Genetic and molecular characterization reveals a unique nucleobase cation symporter 1 in Arabidopsis

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1. Introduction

In plants, nucleobases play a pivotal role in metabolism, growth, and development. Nucleic acid metabolism, carbohydrate, glycoprotein and phospholipid metabolism as well as the biosynthesis of many secondary metabolites, such as cytokinins, theobromine and caffeine, are just a few examples of the myriad of biochemicals dependent upon nucleobases [1–4]. Nucleobase biochemistry is marked by a sophisticated interaction between salvage reactions, de novo synthesis and catabolism pathways. A further trait in nucleobase biochemistry is the high degree of compartmentalization, necessitating extensive intra- and inter-cellular transport. A prime example is the essential interplay of nucleobase biochemical pathways in seed germination and early seedling development necessitating both intercellular transport of metabolites from endosperm to cotyledons as well as intracellular transport between organelles [1–7]. This highly involved nucleobase transport serves the numerous needs for nucleobases in plants and stands in contrast to nucleobase transport in microbes that mainly serves salvaging of external nitrogen sources.

The extent and importance of nucleobase transport is reflected in the Arabidopsis genome which contains six different gene families encoding for nucleobase transporters. Two transporter families are unique to plants and include the purine permease (PUP) with 21 members and ureide permease (UPS) with eight members. Characterized Arabidopsis PUP proteins transport adenine, cytosine or secondary compounds such as cytokinins and caffeine [8–10]. Two transporters in the Arabidopsis UPS family transport uracil, allantoin and the purines xanthine and hypoxanthine [11–13], and an UPS from French bean transports allantoin [14]. The remaining four nucleobase transporter families have orthologs in other prokaryotic and eukaryotic taxa. Nucleobase-ascorbate transporters (NATs) are ubiquitous and collectively transport an array of solutes including purines, oxidized purines (xanthine, hypoxanthine, and uric acid), uracil and ascorbate [15,16]. Although none of the 12 identified Arabidopsis NAT loci are functionally-characterized yet, the maize Leaf Permease 1 is known to transport xanthine and uric acid [17,18]. The AzgA-like transporter family in Arabidopsis facilitates the movement of adenine, guanine and uracil [19]. What appears to be the only member of the nucleobase cation symporter 1 (NCS) is encoded by Arabidopsis locus At5g03555 which encodes a protein with significant amino acid similarity to FUR4, a uracil transporter of Saccharomyces cerevisiae [20].

Abstract
Locus At5g03555 encodes a nucleobase cation symporter 1 (AtNCS1) in the Arabidopsis genome. Arabidopsis insertion mutants, AtNcs1-1 and AtNcs1-3, were used for in planta toxic nucleobase analog growth studies and radio-labeled nucleobase uptake assays to characterize solute transport specificities. These results correlate with similar growth and uptake studies of AtNCS1 expressed in Saccharomyces cerevisiae. Both in planta and heterologous expression studies in yeast revealed a unique solute transport profile for AtNCS1 in moving adenine, guanine and uracil. This is in stark contrast to the canonical transport profiles determined for the well-characterized S. cerevisiae NCS1 proteins FUR4 (uracil transport) or FCY2 (adenine, guanine, and cytosine transport).

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Although each plant nucleobase transporter has a unique solute transport specificity and gene expression pattern, the emerging picture from available data reveals a dense system of overlapping transport throughout the plant. Here we add to this picture by characterizing the sole Arabidopsis thaliana NCS gene (AtNCS1) through in planta radiolabel studies using insertion mutants as well as through heterologous complementation studies in yeast.

2. Materials and methods

2.1. Arabidopsis genetic stocks and growth conditions

Arabidopsis lines WiscDsLox419Co3 [ABRC # CS854962] [21], and Gk-266G10 [ABRC# CS305244] [22], were obtained from Arabidopsis Biological Resource Center (Columbus, OH, USA) and line ET8162 from Cold Spring Harbor Laboratory (Cold Spring Harbor, NY, USA) [23]. Growth conditions were as previously described [19]. For the allantoin growth experiments, MAM was used in which 5 mM allantoin was the sole nitrogen source as previously described [11].

2.2. Molecular manipulations

Arabidopsis DNA was extracted using DNeasy Plant Mini Kit (Qiagen, Valencia, CA, USA) or QuickExtract (Epicentre, Madison, WI, USA). Individual plants from insertion lines Gk-266G10, ET8162 and WiscDsLoc419Co3 were genotyped by multiplex PCR using gene-specific primers (F12E4C, F12E4G, LP-03555 or RP-03555) and T-DNA specific primers (GABITDNA, Ds3-2 or P745) (Table 1). Total RNA was isolated using TRIzol Reagent (Invitrogen, Carlsbad, CA, USA) from tissues at different developmental stages. Reverse transcription was performed using Reverse Transcriptase M-MuLV (Roche, Basel, Switzerland) at 42°C for 2 h and then

Table 1

<table>
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<tr>
<th>Oligonucleotide primers used in this study.</th>
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Fig. 1. Amino acid sequence similarity and phylogenetic relationships of AtPRT1. (A) Alignment of AtNCS1 (GenBank # AA25608.1), a paralog from Vitis vinifera (XP_00264640.1) [VvNCS1] and the uracil transporter from Saccharomyces cerevisiae ScFUR4 (NP_009577) by ClustalW [45]. Black boxes represent amino acid identity while grey boxes indicate amino acid similarity. Predicted transmembrane spanning domains are indicated for AtNCS1 (light gray bars), VvNCS1 (dark gray bars) and ScFUR4 (black bars) using TMHMM [29]. (B) Phylogenetic relationships of select NCS1 proteins (in grey shaded area) from eukaryotes [Chlamydomonas reinhardtii (XP_00169432.1); Physcomitrella patens (XP_00175878.1); Picea glauca (UGCID:3879008); Plesiocystis pacifica SIR-1 (ZP_01906831.1); Oryza sativa (NP_001047676.1); Oryza sativa (XP_01481566.1); Ricinus communis (XP_002508829.1); Solanum lycopersicum (translated from AK323376); Zea mays (NP_001136335.1); and Saccharomyces cerevisiae ScDal4 (NP_012294), ScFUR4 (NP_009577), ScFur7 (NP_013338), ScFuj1 (NP_009311), and prokaryotes [Alcaligenes faecalis (ZP_01111046); Bacillus subtilis (ZP_0111046); bacterium Ellin514 (ZP_03630186.1); Escherichia coli (AP_001159.1); Paracoccus denitrificans (ZP_00631515); Synechococcus sp. PCC 7335 (ZP_05037138.1), Aeropyrum pernix (NP_147909.1)] with representative members of other Arabidopsis nucleobase transporters; AtAZG1 (AAQ65179.1), AAZG2 (BAB00940.1), AtENT1 (NP_564987), AtENT3 (NP_192421), AtNAT1 (Q5SH2.1), AtNAT2 (Q94C70.2), AtPUP1 (NP_174414.1), AtPUP2 (NP_973592.1), AtUPS1 (NP_563033.1), AtUPS2 (NP_178451.2). Red numbers at nodes denote branch support values and the scale bar denotes genetic distance of branch length. Pylogenetic tree constructed using Phylogeny.fr [46] using MUSCLE alignment parameter [47] and Tree construction using maximum likelihood [48–50].
AtNCS1/AtNcs1-1 (973 bp) representative of homozygous wild type individuals, lane 4 heterozygous for the insertion formed for 30 cycles of 94°C amplification with Taq DNA Polymerase (Qiagen) using oligonucleotide primers designed for the At5g03555 locus. (A) Physical map of locus seedlings (lane 6). (E) RT-PCR analysis of inflorescence (lane 4), mature leaf (lane 5), and day-old seedlings (lane 1), 10-day-old seedlings (lane 2), isolated flowers (lane 3), PCR analysis of AtNCS1/AtNCS1 family, an molecular size standards, lanes 2 and 3 show AtNCS and 7 show the genotypes of two homozygous individuals (lanes 4 and 5), while lanes 6 and 7 show the genotypes of two homozygous AtNcs1-1 individual (lane 3), AtNcs1-3 individual plants. (B) Genotype ET8162 (ET) T-DNA insertions. PCR fragment sizes of informative oligonucleotide combinations are given for genotyping and transcript detection. (C) Lanes 2–5 show genotypes of AtNcs1-1 segregating for the insertion AtNcs1-1. (C) Lanes 2–5 show genotypes of AtNcs1-1 segregating family, an AtNCS1/AtNCS1 wild type (lane 2), a heterozygous AtNCS1/AtNCS1-3 individual (lane 3), AtNcs1-1/AtNcs1-3 individuals (lanes 4 and 5), while lanes 6 and 7 show the genotypes of two homozygous AtNcs1-2 individual plants. (D) RT-PCR analysis of AtNCS1 (973 bp) and actin RNA isolated from Columbia wild type 5-day-old seedlings (lane 1), 10-day-old seedlings (lane 2), isolated flowers (lane 3), inflorescence (lane 4), mature leaf (lane 5), and AtNcs1-1/AtNcs1-1 10-day-old seedlings (lane 6). (E) RT-PCR analysis of AtNCS1 (lanes 1 and 3) and actin (lanes 2 and 4) RNA isolated from AtNcs1-3/AtNcs1-3 (lanes 1 and 2) and from Columbia wild type (lanes 3 and 4) 10-day-old seedlings.

Molecular analysis of the WISC line segregating for the insertion. Lane 1 contains his3 Δ:: kanMX4, his3 Δ1, leu2 Δ10, met15 Δ10, ura3 Δ10 [25] and NC122-sp6 [Mat a leu2 fur4-1] [20] were grown in YPD or on Synthetic Complete medium (SC) at 30°C. Yeast transformation was by the lithium acetate method treated with DNA-free (Ambion, Austin, TX, USA). Subsequent PCR amplification with Taq DNA Polymerase (Qiagen) using oligonucleotides LP-03555 and RP-03555 or AraCT2A and AraCT2B was performed for 30 cycles of 94°C 30 s, 55°C 30 s and 70°C 2 min in a Mastercycler gradient machine (Eppendorf, Hamburg, Germany). Genomic DNA from Arabidopsis Columbia wild type was amplified with primers At5g03555YEA and At5g03555YEB and the resulting DNA fragment treated with Xho I and Sph I, ligated with Xho I/Sph I linearized pRG399 [19] to form plasmid pH369. Plasmid pH124 is composed of the pRG399 yeast expression vector containing a 1.8 kb insert consisting of an open reading frame that encodes for the 599 amino acid version of AtNCS1. This plasmid was constructed by PCR-amplifying bacterial artificial chromosome F12E4 containing genomic sequence of At5g03555 with oligonucleotides F12E4D and F12E4E. The resulting DNA fragment was digesting with Xho I and Sph I and ligated into the like cleaved sites on pRG399. The plasmid insert was subject to DNA sequence to verify sequence integrity.

2.3. Yeast cultures and transformation

Saccharomyces cerevisiae strains FY1679-5C [MAT a leu2 Δ1 his3 Δ200 ura3 Δ52] [24] and RG191 [Mat a, fcy2 Δ1:: kanMX4, his3 Δ1, leu2 Δ10, met15 Δ10, ura3 Δ10] [25] and NC122-sp6 [Mat a leu2 fur4-1] [20] were grown in YPD or on Synthetic Complete medium (SC) at 30°C. Yeast transformation was by the lithium acetate method.
Sensitivity to toxic nucleobase analogues was assayed by adding filter-sterilized stock solutions to the growth media.

2.4. Radiolabel uptake experiments

Five- and eleven-day-old seedlings were transferred from MAM onto MAM supplemented with [8-3H]guanine (3.7 kBq ml−1), [2,8-3H]adenine (3.7 kBq ml−1) or [5,6-3H]uracil (12.33 kBq ml−1) (Moravek Biochemicals, Brea, CA) or MAM alone as previously described [19]. After 2 days of growth the plants were carefully removed and weighed. Three 75 mg samples of selected genotype tissue were homogenized in 0.5 ml 20 mM Tris–HCl, pH 8. After homogenization, samples were centrifuged at 11,750 g for a minute. For each sample, the radioactivity in 0.1 ml of extract was determined by mixing with 3 ml EcoLume (MP Biochemicals,Solon, OH) and measured by using a scintillation counter. For radiolabeled uptake experiments in yeast, cells were grown to OD600 = 4 and incubated for 0 and 2.5 min with 0.5 μM of one of the isotopes mentioned above in 100 mM citrate buffer (pH 3.5) with 1% glucose. Fifty microliter aliquots were added to 4 mL of ice-cold water and filtered through a 0.45 μm Metricel membrane filter (Gelman Sciences, Ann Arbor, MI). Filters were then washed with 8 ml of water and radioactivity was measured by scintillation counter.

2.5. Transport kinetics of AtNCS1 expressed in yeast

Time course for the uptake of 0.5 μM [5,6-3H]uracil was performed with yeast cells (OD600 = 4) expressing AtNCS1-1. Substrate saturation kinetics was established for varying concentrations of [5,6-3H]uracil and [2,8-3H]adenine at 2.5 min using yeast cells (OD600 = 4) expressing AtNCS1-1. Radiolabel uptake was assayed as mentioned above.

2.6. Radiou nucleobase uptake by yeast expressing AtNCS1-1 with a protonophore

Yeast suspensions were prepared as described [19]. Equal volumes (50 μL) of yeast suspension (OD600 = 2) and buffer containing 1.0 μM [5,6-3H]uracil (12.33 kBq ml−1) and 100 μM carbonyl cyanide m-chlorophenylhydrazone (CCCP) in 100 mM citrate buffer (pH 3.5) with 1% glucose. For control, CCCP was omitted from the reaction mixture. Assay of radiolabel uptake was performed at 0 and 2.5 min as mentioned above.

3. Results and discussion

3.1. Arabidopsis locus At5g03555 encodes for AtNCS1

The Arabidopsis genome contains a locus (At5g03555) that encodes for a protein with sequence and structural characteristics common with members of the purine related transporters or nucleobase cation symporter 1 (NCS1) superfamily. At5g03555 is therefore renamed as the A. thaliana nucleobase cation symporter 1 or AtNCS1. Despite the extensive evolutionary distance between plants and yeast, AtNCS1 shares substantial amino acid similarity.

Table 2

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<tr>
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<td>*</td>
</tr>
<tr>
<td>8-Azaguanine</td>
<td>*</td>
</tr>
<tr>
<td>5-Bromo-2’-deoxyuridine</td>
<td></td>
</tr>
<tr>
<td>5-Fluorocytosine</td>
<td></td>
</tr>
<tr>
<td>5-Fluoro-2’-deoxyuridine</td>
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</tr>
<tr>
<td>5-Fluororotic acid</td>
<td></td>
</tr>
<tr>
<td>5-Fluourouracil</td>
<td>*</td>
</tr>
<tr>
<td>Pyrithiamine</td>
<td></td>
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<tr>
<td>Allantoin</td>
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</table>

[26] Resistance of AtNcs1-1 mutants to a panel of toxic purine and pyrimidine analogues incorporated in the agar growth medium (+ = resistance; − = sensitive).

Fig. 4. Uptake of [3H]-adenine, -guanine, and -uracil by AtNcs1-1 mutant and Columbia wild type seedlings. Radio-labeled purine uptake was measured in extracts from 11-day-old seedlings (A). Radio-labeled uptake of uracil was measured in extracts of 11-day-old (B) and 5-day-old (C) seedlings. Values shown are the mean of three independent experiments. Error bars indicate the standard error of the mean. Statistical analysis used an independent paired t-test. Significance was measured at P = 0.05.
(20–22% amino acid identity and 52–54% amino acid similarity) to the well characterized S. cerevisiae NCS1 FUR4 family including FUR4 (uracil transporter), DAL4 (allantoin transporter), FUI1, (uridine transporter), THI7, (thiamine transporter) and lesser levels of sequence similarity to the related NCS1 FCY2 family including the cytosine–adenine transporters FCY2, 21 and 22 and vitamin B6 transporter, TPN1 [Fig. 1A] [15,16,27,28]. Secondary structure predictions suggest that AtNCS1 and a closely related orthologous protein from Vitus vinifera, are membrane localized proteins showing twelve predicted membrane spanning domains (Fig. 1A) [29]. Positions of the predicted transmembrane spanning domains in AtNCS1 and VvNCS1 overlap the transmembrane spanning domains of FUR4 (Fig. 1A). In addition, key amino acids conserved among NCS1 proteins are also conserved in AtNCS1. Recently, several tertiary structures were resolved for the benyl-hydantoin transporter (Mhp1), a NCS1 from Microbacterium liquefaciens, aiding in the development of a molecular dynamic simulation model and the identification of likely solute and cation interacting amino acids [30,31]. AtNCS1 shares 30 of 34 amino acids highly conserved among NCS1 proteins as well as four of six solute interacting amino acids and two of five cation interacting amino acids. Together these data solidly place AtNCS1 in the NCS1 super family.

NCS1 members are ubiquitous in nature and the plant kingdom is no exception. Fig. 1B details a representative phylogenetic tree detailing the relationship between AtNCS1 amino acid sequence and likely orthologous proteins from across the plant and microbial kingdoms. The order of AtNCS1 sequence similarity among orthologous plant proteins mirrors the hierarchy of taxa throughout the plant kingdom showing the highest similarity to dicots (Vitus, Rincus, Populus and Solanum), followed by monocots (Zea and Oryza), gymnosperm (Picea), nonflowering moss (Physcomitrella) and chlorophyte algae (Chlamydomonas, Osterococcus). Significant, but more distant similarities are observed between plant NCS1 proteins and orthologous proteins from diverse sources of bacteria (Escherichia, Plesiocystis, Bacillus, Paracoccus, Synchococcus, Alteromonas, bacterium Elinn514), archaeobacteria (Aeropyrum), and S. cerevisiae. All of the NCS1 proteins locate together in a distinct clade. AtNCS1 is different from other Arabidopsis nucleobase and nucleoside transporters here represented by two members each of the Atnat, Atups, Atent, Atpup and Atazg families (Fig. 1B).

3.2. Molecular characterization of the AtNCS1 locus

The AtNCS1 locus is among a minority of nuclear genes in the Arabidopsis genome that appears to have no introns. Further this locus contains two in frame ATG codons which predict either a long 1800 bp open reading frame (orf1) starting from ATG-1 that encodes for a 598 amino acid protein, or a shorter 1509 bp ORF (orf2) with an origin at ATG-2 that encodes for a 501 amino acid protein (Fig. 2A). No verified full length cDNA including the long coding region placed under constitutive transcriptional control (pRH369) were grown on nutrient media with uracil as the sole nitrogen source is presented in (E). RG191 and RG 191 + pRH369 were incubated [3H]-adenine, -guanine, or -uracil in citrate buffer (pH 3.5) and aliquots were taken at 0 and 2.5 min. Values shown are the mean of at least three independent experiments. Error bars indicate the standard error of the mean. Statistical analysis used an independent paired t-test. Significance was measured at *P ≤ 0.05.

Fig. 5. Growth of Saccharomyces cerevisiae expressing AtNCS1 on toxic purine or pyrimidine analogs and uptake of [3H]-adenine, -guanine, and -uracil. The growth pattern of yeast strains: deficient for the adenine-cytosine transporter fcy2 (RG191); deficient for the uracil transporter fur4 (NC122sp6), wild type for FCY2 and FUR4 (FY1679-5C); with an empty expression vector (pRG399); or with the AtNCS1 coding region placed under constitutive transcriptional control (pRH369) were grown on nutrient media with 5-fluorocytosine (A), 8-azaguanine (B), 8-azaadenine (C) or 5-fluorouracil (D). The growth pattern of NC122sp6 alone or containing pRH369 on nutrient media with uracil as the sole nitrogen source is presented in (E). RG191 and RG 191 + pRH369 were incubated [3H]-adenine, -guanine, or -uracil in citrate buffer (pH 3.5) and aliquots were taken at 0 and 2.5 min. Values shown are the mean of at least three independent experiments. Error bars indicate the standard error of the mean. Statistical analysis used an independent paired t-test. Significance was measured at *P ≤ 0.05.
3.3. AtNCS1 null mutants show resistance to certain purine and pyrimidine analogs

Three different homozygous T-DNA insertion mutants in the AtNCS1 locus were tested for their ability to germinate and grow in a range of toxic purine or pyrimidine analog concentrations. In comparison to wild type, homozygous AtNcs1-1 mutants displayed resistance to toxicity of 8-azaadenine, 8-azaguanine, and 5-fluorouracil (Fig. 3A–D). AtNcs1-1 displayed 2.3-fold more resistance to 8-azaadenine than wild type (wild type I50 = 3 μM; AtNcs1-1 I50 = 7 μM) (Fig. 3D). However, AtNcs1-1 displayed lower levels of resistance to 8-azaadenine, only 0.7-fold resistance compared to wild type (wild type I50 = 7 μM; AtNcs1-1 I50 = 10 μM) (Fig. 3C). This differential level of sensitivity to 8-azaadenine and 8-azaadene can be clearly demonstrated at 5 μM and 10 μM. For wild type, at 5 μM, seedling fresh weight was 62% of the control in case of 8-azaadenine and only 10% in case of 8-azaguanine while for AtNcs1-1 it was 70% of the control for 8-azaadenine (data not shown). This result and the presence of an AtNCS1-specific transcript in mutant tissue resulted in no further investigation of this line.

3.4. In planta radio-labeled nucleobase uptake studies

In planta radiolabeled nucleobase uptake experiments were performed and show that the uptake of [3H]-adenine and -guanine by 11- to 13-day-old seedlings was significantly reduced in AtNcs1-1 compared to Columbia wild type (Fig. 4A). Compared to wild type, AtNcs1-1 uptake of [3H]-guanine was reduced by 50% while uptake of [3H]-adenine was reduced by 40% (Fig. 4A). On the other hand, in seedlings of the same developmental stage uptake of [3H]-uracil was significantly reduced in the wild type compared to AtNcs1-1 (Fig. 4B). Uptake of [3H]-uracil by wild type was reduced by about 20% when compared to AtNcs1-1 (Fig. 4B). This seemingly contradictory result suggests that AtNCS1 may not act as the main transport route for uracil at this developmental stage, but that other transporters are then more active. Indeed AtUPS1 and 2 and to a lesser extent AtAgA2 transport uracil [12,19]. Expression data shows that 11- to 13-day-old seedlings have appreciable AtUPS2 expression whereas 5- to 7-day-old seedlings have signific-

![Fig. 6.](image-url) [3H]-uracil uptake by Saccharomyces cerevisiae expressing AtNCS1. Yeast strain NC122-sp6deficient in the uptake of uracil and containing pH369 was incubated with [5,6-3H]uracil in the presence or absence of protonophore CCCP and aliquots were taken at 0 and 2.5 min. Values shown are the mean of at least three independent experiments. Error bars indicate the standard error of the mean. Statistical analysis used an independent paired t-test. Significance was measured at *P = 0.05.*
cantly less expression of AtUPS1 and 2 when AtNCS1 is expressed (Fig. 2D) [12]. Radiolabel uracil uptake experiments were conducted with 5-day-old seedlings and revealed that AtNCS1 accumulates a significantly smaller amount of \([3\text{H}]\text{uracil}\) compared to wild type (Fig. 4C). The result suggests that AtNCS1 is an important source of uracil transport in 5- to 7-day-old seedlings. Interestingly at germination and the first week of seedling development, nucleobase salvage – particularly uracil salvage – is essential for seedling development with strong expression of genes encoding pyrimidine salvage enzymes [6,7,43]. As the seedling develops after this initial stage there is a shift toward de novo pyrimidine synthesis and recycling of pyrimidine nucleotides increases [43,44]. The shift in the rate of nucleobase salvage/synthesis versus catabolism may also be reflected in the presence and activities of different nucleobase transporters. It appears that at 11 days, uracil transport in the seedling relies less on AtNCS1 and more on AtUPS2 or other transporters.

3.5. Heterologous complementation studies confirm that AtNCS1 is a nucleobase transporter

Heterologous complementation studies in S. cerevisiae were performed to test whether AtNCS1 could act in a transport capacity outside of a plant. Yeast expression plasmids containing Arabidopsis At5g03555 sequences which encode for a long AtNCS1 protein of 598 amino acids, pRH124, and sequences encoding for a short AtNCS1 protein of 501 amino acids, pRH369, were transformed into two yeast strains deficient in either fcy2 (adenine–cytosine–guanine transport) or fur4 (uracil transport). The short version encodes for a 502 amino acid protein devoid of predicted chloroplast transit and signal sequences present in the first ~99 amino acids. Yeast strains deficient for fcy2 and harboring either pRH124 (data not shown) or pRH369 (Fig. 5B and C) displayed heightened sensitivity to growth on 8-azaadenine, 8-azaguanine. However, strains harboring pRH369 displayed appreciably more sensitivity than those harboring pRH124, suggesting that either the shorter AtNCS1 protein functions better in yeast and/or that there is a steady state higher level of AtNCS1 short protein compared to long AtNCS1 in the respective strains. Although it is evident that both the long and short AtNCS1 forms function in yeast, further characterization was conducted with yeast strains harboring pRH369.

The yeast expression vector pRH369 containing sequences encoding for a short AtNCS1 protein was transformed into two yeast strains deficient in either fcy2 (adenine-cytosine-guanine transport) or fur4 (uracil transport). Yeast strains were grown on media containing 5-fluorocytosine, 8-azaadenine, 8-azaguanine, 5-fluorouracil or with uracil as the sole nitrogen source. Results shown in Fig. 5 indicate that yeast stains harboring pRH369 regain

![Fig. 7. Substrate kinetics of AtNCS1 expressed in yeast. (A) Time course of the transport of \([5,6-3\text{H}]\text{uracil}\). (B) Transport rate of varying concentrations of \([5,6-3\text{H}]\text{uracil}\). (C) Lineweaver–Burk plot for \([5,6-3\text{H}]\text{uracil}\) uptake shown in (B). (D) Transport rate of varying concentrations of \([2,8-3\text{H}]\text{adenine}\). (E) Lineweaver–Burk plot for \([2,8-3\text{H}]\text{adenine}\) uptake shown in (D).](image-url)
sensitivity to growth on 8-azaguanine, 8-azaadenine and 5-fluorouracil, but not to growth on 5-fluorocytosine when compared to control strains deficient in fcy2 or fcy4 (Fig. 5A–D). The uracil-transport-deficient yeast strain is unable to grow on uracil as a sole nitrogen source but regains this ability with pH369 (Fig. 5E). Additionally, yeast strains deficient in fur4 or fcy2 and transformed with pH369 showed significantly increased uptake of [3H]-guanine, [3H]-adenine as well as [3H]-uracil when compared to the control strain (Fig. 5F–H). These results match the toxic analog growth studies and radio-labeled nucleobase uptakes studies in planta.

In the presence of the protonophore CCCP, [3H]-uracil uptake by the yeast strain deficient in fur4 and harboring pH369 is strongly inhibited (Fig. 6). This result is consistent with AtNCS1 functioning as a nucleobase cation symporter.

A time course experiment for [3H]-uracil uptake by yeast expressing pH369 revealed a linear relationship up to 10 min (Fig. 7A). As we have shown here AtNCS1 displays a distinct solute transport from fungal NCS1 proteins. AtNCS1 transports guanine, adenine, and uracil (Figs. 4 and 5) whereas fungal NCS1 proteins transport either guanine and adenine (FCY2) or uracil (FUR4). Because of that we examined the transport kinetics of [3H]-adenine and [3H]-uracil in yeast cells expressing pH369 at 2.5 min when the uptake of [3H]-uracil was within the linear range (Fig. 7A). Increasing [3H]-uracil and [3H]-adenine concentrations showed an initial linear rate but approached saturation at higher concentrations (Fig. 7B and D). Lineweaver–Burk plots revealed $K_{mapp} = 7.0$ μM and $V_{max} = 0.3$ (nmol/10^9 cells/min) for uracil (Fig. 7C) and $K_{mapp} = 7.9$ μM and $V_{max} = 3.3$ (nmol/10^9 cells/min) for adenine (Fig. 7E).

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References


