Impact of AtNHX1, a vacuolar Na⁺/H⁺ antiporter, upon gene expression during short- and long-term salt stress in Arabidopsis thaliana

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Abstract

Background: AtNHX1, the most abundant vacuolar Na⁺/H⁺ antiporter in Arabidopsis thaliana, mediates the transport of Na⁺ and K⁺ into the vacuole, influencing plant development and contributing to salt tolerance. In this report, microarray expression profiles of wild type plants, a T-DNA insertion knockout mutant of AtNHX1 (nhx1), and a 'rescued' line (NHX1::nhx1) were exposed to both short (12 h and 48 h) and long (one and two weeks) durations of a non-lethal salt stress to identify key gene transcripts associated with the salt response that are influenced by AtNHX1.

Results: 147 transcripts showed both salt responsiveness and a significant influence of AtNHX1. Fifty-seven of these genes showed an influence of the antiporter across all salt treatments, while the remaining genes were influenced as a result of a particular duration of salt stress. Most (69%) of the genes were up-regulated in the absence of AtNHX1, with the exception of transcripts encoding proteins involved with metabolic and energy processes that were mostly down-regulated.

Conclusion: While part of the AtNHX1-influenced transcripts were unclassified, other transcripts with known or putative roles showed the importance of AtNHX1 to key cellular processes that were not necessarily limited to the salt stress response; namely calcium signaling, sulfur metabolism, cell structure and cell growth, as well as vesicular trafficking and protein processing. Only a small number of other salt-responsive membrane transporter transcripts appeared significantly influenced by AtNHX1.

Background

The AtNHX1 gene encodes the most abundant vacuolar Na⁺/H⁺ antiporter in Arabidopsis thaliana, and mediates the transport of both K⁺ and Na⁺ into the vacuole [1,2]. Constitutive over-expression of AtNHX1 and homologues from other plants have been shown to confer significant salt tolerance in a variety of plant species as a result of increased vacuolar sequestration of sodium ions ([3], and references therein). The importance of AtNHX1 to salt stress tolerance was further demonstrated when T-DNA insertional mutant nhx1 'knockout' plants lacking a functional antiporter were shown to be more salt sensitive than wild-type Arabidopsis [4]. Additionally, it was found that nhx1 mutants exhibit an altered phenotype under normal growth conditions, including smaller cells, smaller leaves, and other developmental irregularities,
associated with altered K⁺ homeostasis brought about by the lack of AtNHX1. These results suggested that AtNHX1 is associated with other cellular processes that are not necessarily related to salt tolerance. Subsequently, the AtNHX1 coding region driven by the CaMV 35S promoter was introduced into the nhx1 knockout line. These ‘rescued’ plants (NHX1::nhx1) displayed AtNHX1 activity, and a phenotype similar to that of wild-type plants [4].

The transcriptional profile of the AtNHX1 'knockout' (nhx1) line has been analyzed previously [5]. That study examined the differences in transcript level using the Affymetrix® 23 k 'Full Genome' GeneChips® to look at the differences of expression profiles between wild-type and nhx1 plants grown in the absence of salt stress, and also to examine the difference in relative gene expression changes that occurred after exposure to two weeks of salt stress. It was found that there was little overlap between the two comparisons suggesting that the role of the antiporter as part of the salt stress response machinery is distinct from its role under normal growing conditions. The previous study [5] also suggested that AtNHX1 is important to the expression of several cellular processes, including components of cell structure, protein processing and trafficking, and energy balance, although AtNHX1 did not appear to dramatically affect the expression of many other transporters.

This report further establishes and clarifies the influence of AtNHX1 on gene expression, limiting the analysis to only those transcripts that respond to salt stress, and including an analysis of the influence of both shorter (12 h and 48 h) and longer (one week and two weeks) salt stress treatments. Additionally we have employed an NHX1::nhx1 'rescued' line to determine transcripts whose expression levels correlate with the expression of AtNHX1. This approach provides evidence of the influence of a single gene on the expression of other genes while helping to eliminate some of the non-specific effects that result from the mutation of the antiporter.

**Results and discussion**

Plants have been shown to have a "dual response" to salt stress, with an early response to the osmotic stress brought about by the more negative water potential of a salty soil solution, and a later response due to the Na⁺ toxicity resulting from the relatively slower entry of Na⁺ ions into the leaf tissues [6]. In an effort to include both components of the salt-stress response, we studied the influence of AtNHX1 on gene expression after 12 hours, 48 hours, one week, and two weeks of salt stress. This work is an extension of a previous microarray study that compared wild-type and nhx1 "knockout" plants before and after 2 weeks of salt stress [5]. Here the added shorter salt stress treatments (12 hours, 48 hours, and one week) and the inclusion of the NHX1::nhx1 'rescued' line allowed for a more detailed analysis of the importance of AtNHX1 to the expression of salt responsive genes. Furthermore, the greatly increased number of microarray chips used here (increased from 14 to 48) allowed for the use of a more robust ANOVA-based statistical analysis.

The NHX1::nhx1 plant line used in this study has an average increased expression of 50% of AtNHX1 as compared to the wild-type. This level of expression were sufficient to restore the wild-type phenotype [4], but was insufficient to confer meaningful salt tolerance [1]. Also, because AtNHX1 is normally expressed in all tissues and to a comparable level in all cells, with the exception of meristemmatic cells lacking vacuoles [4,7,8], expression patterns under a constitutive promoter should not differ dramatically from expression under the native promoter. The objective behind using this line was to identify transcripts with expression directly affected by the presence or absence of a functional AtNHX1.

**Overview of salt-responsive transcripts influenced by AtNHX1**

Out of the 17,030 genes that exhibited reliable expression data, 4,027 transcripts met the criteria of salt responsiveness, and 147 of these also showed a significant influence by AtNHX1, as delineated in Materials and Methods. This study focused on transcripts that showed a significant influence by both salt and AtNHX1. Other transcripts also influenced by AtNHX1 but not responding to the salt treatments, or responding to salinity but without restored levels of expression in the NHX1::nhx1 were not considered. The latter transcripts may yet be an important component of AtNHX1-related processes, but due to inherent variation in expression levels or the consequences of constitutive AtNHX1 expression, they did not meet the necessary significance criteria threshold to establish a clear relationship to the presence of the antiporter. Even with an increased statistical filtering, comparisons of more salt treatments, and an analysis of salt responsive transcripts based on absolute values rather than relative values, 42 of the 147 (>28%) transcripts that showed a significant effect of AtNHX1 in this report, were also previously shown to have an influence of AtNHX1 on expression levels [5] (comparison data not shown).

Among the 147 salt-responsive transcripts that were significantly affected by AtNHX1, 102 genes (69%) were up regulated while only 44 genes (31%) were down regulated in the absence of AtNHX1, with one transcript (At3g54810) showing increased expression after one week of salt stress, but decreased expression after two weeks of salt stress. The Genevestigator® database [9,10] was searched and most (88%) of same transcripts were found to have at least a 20% change in expression in response to
scripts were influenced by (genotype × treatment interaction). The latter 89 transcripts showed the influence of AtNHX1 under a specific salinity treatment (genotype × treatment interaction). The latter 89 transcripts were influenced by AtNHX1 typically only in one treatment (three transcripts showed a specific influence of two treatments), with fewer transcripts showing this pattern under control conditions (12 transcripts; e.g. Fig. 1C, D) or after the shortest salt treatment of 12 hours (15 transcripts; e.g. Fig. 1E, F) as compared with longer exposure to salinity (20–24 transcripts per treatment; e.g Fig. 1G–L). The two-factor ANOVA used in this study to determine the influence of AtNHX1 is considered a powerful tool for the analysis of microarray experiments with multiple factors [11], as it utilized all 48 microarray data points to distinguish between an effect of genotypes across all treatments (main effect) and a treatment-dependent effect of lines (genotype × treatment interaction). In order to focus on AtNHX1-influenced salt-responsive genes, a further statistical test was used to identify transcripts with significantly different expression levels in the nhx1 line relative to both wild type and NHX1::nhx1 lines. While AtNHX1 influenced the expression of 58 genes that were not specific to a particular salt treatment, most salt-responsive genes appeared significantly impacted in conjunction with a particular length of salt stress, with more genes influenced as the duration of stress was increased. This pattern would suggest that AtNHX1 has greater impact on the expression of other genes as the influence of salt stress shifts from initial osmotic stress to the ion stress [6].

Various databases were queried [12-14] to determine the most likely functional role of the proteins encoded by the 147 salt-responsive transcripts showing an impact of AtNHX1 on their expression levels. These transcripts were then classified into general functional groups to assist with the analysis. (Figure 2) The largest group of transcripts showing the influence of the AtNHX1 vacuolar antiporter was comprised of 58 genes (40%) with unclear functional classifications (Additional file 1). Interestingly, the percentage of unclassified transcripts was larger among the up-regulated genes (46% of the total increased) than among the down-regulated (26% of the total decreased), suggesting that more novel salt-responsive genes are increasing in the absence of functional AtNHX1.

The remaining 89 transcripts encode proteins from a variety of functional groups. The majority of encoded proteins included signaling elements, DNA binding elements, components of the protein processing and trafficking machinery, and enzymes involved with metabolic and energy balance of the cell. Details of all salt-responsive transcripts that also showed a significant influence of AtNHX1 are presented in Table 1. Specific transcripts of particular interest are discussed in the subsequent sections of this report. The research community is encouraged to explore the data for all transcripts that were found to have meaningful expression levels [15].

**AtNHX1 influences salt-responsive transcripts encoding signaling elements, including several putative calcium-binding proteins**

Thirteen salt-responsive signaling-associated transcripts were significantly influenced by the AtNHX1 antiporter (Table 2A). Nine of these transcripts exhibited significantly increased expression levels in the nhx1 line, while the expression of 4 transcripts showed reduced expression. Six of the up-regulated transcripts showed a genotype × treatment interaction with a significant effect of AtNHX1 being observed only after a week or more of salt treatment, suggesting that cellular signaling was not strongly impacted by AtNHX1 until the later stages of salt stress. The only transcripts that displayed a general trend of increased expression for all salt treatments were three kinases. These included two receptor protein kinases (At4g04540 and At5g56040) and a casein kinase II (At5g67380) all with unknown roles, although a CK2 homolog, with unidentified targets, has been implicated in the response of maize to ABA [16].

A notable feature of the signaling elements influenced by AtNHX1 is the number of transcripts encoding calcium-binding proteins, including 2 of the 9 transcripts that were up-regulated (At5g66210 and At1g52570) and 3 of the 4 transcripts (At3g09960; At2g38750; At4g34150) down-regulated in the nhx1 line. At5g66210 is a calcium-dependent protein kinase with an undetermined role, that is localized at the plasma membrane [17]. At1g52570 is a phospholipase D, shown to have regulatory functions in plant growth and development as well as the stress response (reviewed in [18]). The signaling transcripts with diminished expression in the nhx1 line included a member of the annexin family, ANNEXIN4 (At2g38750/AnnAt4). Annexins are Ca^{2+}-dependent membrane-binding proteins found in most eukaryotic species, playing roles in a wide variety of cellular processes. In Arabidopsis, they have been implicated, though not necessarily limited to, roles in Golgi-mediated secretion [19] which is also one of their key roles in animal systems. Moreover, AnnAt4, along with AnnAt1, have been shown to be important in Ca^{2+}-dependent signaling in response to osmotic stress and to ABA [20]. The other calcium-binding signaling components with diminished expression in
Expression profiles of selected salt responsive transcripts showing a significant influence of the AtNHX1 cation/H+ vacuolar antiporter. Transcripts that were found to be influenced by AtNHX1: [A,B] regardless of specific salt treatment, or [C,D] specifically under control conditions; [E,F] 12 h salt treatment; [G,H] 48 h treatment; [I,J] one week treatment; [K,L] two weeks treatment. Green ♦ = nhx1, Black ■ = wild-type, Red ▲ = NHX1::nhx1. Values are the Mean ± S.D. (n = 4 for control, n = 3 for all other treatments).
the nhx1 line included At4g34150, a transcript encoding a protein that is similar to calcium-dependent protein kinases and contains a C2 domain (Ca$^{2+}$-dependent membrane-targeting module often associated with signal transduction or membrane trafficking, [21]) and At3g09960, a calcineurin-like phosphoesterase family member [22].

The presence of several calcium binding elements provides further evidence of the influence of pH and ion homeostasis on the calcium signaling network. Calcium has been shown to be an important component of the SOS (Salt Overly Sensitive) network, with a calcium-binding protein (SOS3) in conjunction with a kinase (SOS2), influencing both the expression and activity of the SOS1/AtNHX7, a plasma membrane Na$^+/H^+$ exchanger that is important to salt stress tolerance and cytosolic pH homeostasis [23]. A previous microarray study has also shown that Ca$^{2+}$ starvation induced decreased expression of AtNHX1, AtNHX2 and AtNHX5 in Arabidopsis [24], further suggesting a link between vacuolar cation/H$^+$ antiporters and calcium levels in the cell. Moreover, the C-terminal portion of AtNHX1 itself has been shown to bind a calmodulin-like protein, with activity and ion specificity modified by the interaction, in a calcium- and pH-dependent manner [3]. Our results provide further demonstration of the influence of Ca$^{2+}$ on cellular ion and pH homeostasis.

**AtNHX1 influences the expression of DNA binding elements including water deficit responsive transcripts**

The expression of 20 salt-responsive transcripts encoding DNA binding elements (mostly transcription factors) was influenced by AtNHX1 (Table 2B). Similar to the trends

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**Figure 2**

Functional assignments of transcripts influenced by AtNHX1. Pie chart depicting the functional distribution of all 147 transcripts showing a significant influence of the AtNHX1 cation/H$^+$ antiporter.

The nhx1 line included At4g34150, a transcript encoding a protein that is similar to calcium-dependent protein kinases and contains a C2 domain (Ca$^{2+}$-dependent membrane-targeting module often associated with signal transduction or membrane trafficking, [21]) and At3g09960, a calcineurin-like phosphoesterase family member [22].
seen among the signaling elements discussed above, most (80%) of the transcription factors exhibited increased expression in nhx1 plants and the majority of the individual transcripts were influenced by a specific salt treatment. Genes encoding DNA binding elements were affected by AtNHX1 in response to both short and long terms of salt exposure whereas signaling elements were predominately influenced after longer treatments with salt. Several of these genes have been shown to be associated with the plant response to osmotic stress. At4g25490/CBF1 and At1g21910, which displayed increased expression in the nhx1 line are members of the DREB transcription factor family shown to be involved in the response of plants to different environmental stimuli by binding to dehydration-responsive element (DRE) promoter regions of stress-inducible genes [25]. CBF1, also known as DREB1B, has been shown to be involved in increasing tolerance to low temperatures, and shows a response to ABA treatment [26], and was also recently shown to be regulated by the circadian clock [27]. Conversely, expression of At4g27410/RD26 was reduced in the nhx1 line. RD26 is a drought- and salt-induced transcript belonging to the NAC gene family, that is also part of an ABA-dependent stress-signaling pathway [28]. The altered expression of these transcripts highlights the impact of AtNHX1 on known and predicted components of drought stress-related pathways.

Another transcript with an established role in the environmental stress response and influenced by the presence of the AtNHX1 was a transcriptional co-activator, At3g24500/AtMBF1c, that exhibited a 3–4 fold increase in expression as a result of the nhx1 mutation with 12 hours of salt stress. Over-expression of AtMBF1c in Arabidopsis enhanced the tolerance of the plants to different stresses (including osmotic), possibly due to perturbation of the ethylene-response signal pathway [29]. Moreover, plants over-expressing AtMBF1c demonstrated increased expression of several genes (At5g66210, At1g21910, At1g35140, At4g08950, At1g28480, and At2g32150) [29] that were also shown to be significantly influenced by AtNHX1 in this study, suggesting a possible relationship between altered ion homeostasis and stress-induced hormonal responses.

A heat shock transcription family member (At2g26150/AtHsfA2) showed a significant influence of AtNHX1 after 12 hours of salt stress. The altered level of expression of this gene may reflect another aspect of the disrupted response to stress in the nhx1 line. However it is also possible that this gene is part of the protein processing network that is disrupted in the absence of AtNHX1 (see following discussion).

Other AtNHX1-influenced transcripts encoding putative DNA binding elements have not been associated with abiotic stress response previously. At3g56980/OBP3, which increased in expression after 48 hours of salt treatment, is a transcription factor shown to target genes that are inducible by salicylic acid, and is important to normal plant development [30]. At3g54810/BME-ZF has been shown to act as a regulator of seed germination during cold stratification [32], which may reflect a role in the response to environmental stimuli similar to other GATA-type genes.

| Table 1: Functional distribution of the 147 gene transcripts influenced by both salinity and AtNHX1. |
|--------------------------------------------------|--------------------------------|--------------------------------|--------------------------------|--------------------------------|--------------------------------|--------------------------------|--------------------------------|
| Gene classification                              | # of transcripts influenced under each treatment1 | Distribution of decreased/increased transcripts in the nhx1 mutant2 |
|--------------------------------------------------|--------------------------------|--------------------------------|--------------------------------|--------------------------------|--------------------------------|--------------------------------|
| Unclassified                                      | 58                             | 12                             | 15                             | 21                             | 24                             | 20                             | 45                             | 103                            |
| DNA binding                                      | 12                             | 4                              | 4                              | 3                              | 3                              | 5                              | 16                             |
| Membrane Transport                               | 1                               | 0                              | 1                              | 1                              | 0                              | 1                              | 3                              |
| Metabolism/Energy                                | 12                             | 0                              | 3                              | 6                              | 2                              | 16                             | 9                              |
| Structure/Growth                                 | 2                               | 1                              | 1                              | 3                              | 4                              | 2                              | 4                              | 9                              |
| Signaling                                        | 6                               | 0                              | 0                              | 1                              | 4                              | 2                              | 4                              | 9                              |
| Protein Processing                               | 8                               | 1                              | 1                              | 1                              | 1                              | 2                              | 3                              | 11                             |

1 three transcripts were specifically influenced by AtNHX1 under two treatments (At4g17120, At5g47490 – both unclassified, significantly affected by Control and 12 h treatments – and At3g54810 – DNA binding, significantly affected by 1wk and 2wk treatments)

2 one transcript (At3g54810) was up-regulated in one treatment (1wk) and down regulated in a second treatment (2wk)
Table 2: Specific salt-responsive transcripts influenced by AtNHX1, organized by functional category

| Accession | Functional Classes and Gene Descriptions | P(f)a Treatment influenced by AtNHX1b | Transcripts intensity under the influenced treatmentc |
|-----------|------------------------------------------|---------------------------------------|-----------------------------------------------------|
|           |                                          | L □ □ wild-type NHX1::nhx1              |                                                     |
|           |                                          | LxT □ □ nhx1 □ □ □ □ wild-type NHX1::nhx1 |                                                     |
| A. DNA binding elements | | |
| At3g53730 | histone H4                                | Control                               | 2511.1 3461.3 4184.2                                 |
| At5g67580 | myb family transcription factor           | Control                               | 287.4 123.8 149.1                                   |
| At3g55330 | methyl-CpG-binding domain-containing protein | Control                               | 772.7 526.9 508.3                                  |
| At1g4685  | BASIC PENTACYSTEINE 2, BPC2               | Control                               | 555.1 370.5 390.9                                   |
| At2g36960 | myb family transcription factor           | 12 h                                  | 290.0 368.5 476.0                                  |
| At2g26150 | heat shock transcription factor family     | 12 h                                  | 684.1 98.8 181.5                                   |
| At3g24500 | Transcriptional Coactivator Multiprotein Bridging Factor 1c. | 12 h                                  | 1024.0 261.8 393.4                                |
| At1g69010 | basic helix-loop-helix (bHLH) family protein | 48 h                                  | 422.3 278.1 243.7                                  |
| At5g36980 | basic helix-loop-helix (bHLH) family protein | 48 h                                  | 503.2 304.3 130.8                                  |
| At2g5490  | DRE-binding protein (DREB1/CBF/CRE-binding factor 1 (CBF1)) | 48 h                                  | 758.3 522.5 569.9                                  |
| At1g69580 | myb family transcription factor           | 1 wk                                  | 236.4 141.3 135.7                                  |
| At3g54810 | zinc finger (GATA type) family protein1   | 1 wk                                  | 1168.0 571.7 340.6                                 |
| At2g1730  | putative ethylene-responsive protein      | 1 wk                                  | 293.2 147.8 47.1                                   |
| At1g2910  | DREB A-5 subfamily member, ERF/AP2 transcription factor family | 1 wk                                  | 1871.5 771.5 598.6                                 |
| At3g54810 | zinc finger (GATA type) family protein3   | 2 wk                                  | 1018.2 539.9 360.6                                 |
| At5g57660 | zinc finger (B-box type) family protein   | 2 wk                                  | 1108.0 1585.0 1545.2                               |
| At4g00850 | GRF1-interacting factor 3 (GIF3), SSXT family protein | 2 wk                                  | 366.7 273.8 86.5                                   |
| At3g02420 | zinc finger (C3HC4-type RING finger) family protein | 2 wk                                  | 1018.2 539.9 360.6                                 |
| At5g56860 | zinc finger (GATA type) family protein    | 2 wk                                  | 1018.2 539.9 360.6                                 |
| At1g18710 | myb family transcription factor (MYB47)   | All                                   | 257.3 460.3 467.0                                  |
| B. Signaling Elements | | |
| At4g34150 | C2 domain-containing, similar to calcium-dependent protein kinase | 48 h                                  | 2199.5 4215.4 4558.0                               |
| At4g08960 | phosphotyrosyl phosphatase activator (PTPA) family protein | 1 wk                                  | 542.6 377.1 279.9                                  |
| At3g54380 | protein kinase family protein             | 1 wk                                  | 1866.1 1250.7 826.4                                |
| At3g54840 | GTP-binding family protein                | 1 wk                                  | 134.1 60.7 57.2                                    |
| At5g66210 | calcium-dependent protein kinase family protein (CPK28) | 1 wk                                  | 367.5 221.8 213.5                                  |
| At1g32570 | phospholipase D alpha 2 (PLD2)/choline phosphatase 2 | 2 wk                                  | 229.2 99.4 74.5                                    |
| At2g24160 | pseudogene, leucine rich repeat protein family | 2 wk                                  | 349.8 160.4 71.7                                   |
| At2g38750 | annexin 4 (ANN4)                          | All                                   | 511.2 875.1 804.2                                  |
| At3g09960 | calcineurin-like phosphoesterase family protein | All                                   | 59.2 98.7 104.2                                   |
| At4g21370 | putative S-locus protein kinase, pseudogene | All                                   | 63.0 102.8 110.1                                  |
| At4g05450 | protein kinase family/ protein kinase family protein | All                                   | 412.6 289.1 220.3                                  |
| At5g56040 | leucine-rich repeat protein kinase, putative | All                                   | 930.9 748.2 617.4                                  |
| At5g67380 | casein kinase II alpha chain 1            | All                                   | 637.3 496.1 488.6                                  |
| C. Metabolism/Energy Components | | |
| At4g11600 | putative glutathione peroxidase (AtGPX6)  | 12 h                                  | 3636.1 5433.9 4962.7                                |
| At1g68290 | bifunctional nuclease, putative          | 12 h                                  | 105.1 236.2 244.6                                  |
| At3g16050 | putative pyridoxine (Vitamin B6) biosynthesis protein | 12 h                                  | 403.6 121.1 227.3                                  |
| At4g32360 | NADP adrenodoxin-like ferredoxin reductase | 48 h                                  | 102.5 172.0 203.7                                  |
| At2g26560 | putative patatin (PLP2)                   | 48 h                                  | 1647.2 3298.0 3515.3                                 |
| At1g56430 | putative nicotianamine synthase           | 48 h                                  | 995.2 433.8 601.6                                  |
| At3g05350 | phosphoesterase family protein           | 48 h                                  | 208.4 125.2 122.2                                  |
| At5g05960 | protease inhibitor/seed storage/lipid transfer protein (LTP) family protein | 48 h                                  | 754.4 423.3 436.3                                  |
| At3g53440 | FAD-binding domain-containing protein/cytokinin oxidase family protein | 48 h                                  | 224.6 132.2 48.8                                   |
| At4g09555 | amidohydrolase family protein             | 1 wk                                  | 204.2 296.5 465.4                                  |
| At1g63710 | cytochrome P450, putative                | 1 wk                                  | 126.9 67.9 25.9                                    |
| At2g17570 | undecaprenyl pyrophosphate synthetase family protein | 2 wk                                  | 112.3 206.2 310.0                                  |
Table 2: Specific salt-responsive transcripts influenced by AtNHX1, organized by functional category (Continued)

| Accession number | Description | Plant line effect | Treatment | Expression
|------------------|-------------|------------------|-----------|-------------|
| At5g19890        | putative peroxidase | * ** | 2 wk | 212.8 99.3 79.7 |
| At1g28480        | glutaredoxin family protein | *** All | All | 349.1 696.8 1131.5 |
| At2g46650        | cytochrome b6, putative | ** | All | 1075.4 1748.2 1638.4 |
| At5g17220        | glutathione-S-transferase (AtGSTF12, TT19) | * | All | 270.8 402.7 422.1 |
| At4g39940        | adenylylsulfate kinase 2 (AKN2) | ** | All | 1940.2 2675.4 2748.4 |
| At4g0610         | 5′-adenylylsulfate reductase (APR1)/PAPS reductase homolog (PRH19) | * | All | 507.7 1362.0 1284.4 |
| At3g22740        | homocysteine S-methyltransferase 3 (HMT-3) | *** All | All | 622.8 928.0 1155.4 |
| At1g1440         | mutase family protein | ** | All | 2176.8 2373.5 2793.7 |
| At1g06520        | phospholipid/glycerolacyltransferase family protein | ** | All | 69.9 120.1 126.8 |
| At1g16410        | cytochrome P450 family protein (CYP79F1) (bushy1) | *** All | All | 280.2 480.4 492.0 |
| At2g32150        | halocid dehalogenase-like hydrolase family protein | ** | All | 357.9 720.0 857.9 |
| At5g47240        | MutT/nudix family protein | *** All | All | 961.5 1939.5 1471.4 |
| At2g06050        | 12-oxophytodienoate reductase (OPR3)/delayed dehiscence 1 (DDE1) | ** All | All | 804.0 1315.1 1454.0 |

D. Structure/Growth Components

| Accession number | Description | Plant line effect | Treatment | Expression
|------------------|-------------|------------------|-----------|-------------|
| At2g20000        | cell division cycle family protein/CDC family protein | * | Control | 249.7 274.3 260.2 |
| At2g40610        | expansin, putative (EXP8) | * * | 12 hours | 792.4 462.8 381.2 |
| At1g27630        | cyclin family protein | ** | 48 h | 519.1 744.7 913.9 |
| At3g02350        | glycosyl transferase family B protein | * | 48 h | 1160.2 947.2 609.2 |
| At1g19170        | glycoside hydrolase family 28/polygalacturonase (pectinase) family | * | 48 h | 365.0 220.3 177.9 |
| At3g45970        | expansin family protein (EXP1/AtEXLA1) | * | 1 wk | 3780.8 1684.2 1026.0 |
| At3g27220        | galactosyl transferase GMA12/MNN10 family protein | ** | 1 wk | 2237.1 1459.9 790.3 |
| At5g37560        | cell wall-modifying enzyme, endo-xylloglucan transferase (TCH4) | ** | 1 wk | 1349.3 6318.4 6047.0 |
| At4g30470        | cinnamoyl-CoA reductase-related | * | 1 wk | 756.6 478.9 512.6 |
| At1g57590        | putative pectinacetylesterase | * | 2 wk | 143.8 381.3 336.7 |
| At1g6340         | putative 3-deoxy-D-manno-2-octulosonate-8-phosphate synthase | * | 2 wk | 352.0 231.3 48.7 |
| At4g16590        | glucosyltransferase-related | * | All | 194.5 578.7 602.4 |
| At1g24070        | glycosyl transferase family 2 protein (AtCSLA10) | ** | All | 280.2 509.7 495.6 |

E. Protein Processing

| Accession number | Description | Plant line effect | Treatment | Expression
|------------------|-------------|------------------|-----------|-------------|
| At3g17970        | chloroplast outer membrane translocon subunit, putative | * | Control | 252.5 115.4 162.0 |
| At2g20560        | DNAJ heat shock protein | * | 12 h | 413.3 159.6 155.0 |
| At1g08780        | prefoldin, putative | * | 48 h | 476.4 303.2 182.0 |
| At2g47440        | DNAJ heat shock N-terminal domain-containing protein | * | 1 wk | 3769.5 2247.3 1271.8 |
| At1g08730        | myosin heavy chain (PRC43) (XIC) | *** *** | 2 wk | 25.4 147.4 204.6 |
| At5g58810        | subtilisin-like serine protease, similar to prepro-cucumisin | ** *** | 2 wk | 24.0 148.8 201.3 |
| At5g59730        | exocyst subunit EXO70 family protein | * | All | 768.8 1096.8 1199.6 |
| At3g25150        | nuclear transport factor 2 (NTF2) family protein | *** | All | 693.3 527.6 439.1 |
| At5g4760         | 26s proteosome regulatory subunit, putative (RP5S) | ** | All | 419.3 340.3 316.2 |
| At1g22740        | Ras-related protein (RAB7/AtRab75/small GTP-binding | *** | All | 1324.9 892.1 703.5 |
| At2g22040        | transducin family protein/WD-40 repeat family protein | *** | All | 362.1 283.2 231.5 |
| At5g47820        | kinesin-like protein (FRA1) | ** | All | 389.3 303.8 259.8 |
| At4g34980        | subtilase family protein (SLP2) | ** | All | 1133.4 899.5 930.4 |
| At3g23670        | phragmooplast-associated kinesin-related protein, putative | ** | All | 138.2 101.9 75.6 |

F. Membrane Transport

| Accession number | Description | Plant line effect | Treatment | Expression
|------------------|-------------|------------------|-----------|-------------|
| At2g23980        | cyclic nucleotide-regulated ion channel (CNGC6) | * * | Control | 393.3 259.6 180.7 |
| At2g47830        | cation efflux family/metal tolerance (MTPc1) | ** | 48 h | 73.9 149.7 166.0 |
| At1g34170        | nodulin-related | * | 1 wk | 223.1 148.5 110.3 |
| At2g25520        | phosphate translocator-related | ** | All | 999.7 790.3 773.9 |
The nhx1 plants have been shown to have altered leaf development, in addition to increased salt sensitivity [4], and the expression of several transcription factors associated with leaf morphology and development were influenced by AtNHX1. While most developmental genes are expected to be independent of salinity effect, two genes were significantly influenced by AtNHX1 under specific salt treatments. The expression of At2g36960, encoding the TOUSLED gene, was decreased in the nhx1 line after 12 hours of salt stress. TOUSLED interacts with chromatin regulators and its expression normally increases in dividing cells [33]. In addition, At4g00850/AtGIF, involved in leaf growth and morphology [34] showed a significant effect of AtNHX1 after two weeks of salt stress. Possibly, these factors contribute to the altered gene expression that is associated with the nhx1 phenotype [4].

**AtNHX1 is associated with sulfur metabolism**

Of the 89 AtNHX1-influenced transcripts with an assigned or putative function, 25 transcripts, found on Table 2C, encode genes with metabolism or energy functions not directly associated with cell structure or cell growth (discussed in the next section). The majority of these transcripts had significantly lowered expression in the nhx1 line, in contrast to the overall patterns of genes showing mostly increased expression in the absence of AtNHX1. This pattern would suggest an overall decrease of metabolism- and energy processes-related genes in the knockout plants.

Twelve of the 18 metabolism/energy-related transcripts down-regulated in the nhx1 plants were generally decreased in the nhx1 line over all treatments. On the other hand, the transcripts with increased expression in nhx1 plants were responsive to particular lengths of salt stress. These results indicated that, though in general gene expression was enhanced in the nhx1 line to compensate for altered ion homeostasis, metabolic and energy processes were compromised in the absence of AtNHX1.

At least 5 of the 12 transcripts with diminished expression over all salt treatments in the nhx1 line appeared to be associated with sulfur/sulfate metabolism pathways. Transcripts encoding adenosine-5'-phosphosulfate-kinase (At4g39940/AKN2), a 5'-adenyllysulfate reductase/PAPS reductase homolog (At4g04610/APR1/PRH19), and a homocysteine methyltransferase (At3g22740/HMT3) have well established roles in sulfur metabolism [35]. The diminished expression of these transcripts would suggest a decrease in the synthesis of both glucosinolates and methionine within the leaves of the nhx1 plants. Other sulfur-related transcripts were also diminished over all treatments in the nhx1 line, encoding a glutathione S-transferase (At5g17220/AtGSTF12) a putative glutaredoxin (At1g28480), and CYP79F1 (At1g16410) a protein that mediates the formation of glucosinolates that are derived from methionine [36]. Additionally, a glutathione peroxidase (At4g11600/AtGPX6), which is known be regulated by abiotic stress [37], was down-regulated in the nhx1 line specifically with 12 hours of salinity stress.

There are several other down-regulated transcripts that are also likely to play a role in sulfur assimilation pathways. OPR3 (At2g06050) catalyzes the middle step in jasmonic acid biosynthesis, has been associated with the plant response to environmental stresses, and influence the sulfur metabolic pathway [38]. These results highlight a link between S-assimilation/metabolism and the expression levels of the AtNHX1 antiporter, as also suggested by a study using transgenic Brassica plants overexpressing AtNHX1 [39].

**AtNHX1 influences cell wall metabolism and components of cell growth**

Thirteen salt-responsive, AtNHX1-influenced transcripts, were associated with cell wall metabolism and cell growth (Table 2D). Nine of these exhibited increased expression in the nhx1 plants, mostly after exposure to salt stress of two days or longer. The up-regulated cell wall-associated genes included At5g57560/TCH4 – encoding an endoxylloglucan transferase that has been shown to be rapidly up-regulated in response to many environmental and hormonal stimuli [40], a galactosyltransferase (At5g62720), a galacturonosyltransferase (At3g02350), a polygalacturonase family member (At1g19170), a putative cinnamoyl-CoA reductase (At4g30470), and a 3-deoxy-D-manno-octulosonate 8-phosphate synthase (At1g16340). Transcripts encoding proteins with cell-wall associations also had diminished expression in the nhx1 line, including two cellulose synthase-like genes (At4g16590 and At1g24070) that were diminished with all treatments, and a pectinacylesterase (At1g57590) transcript that was diminished after two weeks of salt stress.

The altered expression of the above-mentioned transcripts associated with cell size and structure, in addition to some of the transcription factors mentioned earlier, are likely to be involved in the altered developmental phenotype of the nhx1 line, showing smaller cells, smaller leaves and diminished growth [4]. There are also four salt responsive transcripts displaying altered expression levels in the absence of the AtNHX1 that are part of cell expansion and growth. Under control conditions a cell division gene (At2g20000/HBT) has increased expression in the nhx1 line whereas with 48 hours of salt stress a cyclin family protein (At1g27630) shows decreased expression. Two putative expansins also show increased nhx1 expression levels (At2g40610/AtExpA8 and At3g45970/AtExpA1A) at 12 hours and one week of salt stress, respectively. Intracellular ion and pH homeostasis is important to the regu-
The impact of stress treatments, specific to a particular salt stress treatment, suggesting a small GTP-binding Ras-related protein, in the absence of functional AtNHX1 can change the rate of cell proliferation remains to be demonstrated.

**AtNHX1 influence the expression of protein processing and trafficking components in response to salt stress**

Fourteen of the AtNHX1-influenced salt-responsive genes appeared to play roles in the processing and trafficking of other cellular components and proteins (Table 2E). Nhx1, the yeast orthologue of AtNHX1, has been shown to play an important role in protein trafficking in yeast [45,46], and the regulation of endosomal pH by Nhxl controls the vesicle trafficking out of the endosome [47].

Eleven of the salt-responsive protein processing/trafficking components had increased expression due to the absence of AtNHX1, with seven of these transcripts not specific to a particular salt stress treatment, suggesting an influence of AtNHX1 over the entire range of the studied stress treatments.

The impact of AtNHX1 on vesicular trafficking is reflected by the altered expression of At1g22740, encoding RAB7, a small GTP-binding Ras-related protein, in the nhxl line. Rab GTPases are part of the organization of intracellular membrane trafficking, including vesicle formation, vesicle motility, and vesicle tethering [48], and Rab7-related genes are important for the regulation of the late steps of endocytotic pathway. The overexpression of a Rab7 homolog stimulated endocytosis and conferred tolerance to salinity and oxidative stress in *Arabidopsis* [49,50]. Also a rice homologue of this gene was differentially regulated by both ABA and salinity and was implicated in vesicular traffic to the vacuole [51].

The altered expression pattern of an exocyst subunit EXO70 family protein (At5g59730) may be a further indication of the role of AtNHX1 in vesicular trafficking. Though not yet fully characterized in higher organisms, the EXO70 family members are important to vesicle docking and membrane fusion as well as regulation of actin polarity and transport of exocytic vesicles in yeast [52,53]. Also two kinesin-related transcripts (At5g47820 and At5g23670) showed an altered expression pattern. Kinesins are key to the intracellular transport system ([54] and references therein).

Four salt-responsive transcripts with roles in protein processing that are influenced by AtNHX1, emphasize the role of ion homeostasis on the proper folding and function of other proteins. These include two Dnaj-type genes (At2g20560 and At2g47440), a prefoldin (At1g08780), and a transducin/WD-40 repeat containing gene (At2g22040). The altered expression of these genes would suggest that the absence of AtNHX1 induces the instability of other proteins. Also, the altered expression of subtilases (At5g58810 and At4g34980) and a 26S proteasome regulatory subunit (RPN5/At5g64760) suggest a possible influence on protein degradation pathways.

A salt-responsive myosin XI subunit was also influenced by AtNHX1 (PCR43/XIC/At1g08730). Myosin XI mutants have been shown to be defective in both organelle movement and polar auxin transport [55] through the action on several vesicle-mediated processes. The altered expression of both a nuclear transport factor (NTF2/At3g25150) and a chloroplast outer membrane translocon subunit (At3g17970) would suggest a potential influence of AtNHX1 on trafficking of cellular components to organelles. Additionally, AtNHX1-influenced transcripts in other functional categories may also be related to a role of the antiporter as part of vesicular trafficking. For example, At2g17570, encoding a member of the undecaprenyl pyrophosphate synthetase family (Table 2C – Metabolism) is homologous to the yeast gene RER2, was shown to be important to vesicular processes and organelle integrity [56].

**Most salt-responsive transporters genes are not significantly influenced by AtNHX1**

The Arabidopsis NHX family is comprised of 6 endomembrane (AtNHX1-6) and 2 plasma membrane-bound (AtNHX7/SOS1 and AtNHX8) members and in the absence of AtNHX1, compensation by the other AtNHX members might be expected, in particular when the plants are exposed to salt stress. However, our data did not show significant changes in the expression of any of the AtNHX2-8 transcripts either in nhxl or NHX1::nhxl plants in response to salt. Additionally, though the differences of AtNHX1 signal detection were at 27% and 160% of wild-type levels (p < 0.0001) for the nhxl and NHX1::nhxl lines, respectively, the other transporter genes did not show a significant difference of expression levels between lines regardless of the salt treatment used (data not shown).

A few salt-responsive transporters did show an apparent affect of AtNHX1 on expression levels (Table 2F). A putative phosphate transporter (At2g25520) showed an overall increased level of expression in the nhxl plants, possibly as a result of an imbalance of phosphate ions as proton efflux from the vacuole is changed in the nhxl line.
A cyclic nucleotide-regulated ion channel (At2g23980/CNGC6) also showed increased expression in the nhx1 line. CNGCs comprise a family of 20 members in Arabidopsis, activated by direct binding of cyclic nucleotides and regulated by CaM [57]. They can provide a significant pathway for the non-selective uptake of ions (Na+, K+ or Ca2+) and several family members were up-regulated or down-regulated by salt stress [58]. Since an increase in cellular cGMP was shown to occur during salt and osmotic stress [59], and the expression of AtCNGC6 was shown to be up-regulated in plants exposed to cGMP [60], it could be hypothesized that the overexpression of AtCNGC6 is related to the Na+-induced K+-deficiency. Lastly, the expression of a nodulin-related gene (At1g31470) was increased in the nhx1 line with one week of salt stress, and the expression a cation efflux/metal tolerance family gene (At2g47830) was decreased with 48 hours of salt stress. The role of these putative transporters has yet to be elucidated.

Little is known about the influence of AtNHX1 on the expression/activity of other transporters within the plant cell. Previous work showed that AtNHX1 influenced the expression of a few genes encoding putative transporters [5]. However, as noted by Gong, et al. [61], previous microarray studies of salt stress in Arabidopsis (eg. [62,63]) did not demonstrate significantly altered expression of transporters, such as AtNHX1 or SOS1, which are known to contribute to ion homeostasis and salt tolerance [1,64]. Furthermore, a wide survey of available Arabidopsis microarray data suggested that only approximately 40 transcripts encoding putative cation transporters showed a significant response to salt or drought stress, with less than a 10% overlap between studies [58]. This emphasizes the influence of the experimental design on the expression profiles, suggesting a high level of inherent variability. Several factors might interfere with the detection of transcriptional changes in the genes encoding these transporters during salt stress, such as relatively low levels of expression or post-translational mechanisms that can modify the transporters affinity, selectivity, and/or its kinetics without affecting transcript expression [3,65].

**Conclusion**

A unique feature of this study is the utilization of both an nhx1 'knockout' line and a 'rescued' mutant line (NHX1::nhx1) to identify transcripts with expression changes directly related to the presence of a single gene, AtNHX1. A previous study of the influence of AtNHX1 [5] on gene expression, was limited to only the nhx1 line in comparison to wild-type before and after the exposure of the plants to long-term (two weeks) salt stress. This work is a logical extension of the findings from the previous publication, because it provides novel aspects of the influence of the antiporter, especially as part of the salt stress response. We have provided evidence that AtNHX1 has a larger effect on salt responsive transcripts with increased salt stress duration rather than during the early exposure, emphasizing the increased importance of the antiporter during the later ionic effects of salt stress. Nonetheless the detection of AtNHX1-influenced salt-responsive transcripts during the earlier salt stress treatments, and the presence of 57 transcripts that appeared influenced regardless of any particular stress treatment, also highlights the role(s) of AtNHX1 throughout salt stress exposure. The use of short- and long-terms of sub-lethal levels of salt stress, together with the NHX1::nhx1 line, facilitated the elucidation of adaptive responses that are influenced by the vacuolar antiporter. In line with its importance to salt stress tolerance, our results demonstrate that AtNHX1 influenced transcripts with known roles in the response to water deficit stress. We have additionally provided further evidence that AtNHX1 impacts the expression of other components of the response of Arabidopsis to stress. Recently, it has been shown that AtNHX1 activity can be modulated by calcium levels within the cell [3], and our results demonstrated that several Ca2+-binding elements were also affected transcriptionally by the presence of the antiporter protein. Furthermore, in addition to many uncharacterized transcripts, AtNHX1 also showed an impact on the transcription of several other key cellular processes including: sulfur metabolism, vesicular trafficking, protein processing, energy transfer processes, and cell growth/structure.

Up-regulation of most of the AtNHX1-influenced salt-responsive transcripts in the absence of AtNHX1 would suggest the activation of compensatory mechanisms in the nhx1 plants. Nevertheless, the decreased expression in transcripts encoding proteins with roles in metabolism and energy transfer would correlate with the phenotype displayed by the knockout plants, i.e. reduction of leaf area, smaller plants, and increased salt-sensitivity [4]. Also, the influence of AtNHX1 on vesicular trafficking and protein processing did not appear to be associated with any particular salt stress treatment, but rather appears to be an expression phenotype of the nhx1 plants, further indicating that, similar to its homolog in yeast [45-47], AtNHX1 plays an important role in ion and pH homeostasis of the cell endosomes.

The relatively small effect of AtNHX1 on the expression of other transporters during salt stress is noteworthy. Other microarray studies have also shown little impact of salt stress on the expression of ion transporters [62,63]. It could be argued that the non-lethal salt concentrations used here and in previous studies precluded the detection of significant changes in expression of transporters, and that under these conditions ion transport may be regu-
lated primarily at the level of activity. Nevertheless, the nhx1 plants, in addition to being more sensitive to salt stress, are decreased in size, show developmental changes, and have decreased vacuolar H+-coupled cation transport [4]. This would indicate that any possible compensatory transport mechanism in the knockout plants was insufficient to maintain ion homeostasis at wild-type levels.

Methods

Plant materials and growth conditions

Three lines of Arabidopsis thaliana were used for this study, wild-type line (ecotype Wassilewskija; 'WS'), a 'knockout' line (nhx1) with a T-DNA insertion in the ninth exon of the AtNHX1 gene, and a 'rescued' line (nhx1::NHX1) with a single copy of the AtNHX1 coding sequence driven to constitutive expression by the 35S CMV promoter using the nhx1 line as the genetic background [4]. Seeds were surface-sterilized with bleach and plated at an even density (~1 seed cm\(^{-2}\)) in Petri dishes containing a modified MS growth medium supplemented with 8% agar and 5% sucrose. Seeds were germinated in an incubator (Model CU-36L; Percival Scientific, Perry, IA, USA) at 22°C under a 12-h photoperiod. Two weeks after sowing, seedlings of uniform size were selected and were transplanted into 100 ml pots (five seedlings per pot) containing moist soil mixture (MetroMix 200; Scotts Sierra Horticulture Products, Marysville, OH, USA). The pots were covered with a transparent plastic cover and placed in a growth chamber (Model AC-40 Controller 6000; Enconair, Winnipeg, MB, Canada) at 22°C under a short-day cycle (8 h light, 16 h dark) in order to delay bolting and enhance leaf development. Inflorescence tissues were removed nine days later (one week before harvest) to further emphasize leaf growth and to minimize developmental differences among plant lines and treatments. Plant were allowed to acclimatize for two days after transplanting and the soil was then saturated with the modified MS medium without or with supplemental 100 mM NaCl, as required. The watering solution was applied to the soil surface, allowed to drain and drainage was immediately removed to avoid salt accumulation.

Plants were subjected to salt stress for durations of 12-hours, 48-hours, 1-week or 2 weeks. The 2-week treatment was initiated after a 2-days acclimatization period, whereas other treatments were initiated afterwards at different times so that all treatments were harvested concurrently at the same age, 30 days after sowing. Plant material, excluding root and inflorescence tissues, was immediately frozen in liquid nitrogen for later expression analyses. Subsets of 25 plants (5 pots) of the same treatment and plant line were pooled to form an independent biological replicate. Four samples for control plants and three for each of the salt-stressed treatments were collected from each plant line.

RNA extraction and GeneChip® hybridization

Frozen plant samples were ground to a fine powder and RNA was extracted by a modification of the hot-phenol method [66]. After quality confirmation by agarose gel electrophoresis, the extracted RNA was prepared for array analysis as suggested by the manufacturer [67]. Briefly, ds-cDNA was made from total RNA, followed by formation of biotin-labeled cRNA, which was purified and fractionated prior to hybridization on individual gene chips. After overnight hybridization, the chips were stained with streptavidin-phycoerythrin and biotinylated anti-streptavidin antibody, then scanned by laser, producing an image file, the basis for quantifying and comparing relative transcript levels. Quantification of the data depends on a number of mathematical factors as optimized by Affymetrix® [68] but is primarily based on the hybridization of experimental RNA to probe sets, each consisting of 11 representative 25-mer perfect match probes complementing unique portions of different transcripts and 11 corresponding single mismatch oligomer sequences. For this study, the Affymetrix® ATH1-121501 Genome Array GeneChip® was used, containing probe sets for 22,746 predicted and known expressed Arabidopsis genes.

Data analysis

The data images produced by the microarray scanning were interpreted by Affymetrix® Microarray Suite 5.0 (MAS 5.0) software with scaling of all probe sets to a target value of 500. The purpose of this chip-wide scaling was to minimize chip-to-chip difference in overall hybridization intensities [69]. A numerical file of all the data was produced and any transcript that did not generate a detection P-value <0.05 [70] for at least one chip was removed from the analysis (the default P-value cut-off for a 'present' expression call is 0.065). This filter eliminated 5,716 genes with unreliable expression data; because of low detection levels or non-specific probe sets. This also eliminated a large majority of transcripts with non-normal distribution of detection value data generated by MAS 5.0 algorithms [71]. Data from the remaining 17,030 genes were first normalized to an invariant set using dChip v1.2 computer software [72,73] and exported into Microsoft® Excel® (Microsoft Corp., Redmond, WA, USA) for further processing and analyses. Two statistical methods were used to identify salt responsive genes. A cross-wise log2 ratio analysis was performed with a cut-off threshold for significance set at two standard errors from a log change ratio of 0.585, corresponding to a 95% probability that the true mean represents at least a 50% deviation from the control treatment. In addition, one-tailed Student's homoscedastic t-tests with cutoff of P < 0.05 were used to evaluate the statistical significance of the difference between gene expression data under each salinity treatment vs. under the respective control. Twelve (3 plant lines × 4 salinity treatments) comparisons were made for
each of the 17,030 genes. Only transcripts with comparisons that satisfied both statistical conditions under at least one salt treatment in at least one of the lines were retained for further analysis. These comparisons limited further analysis to the 4,027 transcripts that showed a significant response to salt treatment for at least one comparison. This approach to determine salt responsive transcripts, was used in a previous study of salt-treated Arabidopsis and verified by quantitative real-time PCR demonstrated the consistency of the method [5].

The selected salt-responsive genes were further analyzed to discover those that were influenced most strongly by the AtNHX1 antiporter. Of particular interest were gene transcripts either up- or down-regulated in the nhx1 'knockout' line, as compared to the wild-type line, with recovered expression levels in the NHX1::nhx1 'rescued' line. The 4,027 salt-responsive gene transcripts were subjected to a two-factor model analysis of variance using JMP software (SAS Institute, 2005). Gene transcripts showing a significant (p(F) < 0.05) line x treatment interaction (indicating treatment-dependent effect of plant line) or a significant main effect of the plant line (indicating difference between lines across all treatments) were subjected to means comparison by Student's t-test. Transcripts expression levels of the three plant lines were compared, either under each environment separately (for those showing significant interaction) or averaged across environments (for those showing a significant plant line effect but no significant interaction). A transcript was considered to have an unclear cellular function if it exhibited a significantly decreased (down-regulated) or increased (up-regulated) level of expression in the nhx1 line relative to both the WS line and the NHX1::nhx1 line.

Authors’ contributions

JBS carried out the microarray studies, analyzed the data and drafted the manuscript; YS contributed to the data analysis and the preparation of the manuscript; EB contributed to the experimental design, data analysis and the final preparation of the manuscript. All authors have read and have approved the final manuscript.

Additional material

Additional file 1

Specific salt-responsive transcripts influenced by AtNHX1 that have an unclear cellular function Description: The 58 transcripts that met the same criteria as those found in Table 1 but that currently have an unclear functional classification

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