SINHX7* and *SINHX8* genes (Figure 5b).

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3.5 | Analysis of *cis*-regulatory elements of *SINHX* genes

Analysis of *cis*-regulatory elements in the 1500 bp upstream promoter region of *SINHX* genes was performed with the PlantCare tool (Figure 6). The results indicated that the *SINHX* genes family contained





FIGURE 4 The gene structure of Na^+/H^+ antiporter (NHX) family member in the tomato. CDS, coding sequence; UTR, untranslated region. The scale is given as bp.

a total of 502 cis-regulatory elements, and they were classified in a total of 32 types, which were mainly related to light-responsible elements. development-related elements, hormone-responsible elements, promoter-related elements, site-binding-related elements, and environmental stress-related elements (Figure 6). Of those, the CAAT and TATA-boxes of promotor-related elements were too common and were included an average of 12.14 and 55.71 times, respectively, in the defined promotor regions of all SINHX genes and did not provide a unique pattern as cis-regulatory elements, respectively (data not shown). On the other hand, the light-responsible elements included 3-AF1 binding site, AT1-motif, ATCC-motif, ATC-motif, ACE, AE-box, Box 4, chs-CMA1a, GA-motif, GATA-motif, G-Box, GT1-motif, G-box, I-box, LAMP-element, MRE, and TCT motif (Figure 6a). Developmentrelated elements included O2-site, CAT-box, and circadian motifs (Figure 6a,b). Hormone-responsible elements included ABRE, TGACGmotif, GARE-motif, TATC-box, and CGTCA-motif (Figure 6a,b). Sitebinding-related elements included AT-rich element, Box, and Unnamed_1 (60K protein binding site). Environmental stress-related elements included LTR, MBS, TC-rich repeats, and ARE (Figure 6a,b). The ABRE (ABA, the abscisic acid), TGACG-motif (MeJA), GARE-motif (GB, Gibberallin), TATC-box (MeJA), and CGTCA-motif (MeJA) were determined to be hormone-related cis-acting regulatory elements (data not shown). Of those elements, MeJA-related cis-acting regulatory elements (twice of TGACG-motif and CGTCA-motif and single TATC-box) were located on the SINHX6 (data not shown).

3.6 | PPI prediction

The PPI network was built by the STRING database using the K-means Clustering algorithm (network is grouped to a predefined number of clusters) to further investigate the probable role of SINHXs during possible interactions with other proteins. Based on the

clustering analysis, the network is divided into three clusters based on the clustering analysis. Cluster 1 (red Bubble) contains the proteins CBL10, SINHX7, CIPK, and SolycO6g051970; Cluster 2 (Green Bubble) contains the proteins SINHX1, SINHX2, SINHX4, SINHX6, SolycO7g014680, and SolycO8g065360; and Cluster 3 (Blue Bubble) contains the proteins SINHX3 and SINHX8. The analysis results showed that the proteins SINHX7 and SINHX8 hypothetically interacted with Calcineurin B-like (CBL) and CBL-interacting protein kinases (CIPK) (Figure 7). SolycO7g014680 (Putative High-affinity K⁺ transporter 1) interacted with all proteins in the network. In addition, SolycO6g051970 (Calcineurin B-Like Protein 4) also interacted with all proteins apart from CBL10 (Calcineurin B-like Protein 10) in the tomato genome network.

3.7 | Three-dimensional (3D) structure prediction of SINHX proteins

The 3D model structures of SINHX proteins were obtained by using the I-TASSER tool and visualized using the PDB 3D Viewer (Figure 8). The C-score was used to estimate the confidence of the models created in the analysis for SINHX proteins (Roy et al., 2010). The C-score is a confidence score for estimating the quality of predicted models by I-TASSER (Yang et al., 2013). The C-score generally varies in the range of [-5, 2], and the higher the score, the higher the reliability of the model (Yang et al., 2013). TM-score was proposed to scale for measuring the structural similarity between two structures, and it ranges from 0 to 1 and a value of 1 indicating a perfect match between the two constructs (Zhang, 2008). The root-mean-square deviation (RMSD) varies between 0 and 30 angstrom (Å) (Roy et al., 2010). The analysis results revealed that C-scores of the SINHX proteins ranged from -1.33 to .35 (Table 4). The modeling estimate of the SINHX4 protein gave the highest C-score at .35, whereas the SINHX7 and SINHX8 proteins had the lowest C-score at 1.33 (Table 4). The length of the NHX domain in SINHX proteins ranged from 418 (SINHX3) to 404 (SINHX6) amino acids (Table 4).

3.8 | RNA-Seq gene expression profiles of SINHX genes

The high-throughput gene expression data analysis was retrieved from the TomExpress database, and the Microtom cultivar was selected to determine the expression levels of tissue-specific *SINHX* genes. Seven *SINHX* genes and types of tissues including root, flower, leaf, and flower buds were selected to generate the heatmap using TBtools (Figure 9). According to expression patterns, all *SINHX* genes except the *SINHX1* gene were expressed in all tissues (Figure 9). The *SINHX2* gene was expressed at high levels in all tissues, whereas the *SINHX1* gene showed the lowest expression levels (Figure 9). In addition, the heatmap revealed that the *SINHX2* and *SINHX6* genes were highly expressed in flower tissue while the *SINHX3* and *SINHX1* genes showed the lowest expression levels.



FIGURE 5 Conserved motif analysis of *SINHXs* genes. (a) Conserved motifs of *SINHXs* genes, (b) motif 1 was found in all *SINHX* genes, and it had an amiloride binding site [FFIYLLPPI]. The scale showing the sequence length of the proteins was shown below (up to 1000 aa).

3.9 | Expression profiles of the SINHX genes under salt shock

Relative expression levels of SINHX genes were determined in the open flower tissues collected at the given time intervals (0th, 6th, 12th, 24th h) after 240-mM NaCl shock was applied. The expression of SINHX genes was normalized with the EXPRESSED (Solyc07g025390) gene (Figure 10). The RT-qPCR data indicated that salt shock significantly changed the relative expression levels of all SINHX genes in all given time intervals with some exceptions (Figure 10). For instance, salt shock significantly changed the relative expression levels of SINHX3, SINHX4, SINHX7, and SINHX8 genes in all given time intervals compared with control, whereas significant levels of the relative expression levels of SINHX1, SINHX2 and SINHX6 genes were various based on given time intervals (Figure 10). An increased relative expression level of the SINHX6 gene was delayed until the 12th hour of salt shock, whereas SINHX1 and SINHX2 genes showed early significant upregulations after 6 h of salt shock (Figure 10). Among SINHX gene family members, SINHX1, SINHX2, and SINHX8 genes were substantially upregulated in the open flower tissues at the 6th hour of salt shock, whereas significant downregulations were determined for the relative gene expressions levels of SINHX3 and SINHX7 genes at the same time interval (Figure 10). In contrast to the other SINHX gene family members, the relative expression levels of SINHX6, SINHX7, and SINHX8 genes were the highest at the 24th hour of salt shock (Figure 10).

4 | DISCUSSION

Abiotic environmental factors such as salinity directly affect plant growth and development and cause severe agricultural yield losses (Van Zelm et al., 2020). The *NHX* genes are significant in maintaining

Na⁺ ion homeostasis in plant tissues, and many reports have revealed that these genes provide salt tolerance to plants (Li et al., 2017). The availability of high-quality de novo genome assemblies and annotations (ITAG4.0) for the tomato reference genome (SL4.0) has opened up new opportunities for precision genome-wide studies in the tomato (Hosmani et al., 2019). In this study, a total of seven NHX genes were identified in the tomato, but the orthologue of the NHX5 gene was not identified in the tomato genome. Although the Solyc04g056600 accession showed identity to both AtNHX5 (80.6%) and AtNHX6 (81.6%) genes of Arabidopsis at the amino acid levels, it was defined as the SINHX6 gene in the present study because of its higher identity score (%81.6) and no presence of AtNHX5 gene orthology in the current tomato genome (Table 2). A similar case was also reported in the M. truncatula genome (Sandhu et al., 2018), suggesting that the NHX5 gene does not share a common ancestor with S. lycopersicum and M. truncatula genomes by speciation although orthologous genes retained the same function in the course of evolution. Single knockout mutants of the nhx5 and nhx6 genes in A. thaliana have shown unaltered salt sensitivity, whereas double knockout mutants of the nhx5 and nhx6 genes have exhibited greater salt sensitivity than the wild-type plant (Bassil et al., 2011). This suggests that the absence of the AtNHX5 gene orthologue in the tomato genome may not have a significant effect on the development of salt tolerance in the tomato (Sandhu et al., 2018). Although Hussain et al. (2022) previously reported seven SINHXs in the tomato genome as in the present study, the genes identified are completely different from each other. Interestingly, the Solyc00g021510 gene, which Hussain et al. (2022) identified as SINHX1, is not present in the current tomato reference genome (SL4.0). The orthologue of AtSOS1 in the tomato genome was identified as SISOS1 (CAG30524.1) by Olías et al. (2009), and this gene is identical to the SINHX7/SISOS1 (Solyc01g005020) gene identified in the present study.



FIGURE 6 *Cis*-regulatory elements in the 1500 bp upstream promoter region of *SINHX* genes. (a) Predicted *cis*-acting elements in the promoters of *SINHX* genes, (b) different colored histograms represent categorical *cis*-acting elements in *SINHX* genes.

Phylogenetic analysis revealed that 23 out of 35 NHX proteins and the majority of SINHX (SINHX1, -2, -3, and -4) proteins bind to vacuolar membrane, and the PM and Endo classes contained 6 and 8 proteins, respectively (Figure 1). A similar pattern of cluster was also reported for other plant species including cotton (Fu et al., 2020), sugar beet (Wu et al., 2019), and honeysuckle (Huang et al., 2022). *LeNHX2* was previously located in the Endo class (Class II), which includes *AtNHX5* and *AtNHX6* (Gálvez et al., 2012; Huertas et al., 2013; Rodríguez-Rosales et al., 2008; Venema et al., 2003), whereas the *SINHX2* member identified in this study was located in the Vac class (Class I), which includes *AtNHX1*, *AtNHX2*, *AtNHX3*, and *AtNHX4* (Figure 1). Similarly, *MtNHX2* in the *M. truncatula* genome (Sandhu et al., 2018), *BvNHX2* in the *B. vulgaris* genome (Wu et al., 2019), and *PgNHX2* in the *P. granatum* genome were also localized in Vac class (Dong et al., 2021). As previously reported for other plant species (Dong et al., 2021; Fu et al., 2020; Huang et al., 2022; Sandhu et al., 2018; Wu et al., 2019), the *SINHX* gene family was divided into three subgroups (Figure 1) although Hussain et al. (2022) provided two subgroups for the same gene family designated as N1 and N2. Moreover, Hussain et al. (2022) included the *SINHX1* and *SINHX2* genes in the same subgroups as *AtNHX7* and *AtNHX8*, which belong to the PM class.

The transmembrane domain numbers and amiloride binding domains of SINHX proteins were similar to most *NHX* family members



FIGURE 7 Protein-protein interaction (PPI) network of SINHXs. The lines (gray) represent the shared physical complex, and as the thickness of these lines increases, the confidence of interaction also increases. Each node represents all the proteins produced by a single, protein-coding gene locus. Empty nodes represent proteins with an unknown three-dimensional model. Filled nodes show that a three-dimensional model is known or predicted.



FIGURE 8 Three-dimensional structure prediction of seven SINHX proteins.

TABLE 4 Quantitative outputs of I-TASSER modeling for SINHX proteins and sequence location information of the Na+/H+ antiporter domain and amiloride binding sites.

Protein	C-score	TM-score	RMSD (Å)	Peptide sequence	Na ⁺ /H ⁺ antiporter domain (start to end)	Amiloride binding site (start to end)
SINHX1	23	.68 ± .12	8.0 ± 4.4	538	29-443	85-93
SINHX2	16	.69 ± .12	7.8 ± 4.4	537	29-442	85-93
SINHX3	.04	.72 ± .11	7.3 ± 4.2	526	27-444	81-89
SINHX4	.35	.76 ± .10	6.7 ± 4.0	535	27-442	84-92
SINHX6	16	.69 ± .12	7.8 ± 4.4	532	34-437	89-97
SINHX7	-1.33	.55 ± .15	12.5 ± 4.3	1,152	29-441	-
SINHX8	-1.33	.55 ± .15	12.5 ± 4.3	983	55-462	-

Abbreviation: RMSD, root-mean-square deviation.



FIGURE 9 RNA-Seq gene expression profiles of *SINHX* genes in different tissues.

of other species reported in the literature (Chen et al., 2015; Dong et al., 2021; Fu et al., 2020; Huang et al., 2022; Tian et al., 2017; Wu et al., 2019). The amiloride binding domain is either completely or highly conserved in plant species such as A. *thaliana*, M. *truncatula*, *P. granatum*, and *P. trichocarpa* (Aharon et al., 2003; Dong et al., 2021). The results of this study showed that the amiloride binding domain is fully conserved in *SINHX1-3* genes and highly conserved in *SINHX4* and *SINHX6* genes, whereas this site is absent in *SINHX7* and *SINHX8* genes (Figure 5b), which was in agreement with a previous report (Sandhu et al., 2018). In contrast, the *SINHX1* and *SINHX2* genes identified by Hussain et al. (2022) do not have an

amiloride binding domain, whereas, interestingly, *SINHX7* has this domain. Unlike the results reported by Hussain et al. (2022), the results of the present study were able to fully explore the amiloride binding domains of SINHX proteins.

The results of synteny analysis revealed that there are four homologous pairs between tomato and pepper and four homologous pairs between tomato and Arabidopsis (Figure 3). These results indicated that SINHX genes are phylogenetically related to NHX genes of different plant species. Generally, the Ka/Ks ratio of SINHX genes was lower than 1, indicating a selection pressure on protein coding sequences during the evolution (Table 3) (Hanada et al., 2007). Similar low Ka/Ks ratios were also reported for honeysuckle (Huang et al., 2022), radish (Wang et al., 2020) and cotton (Fu et al., 2020). The protein-protein network of SINHXs revealed that SINHX7 and SINHX8 interact physically with the CBL and CIPK proteins, unlike the other SINHX proteins (Figure 6). It was previously shown that the NHX7 gene, which provides tolerance by transporting Na⁺ ions under salt stress in plants, was activated as a result of the interaction between CBL and CIPK proteins (Weinl & Kudla, 2009). Comparably, the interaction of CIPK and CBL proteins with NHX7 was also predicted in plant species such as poplar, sugar beet, and pomegranate (Dong et al., 2021; Tian et al., 2017; Wu et al., 2019). These results revealed that SINHX7 and SINHX8 genes could play a more important role than the other SINHX genes in responses to salt stress in the tomato genome.

The results of RT-qPCR revealed that *SINHX6* and *SINHX8* genes showed significant and elevated upregulation by salt stress in the open flower tissues as the stress expanded to 24 h, and an early salt stress response (at 6th h) was provided by the *SINHX1* and *SINHX2* genes (Figure 10). These results were comparable with RNA-Seq gene expression profiles of *SINHX* genes determined in different tissues (Figure 9). A similar gene expression pattern of *SINHXs* could also be possible in other tissues at various developmental stages of tomato (Figure 10). In addition, presence of the higher number of MeJA-related *cis*-acting regulatory elements on the promotor regions of *SINHX6* (TATC-box, TGACG, CGTCA) and *SINHX8* (TGACG, CGTCA), compared with the other promotors of *SINHXs*, may also be attributed to the significant and elevated upregulation of these genes (Figure 10). These findings suggested that *SINHX* genes in *S. lycopersicum* might provide



FIGURE 10 Relative gene expression profiles of SINHX gene family members at different time intervals under 240-mM NaCl shock. Error bars show the standard error of three biological replicates mean (n = 3) (ns: no significant; *p < .05; **p < .01; ***p < .00].

orchestrated and time-dependent responses based on the duration and severity of salt stress (Figure 10). In the LeNHX isoforms induced by salt treatment, the greatest induction was in the LeNHX4 gene in the leaf (Gálvez et al., 2012), whereas in the present experiment, the greatest induction was in the SINHX8 gene in the open flower tissue (Figure 10). It has been previously reported that transgenic tomatoes overexpressing LeNHX2 have increased salt tolerance (Huertas et al., 2013). This gene corresponds to the SINHX6 gene, which was significantly upregulated at all time intervals after salt shock treatment in the present study (Figure 10). These findings point to the importance of the SINHX6/ LeNHX2 gene in improving salt stress tolerance in the tomato. Jabeen et al. (2022) reported that there was no change in the expression level of HvNHX1 in the leaf tissue of the Gairdner barley cultivar under 300-mM NaCl compared with the control. A similar finding was observed for the expression pattern of the SINHX1 gene at all time points in the open flower tissue, except at 6 h (Figure 10). A previous report also showed that all members of MtNHX genes were also upregulated under salt stress in flower tissues in M. truncatula (Sandhu et al., 2018) although some differences in terms of gene expression levels stand out in flower tissues of S. lycopersicum and M. truncatula (Sandhu et al., 2018). For instance, the SINHX7 gene was highly expressed in flower tissue after 24 h, whereas low expression profile of MtNHX7 gene was reported in flower tissue of M. truncatula (Sandhu et al., 2018). Conversely, the BvNHX5/BvSOS gene in B. vulgaris under salt stress was significantly upregulated in leaves (Wu et al., 2019). Moreover, it has been determined that PgNHX genes showed different functions depending on the tissue in pomegranate roots and leaves (Dong et al., 2021). Taken together, these results suggested that the expression patterns of orthologous genes of the NHX family in different plant species might differentially be tuned in response to salt stress based on severity of the stress and type of plant tissue.

CONCLUSION 5

In this study, a total of seven SINHX genes were identified, and the phylogenetic analysis provided three classes based on their subcellular localizations. Members of the SINHX family were grouped with SINHX1-4 on vacuole membranes, SINHX6 on the endosomal region, and SINHX7-8 on the PM. The amiloride binding site domain [FFIYLLPPI] was fully conserved in SINHX1-3, whereas this domain was highly conserved in SINHX4 and SINHX6. The cis-acting element analysis revealed that SINHX6 and SINHX8 were involved with the stress-related hormone MeJA in response to salt stress signaling. The PPI network analysis results indicated that the proteins SINHX7 and SINHX8 hypothetically interacted with the CBL and CIPK pathway in salt stress response. The RT-qPCR analysis showed that all SINHX genes in the open flower tissues were significantly upregulated under salt shock, although the expression pattern was time dependent. The results pointed out the importance of the SINHX genes in relation to salinity stress tolerance in the tomato. Our results suggest that the SINHX genes identified using the current tomato genome information provide novel reference genes for future studies on salt stress in the tomato.

AUTHOR CONTRIBUTIONS

I.T. conceived the idea, planned the experiments, interpreted the data, and designed the figures. U.S. and E.C. performed the bioinformatic analysis and RT-qPCR analysis and drafted the manuscript. All authors read and approved the manuscript.

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CONFLICT OF INTEREST STATEMENT

The authors declare that they have no conflict of interest.

DATA AVAILABILITY STATEMENT

Data sharing is not applicable to this article as no new data were created or analyzed in this study.

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