NO$_3^-$/$H^+$ Antiport in the Tonoplast of Cucumber Root Cells Is Stimulated by Nitrate Supply: Evidence for a Reversible Nitrate-Induced Phosphorylation of Vacuolar NO$_3^-$/$H^+$ Antiport

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Abstract

Studies in the last few years have shed light on the process of nitrate accumulation within plant cells, achieving molecular identification and partial characterization of the genes and proteins involved in this process. However, contrary to the plasma membrane-localized nitrate transport activities, the kinetics of active nitrate influx into the vacuole and its adaptation to external nitrate availability remain poorly understood. In this work, we have investigated the activity and regulation of the tonoplast-localized H$^+$/NO$_3^-$ antiport in cucumber roots in response to N starvation and NO$_3^-$ induction. The time course of nitrate availability strongly influenced H$^+$/NO$_3^-$ antiport activity at the tonoplast of root cells. However, under N starvation active nitrate accumulation within the vacuole still occurred. Hence, either a constitutive H$^+$-coupled transport system specific for nitrate operates at the tonoplast, or nitrate uses another transport protein of broader specificity to different anions to enter the vacuole via a proton-dependent process. H$^+$/NO$_3^-$ antiport in cucumber was significantly stimulated in NO$_3^-$-induced plants that were supplied with nitrate for 24 hours following 6-day-long N starvation. The cytosolic fraction isolated from the roots of NO$_3^-$-induced plants significantly stimulated H$^+$/NO$_3^-$ antiport in tonoplast membranes isolated from cucumbers growing on nitrate. The stimulatory effect of the cytosolic fraction was completely abolished by EGTA and the protein kinase inhibitor staurosporine and slightly enhanced by the phosphatase inhibitors okadaic acid and cantharidin. Hence, we conclude that stimulation of H$^+$/NO$_3^-$ antiport at the tonoplast of cucumber roots in response to nitrate provision may occur through the phosphorylation of a membrane antiporter involving Ca-dependent, staurosporine-sensitive protein kinase.

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Introduction

Nitrate variation in soils strongly influences plant growth and development, so plants had to evolve mechanisms allowing them to react and respond to frequent changes of the NO$_3^-$ level. Namely, plants acquire NO$_3^-$ ions through different types of uptake systems operating in their roots: low (LATS) and high (HATS) affinity transport systems which have both constitutive and inducible components [1-6]. All four nitrate transport systems coexist within the cell and each exhibits diffusionality to nitrate and operates at a different external NO$_3^-$ concentration range. The constitutive high-affinity transport system (cHATS) operates at low external nitrate concentrations and is complemented by the inducible high-affinity system (iHATS), which is activated by very low NO$_3^-$ supply [3], [7]. When nitrate supply reaches greater values, the low affinity system (LATS) takes over [8], [9]. For the last decades much attention has been given to understanding how these transport systems are regulated at the molecular level. The passage of NO$_3^-$ through the plasma membrane of root cells has been particularly intensively studied. It is well known that nitrate uptake into the root cells occurs via energy dependent symport with two protons translocated along with one NO$_3^-$ ion against transmembrane gradient of nitrate [10], [11]. The energy for this process comes from the electrochemical gradient generated by plasma membrane proton pump H$^+$-ATPase [10–12]. In the plasma membranes of root cells a few transporters belonging to NRT1 (Nitrate transporter 1) and NRT2 (Nitrate transporter 2) families use the energy provided by H$^+$-ATPase to transport nitrate into the cells. Two Arabidopsis NRT1 proteins (AtNRT1.1 and AtNRT1.2) and two NRT2 proteins (AtNRT2.1 and AtNRT2.2) are directly involved in nitrate uptake but they participate in different-affinity uptake systems [13–16]. AtNRT2.1 and AtNRT2.2 are involved in high-affinity NO$_3^-$ influx, whereas AtNRT1.2 is a low-affinity nitrate transporter [4], [14], [17]. AtNRT1.1 functions as a dual-affinity transporter: the phosphorylated form of the protein is involved in high-affinity nitrate uptake, whereas the dephosphorylated form contributes to low-affinity NO$_3^-$ influx [5], [6], [18], [19]. The posttranslational modification relying on phosphorylation/dephosphorylation of the nitrate transport proteins is not surprising since the putative amino acid sequences of NRT1 and NRT2 families
contain a number of conserved protein kinase C recognition motifs in their N- and C-terminal domains or in the central loops [20]. This may indicate the importance of phosphorylation and dephosphorylation events in the regulation of other members of NRT families.

Although the mechanism of regulation of nitrate influx into the cell has been extensively studied, the way by which nitrate sequestration within the vacuole is controlled is still unknown, and the question arises whether similar regulation mechanisms occur for the tonoplast and plasma membrane nitrate transport activities. It was earlier shown that plants provided with unlimited supplies of NO\textsubscript{3}\textsuperscript{-} stored most root and shoot nitrate within the vacuoles [21]. Nitrate was also accumulated in the central vacuole of leaf cells [22] or root tip cells [23] and the vacuolar NO\textsubscript{3}\textsuperscript{-} concentration increased in response to high external nitrate concentrations [2, 23]. The mechanism of NO\textsubscript{3}\textsuperscript{-} transport across the tonoplast has been recently more extensively studied. It was earlier assumed that since the ∆Ψ between the cytosol and the vacuole is not sufficient to drive the NO\textsubscript{3}\textsuperscript{-} accumulation within the vacuole, the flux of nitrate across the tonoplast must occur via both passive and active ways. Indeed, the experiments using direct and indirect assay of NO\textsubscript{3}\textsuperscript{-} or H\textsuperscript{+} transport have provided evidence for an NO\textsubscript{3}\textsuperscript{-}/H\textsuperscript{+} antiporters in the tonoplast [24–27]. There are two proton pumps (H\textsuperscript{+}-ATPase and H\textsuperscript{+}-PPase) on tonoplast which pump H\textsuperscript{+} into vacuole and thus contribute to building an electrochemical proton gradient between cytoplasm and vacuolar lumen [28–30]. The pH gradient generated by both pumps serves as a driving force for ion antiport at tonoplast membrane [30]. V-ATPases are highly conserved, multisubunit proton pumps that are built of two domains: the membrane-integral V0 complex of subunits VHA-a, -c, -ε, -c′, -d, and –e, which is responsible for proton translocation into the lumen of endomembrane compartments and the peripheral V1 complex of subunits VHA-A to -H, involved in ATP hydrolysis [31]. In contrast, the VPPase functions as a homodimer of a single polypeptide, and uses energy from the pyrophosphate (PPi) hydrolysis to drive proton transport across membranes [32], [33]. However, since PPi is a by-product of several biosynthetic processes, it has been suggested that the V-PPase may be the major proton pump responsible for the acidification of vacuoles of young, growing cells [34]. It has also been proposed that V-PPase may function as a supporting system for V-ATPase under stressful conditions limiting the availability of ATP [33]. Despite the functional relation of these two pumps remains to be established, it has been already shown that a proton motive force generated by V-ATPase can be directly used to determine transport activity. The high purity and activity of the isolated vacuoles permitted the investigation of nitrate transport in vacuoles [23] and enabled the determination of NO\textsubscript{3}\textsuperscript{-} transport rates in intact vacuoles [21]. Studies on the effect of nitrate, nitrogen induction or nitrogen starvation on the rate of NO\textsubscript{3}\textsuperscript{-} transport across the tonoplast reveal that nitrate itself may regulate the activity of nitrate transporters localized at the vacuolar membrane. We also show that phosphorylation/dephosphorylation events may be involved in regulation of the nitrate transport process leading to NO\textsubscript{3}\textsuperscript{-} accumulation within the vacuole lumen.

Materials and Methods

Plant Material

All experiments were performed using the roots of 7-day-old seedlings of Cucumis sativus L var. Krak. Seeds were germinated in darkness and then grown on nutrient solution, pH 6.0, with or without nitrate. The experimental material included plants grown on NO\textsubscript{3}\textsuperscript{-}-rich (NO\textsubscript{3}\textsuperscript{-}-grown plants) or N-free (N-deprived plants) medium for 7 days and plants grown on N-free medium for 6 days and transferred on NO\textsubscript{3}\textsuperscript{-}-rich medium for 24 hours (NO\textsubscript{3}\textsuperscript{-}-induced plants). The standard medium contained the following macroelements: 1 mM K\textsubscript{2}SO\textsubscript{4}, 0.7 mM CaSO\textsubscript{4}, 0.33 mM Ca(H\textsubscript{2}PO\textsubscript{4})\textsubscript{2}, 0.33 mM MgSO\textsubscript{4} with 5 mM KNO\textsubscript{3} (NO\textsubscript{3}\textsuperscript{-}-rich medium) or 2.5 mM K\textsubscript{2}CO\textsubscript{3} (N-free medium). Microelements added to the medium included: 25 μM ferric citrate, 3 μM MnSO\textsubscript{4}, 1.7 μM H\textsubscript{2}BO\textsubscript{3}, 0.3 μM CuSO\textsubscript{4}, 0.003 μM ZnSO\textsubscript{4} and 0.017 μM Na\textsubscript{2}MoO\textsubscript{4}. All plants were grown under a 16-h photoperiod (180 mmol m\textsuperscript{-2} s\textsuperscript{-1}) at 25°C during the day and 22°C during the night.

Tonoplast Isolation

The membranes and the soluble (cytosol) fraction were isolated after homogenization of fresh roots in 70 mM TRIS-MES (pH 8.0), containing 500 mM sorbitol, 10 mM sodium glycerophosphate, 280 mM choline chloride, 2 mM salicylhydroxamic acid, 3 mM Na\textsubscript{2}EDTA, 26 mM K\textsubscript{2}SO\textsubscript{4}, 5 mM DTT, 0.2% BSA and 0.5% PVPP, followed by 15-min centrifugation at 10 000 g. The total microsomal membranes were collected through the subsequent ultracentrifugation at 80 000 g for 30 min. The supernatant obtained after this centrifugation was used as a soluble cytosolic fraction. Vascular membranes were separated by centrifugation of microsomes on a discontinuous sucrose density gradient (20, 28, 32 and 42% w/w) as described earlier [35]. The fraction enriched in tonoplast displaying the highest activity of PPase and V-ATPase (band of 20% sucrose) was diluted 4 times with 5 mM Tris-MES (pH 7.2), 1.1 M glycerol, 1 mM Na\textsubscript{2}EDTA and 1 mM DTT, and further recentrifuged at 80 000 g for 30 min. The final pellet of vascular membranes was resuspended in a 20 mM Tris-MES containing 250 mM sucrose and immediately used to determine transport activity. The high purity and.

The research on the molecular identity of antiporters that use the electrochemical gradient generated by V-ATPase to drive nitrate transport into vacuole are now in progress. The evidence has been presented that some proteins of the chloride channel family (CLC) reside at tonoplast and act as NO\textsubscript{3}\textsuperscript{-}/H\textsuperscript{+} exchangers [18], [37], [38]. Specific nitrate transporters (NRT1 and NRT2) may also be implicated in nitrate accumulation into the vacuole. It was shown that tonoplast-localized ANRT1.2.7 protein may be important for nitrate loading into the vacuole of dry seeds during seed maturation [39]. Since nitrate transporters are encoded by multigene families and the information about the localization of nitrate transporters in other plants is still scarce, we cannot exclude the contribution of other proteins of both NRT families in nitrate accumulation within the vacuole.

Although protein candidates have been proposed that could catalyze active nitrate influx into the vacuole, the regulation of this process remains poorly understood. The storage of nitrate within vacuoles is significant for both osmotic purposes and redistribution to the cytoplasm under starvation [22]. Since active accumulation of NO\textsubscript{3}\textsuperscript{-} in the vacuole under unlimited NO\textsubscript{3}\textsuperscript{-} supply may be considered as one of the mechanisms enabling plants to survive further NO\textsubscript{3}\textsuperscript{-}-limited conditions, it is highly desirable to understand the regulation of this process. In this work, we try to elucidate the mechanism of regulation of nitrate accumulation in tonoplast membranes isolated from cucumber root cells. Our studies on the effect of nitrate, nitrogen induction or nitrogen starvation on the rate of NO\textsubscript{3}\textsuperscript{-} transport across the tonoplast reveal that nitrate itself may regulate the activity of nitrate transporters localized at the vacuolar membrane. We also show that phosphorylation/dephosphorylation events may be involved in regulation of the nitrate transport process leading to NO\textsubscript{3}\textsuperscript{-} accumulation within the vacuole lumen.
the right-side out orientation of tonoplast membranes were determined earlier [40].

Assays of ATP Hydrolysis and ATP-dependent Proton Transport

The rate of ATP hydrolysis was determined from the differences in the rate of Pi release in the absence and presence of specific V-ATPase inhibitor 50 mM KNO₃. Inorganic Pi release was determined spectrophotometrically as described by Gallagher and Leonard [41] and Ames [42]. ATP-dependent proton transport activity was measured as a drop of acridine orange absorbance at 495 nm as described earlier [35]. After 5 min incubation of 50 µg vesicle protein with 20 mM TRIS-MES (pH 7.2), 0.25 M sucrose, 50 mM KCl, 1 mM DTT and 10 µM acridine orange, the reaction was initiated by addition of 3 mM MgATP. In control assay, 3 mM MgSO₄ instead of 3 mM MgATP was added to the reaction medium to determine the passive permeability of tonoplast membranes to protons (Figure S1A). In the assay including MgATP, bafilomycin, KNO₃ and gramicidin were used to confirm that the observed ΔA₄₉₅ changes result from V-ATPase activity (Figure S1B). 5 mM KNO₃ was added to MgATP-energized vesicles to initiate H⁺-coupled NO₃⁻ influx into membranes accompanied by the increase in acridine orange absorbance at 495 nm, in order to confirm the presence of nitrate antiport in tonoplast isolated from cucumber root cells (Figure S2).

Assays of ΔpH-dependent Proton Transport

To determine anions antiport activity in vacuolar membranes tonoplast vesicles were suspended in pH 6.0 (to impose transmembrane pH gradient, Figure S5) or 8.0 (to maintain equal transmembrane pH, Figure S3) and introduced to 20 mM TRIS-MES (pH 8.0), containing 0.25 M sucrose, 1 mM DTT and 10 µM acridine orange (absorbance assay) or 10 µM quinacrine (fluorescence assay, Figure S5). After 3 min incubation, the reactions were initiated by the addition of different concentrations of KNO₃, K₂SO₄ or KCl to the assay and monitored for the next three minutes. The changes in acridine orange absorbance were measured at 493 nm whereas the quinacrine fluorescence was determined with excitation and emission wavelengths of 420 nm and 500 nm, respectively, using a TD-700 fluorometer (Turner Designs).

The results obtained using two different pH-sensitive probes were comparable, hence only the acridine orange absorbance assay was used in the further studies on nitrate antiport activity. To determine whether tonoplast nitrate transporters and proton pump undergo phosphorylation/dephosphorylation event, 50 µl of soluble cytosolic fractions, 5 µM staurospermine and 2 µM phosphatases inhibitors (okadaic acid (OA) or cantharidin) or 5 mM EGTA were applied to the reaction media at the beginning of 3-min incubation of vesicles with acridine orange - containing mixture. In the experiments where the inhibitors of protein kinases or phosphatases were used, an equal amount of their solvent, DMSO was applied to control assays. In the experiments including cytosolic fraction, an equal volume of water was added to control assays.

Analysis of Tonoplast Nitrate content by HPLC

Nitrate accumulation within tonoplast vesicles imposing artificial pH gradient was also determined using HPLC-ion chromatographic assays. The reaction mixture was composed as described above except there were no pH-sensitive probes included. Different concentrations of nitrate were introduced into the assays and after 5 min incubation, the reactions were filtered through 0.45 µm nitrocellulose membranes (Millipore) under vacuum to separate vesicles from the medium. Nitrate content in the filtrate was determined using High Performance Liquid Chromatography (HPLC) according to Thayer and Huffaker [43] using a PARTISIL 10 SAX column, with 30 mM NaH₂PO₄ buffer, pH 3.0, as the mobile phase. H⁺-coupled nitrate accumulation within tonoplast vesicles was calculated from the difference between nitrate determined in vesicles imposing pH gradient and nitrate measured in vesicles without transmembrane ΔpH.

Western Blot

Membrane protein (15 µg) was mixed with the 10 mM TRIS buffer containing 2% (w/v) SDS, 80 mM DTT, 40% (w/v) glycerol, 5 mM PMSE, 1 mM EDTA, and 0.05% (v/v) bromophenol blue and incubated at room temperature for 30 min. The samples were separated by SDS/PAGE using 10% acrylamide gel (Sigma) and stained with Coomassie Blue (Figure 1D) or transferred onto a nitrocellulose membrane (Schleicher & Schuell BioScience). The membrane was incubated with polyclonal antibodies against the subunit a of tonoplast V-ATPase (Agrisera) at room temperature for 1 h. After repeated washing, the membranes labeled with primary antibodies were further incubated for 1 h with secondary antibodies (conjugated to horseradish peroxidase, Agrisera) and visualized by staining with DAB.

Protein Determination

Protein was measured using the Bradford [44] method with BSA as a standard.

Database Searching

The complete genomic sequence from cucumber has recently been made available to the public in GenBank with accession code ACHRI01000000 [45]. NCBI database (whole-genome shotgun reads) was used for Blast searches of cucumber sequences with homology to the previously annotated Arabidopsis thaliana CLC-a, CLC-c and CLC-g sequences. Full cDNAs as well as the protein sequences encoded by the newly identified cucumber CLCs were generated using FGENESH [46] and Eukaryotic GeneMark.hmm [47]. The primers used for PCR and qPCR were designed to amplify highly specific products. The specificity of all PCR products was confirmed through sequencing and the partial ESTs were submitted into GenBank; they are available under the following accession numbers: JK525488 (CsCLCa), JK525489 (CsCLCb) and JK525490 (CsCLCc).

Expression Analyses

Total RNA was extracted from cucumber roots with TRI Reagent (Sigma) according to manufacturer’s instructions. The quantity and quality of RNA were determined using a NanoDrop spectrophotometer (Nyxor, ND-100). RNA integrity was checked on 2100 Bioanalyzer (Agilent) using RNA kits and the RNA Integrity Number (RIN) algorithm (Figure S4). Two microgramms of DNase-digested total RNA were used for cDNA synthesis using random primers and High Capacity cDNA synthesis Kit (Applied Biosystems). 1 µl of cDNA was used as template for standard PCR with Marathon Polymerase (Aka Biotechnology) according to the following programme: 94°C for 5 min followed by 30 cycles of 94°C for 30 s; 56°C for 30 s and 68°C for 1 min, with final extension at 72°C for 5 min. PCR products were ligated to pGEM T-Easy (Promega) and sequenced 3–4 times. The same cDNA used for gene isolation was used for real-time PCR analyses. PCR
amplification was carried out using LightCycler 2.0 (Roche) and SYBR Green DNA dye, in a total volume of 10 µl containing 1 µl of 5-fold-diluted cDNA template, 1 µl of each primer (10 µM), 2 µl of nuclease-free water and 5 µl of 2× Master SYBR® Green I B (A&A Biotechnology). The thermal cycling conditions were 95°C for 10 min followed by 45 cycles of 95°C for 10 s, 56°C for 10 s, 72°C for 15 s. The gene encoding clathrin adaptor complex subunit (CAG) was used as internal control with the following forward 5′-TTGCTTCTTTCTGTAAATGC-3′ and reverse 5′-TGAACCTCGTCAAATTTACACA-3′ primers. A negative control without cDNA template was included in the same PCR run for each primer pair. To confirm the specificity of amplification, melting curve analysis was performed allowing to identify putative unspecific PCR products (e.g., primer dimers, amplification, melting curve analysis was performed allowing to run for each primer pair. To confirm the specificity of control without cDNA template was included in the same PCR reaction mix contamination) and the RT-PCR products were sequenced. Successive dilutions of the sample with the lowest Ct were used as a standard curve. Amplification efficiency was around 2. For each of the two independent RNA extractions, measurements of gene expression were obtained in duplicate. Specific primers were carefully designed using LightCycler Probe Design Software 2 (Roche): CLCa(forward, 5′-TATTAGGGTTATTTCTCTGATCC-3′; reverse, 5′-TTTCAAGGAAATATGCACACAAAAGAAAAA-3′), CLCb(forward, 5′-CGTGAAGATGTTTTTCCTACTGTTGC-3′; reverse, 5′-CATGAAATTTGAGGAAATATTAG-3′), and CLCg(forward, 5′-TTCTCTCTAGTATGGAGTATGAGAGAC-3′; reverse, 5′-TTTGAACCTGGAATTTGCATTTAACC-3′).

Statistical Analysis

Data on proton and nitrate transport activities were analysed by one-way completely randomized ANOVA and means comparison was performed by t test at significance level of 0.05. A GraphPad Prism program (GraphPad Software, Inc) was used to fit the data directly to the Michaelis-Menten equation using nonlinear regression and to display data with Lineweaver-Burk plots. The qPCR data were analyzed by the ∆∆CT - method using the LightCycler® Software 4.1 (Roche). Paired student’s t test and ANOVA (Excel) were used to confirm statistical significance of difference in CqCLCs expression between plants grown under different nitrate supply. The quantitative calculation of Western Blot results was performed using ImageJ software (rsb.info.nih.gov/ij).

Results

The Effect of different Nitrate Supply on the Hydrolytic and Transport Activity of Tonoplast V-ATPase

In most research on membrane antiport activities, primary proton pumps have been used to provide the energy to drive ion transport across the lipid bilayer. In the preliminary assay, we studied the effect of varying nitrate availability or deprivation on tonoplast V-ATPase to gain an insight into the rate of proton gradient formation in membranes isolated from plants growing under different nitrate regimes. The generation and maintenance of proton gradient in isolated tonoplast vesicles were monitored using acridine orange as a ΔpH-sensitive probe. Upon addition of MgATP, the formation of a ΔpH in the tonoplast was observed as a decrease of acridine orange absorbance at 495 nm (Figures 1A and 1B, Figures S1A-B). ΔpH generation across membranes was inhibited by bafilomycin (500 nmol) or KNO3 (50 µM), specific inhibitors of tonoplast V-ATPase, confirming that the effect resulted from vacuolar pump activity (Figure S1B). The rate of proton gradient formation was significantly different between tonoplast membranes isolated from plants grown under various nitrate nutrition. As shown in Figures 1A and 1B, the highest level of proton gradient was generated in membranes isolated from NO3−-grown plants. In comparison, the rate of ΔpH formation in tonoplast obtained from NO3−-grown and N-deprived plants decreased to 75% and 60%, respectively. Despite significant stimulation of proton transport activity of tonoplast V-ATPase, the hydrolytic activity of this enzyme was only slightly affected (decreased to 90%) and the level of proton pump protein was decreased up to 60% in NO3−-induced plants (Figures 1C and 1E). In comparison, the hydrolytic activity and proton pump protein level in membranes isolated from N-deprived plants were much more reduced to almost 50% and 40%, respectively (Figures 1C and 1E). Therefore, the availability of nitrate in nutrient solution had a significant impact on the amount and activity of tonoplast V-ATPase and consequently on the generation of a transmembrane ΔpH, an essential force for the transport of various compounds, e.g. nitrate, between cytosol and vacuoles. In consequence, the N-deprived plants displayed markedly reduced growth of roots, hypocotyls and cotyledons in comparison to NO3−-grown seedlings, which even developed the first leaf after 6 day-long cultivation on NO3−-enriched medium. The rate of antiport activity strictly depends on the magnitude of the proton motive force; hence to equip tonoplast prepared from different plants with the same rate of transmembrane ΔpH, we imposed an artificial pH gradient on the vesicles (Figure S8). Thus, we could measure the direct effect of nitrate availability on vacuolar antiporters.

Uptake of NO3− into Tonoplast Vesicles

The uptake of NO3− by isolated tonoplast vesicles was measured either indirectly, through the measurement of proton fluxes reflecting nitrate antiport, or directly, by measurement of nitrate accumulation inside vesicles. In the assays employing pH-sensitive probes (acidine orange and quinacrine) different concentrations of KNO3 were added to the reaction media containing tonoplast membranes with imposed ΔpH and then acidine absorbance or quinacrine fluorescence was monitored for the next 3 minutes. The transport of nitrate into vesicles imposing a pH gradient caused an instant, ΔpH-dependent increase of quinacrine fluorescence (Figures S5A-C) or acidine absorbance (Figures 2A-C), which was not observed in the vesicles without transmembrane ΔpH (Figures S6A-F). The dissipation of ΔpH was dependent on both the source of tonoplast membrane and the concentration of nitrate added to the vesicles. As shown in Figures 2A-C and S5A-C, to initiate the increase of acidine orange absorbance or quinacrine fluorescence, different nitrate concentrations were required for the membranes obtained from plants differently supplied with nitrate. The first absorbance/fluorescence change induced by nitrate was observed at a similar 2.5 mM concentration for both membranes isolated from N-deprived and those from NO3−-grown roots, whereas in the case of tonoplast isolated from NO3−-induced roots, lower nitrate concentration (0.5–1 mM) was sufficient to induce proton efflux from the vesicles. Tonoplast isolated from NO3−-induced roots displayed not only higher sensitivity but also higher capacity for nitrate transport, showing the maximum increase of absorbance/fluorescence approximately 2–3 times higher (100%), when compared with tonoplast isolated from N-deprived (35%) or NO3−-grown (60%) plants (Figures 2 and S5). Interestingly, the maximal rate of absorbance recovery for membranes isolated from plants growing under all nitrate regimes was obtained with 10 mM KNO3. Therefore, the active nitrate transport across tonoplast membranes from roots of NO3−-induced plants was induced at different but saturated at similar nitrate concentration compared.
The Properties of NO$_3^-$ / H$^+$ Tonoplast Antiport

Figure 1. V-ATPase activity and quantity in tonoplast vesicles isolated from roots of cucumber plants grown under different nitrate supply. A–B. The rate of MgATP-dependent, bafilomycin-sensitive proton transport in tonoplast under constant (squares, grey bars) or temporary (triangles, dark bars) NO$_3^-$ supply and under N-deprivation (circles, white bars). Figure A is representative for the results obtained in six independent experiments. Figure B presents the average values ± SD of four independent experiments. Asterisks indicate a significant difference (P<0.05) between V-ATPase activity under different nitrate regime. NO$_3^-$-induced (dark grey bars) and N-deprived (white bars) plants. Data are the average values ± SD of three independent experiments. Asterisks indicate a significant difference (P<0.05) between V-ATPase activity under different nitrate regime. 15 µg of tonoplast proteins were separated by SDS-PAGE on a 10% linear acrylamide gel. The positions of PAGE molecular mass markers are shown in kilodaltons on the right of the gel image. E. Immunoblot of V-ATPase subunit a on tonoplast membranes obtained from plants grown under different nitrate regime. Presented picture is representative for the results obtained in three to four independent experiments.

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to the relevant process observed in the membranes isolated from NO$_3^-$-uninduced plants. Since the phenomenon of nitrate transport into tonoplast vesicles exhibited different saturation kinetics in the membranes isolated from plants differently supplied with nitrate, the apparent Km values also varied in a way dependent on the membranes tested. However, Michaelis–Menten kinetics of nitrate transport were the same in the tonoplast of both NO$_3^-$-induced and NO$_3^-$-grown plants with the apparent Km of ~5±1.2 mM (Figures 3A and S5D). In contrast, the nitrate antiport kinetics into membranes isolated from N-deprived plants were characterized by the apparent Km of ~10±2.12 mM (Figures 3A and S5D). The results clearly show that the overall kinetics of acridine absorbance and quinacrine fluorescence recovery were different after the nitrate additions to the vesicles obtained from differently growing plants. In all the membranes the rate of NO$_3^-$-induced ΔpH dissipation was the highest during 120 s after addition of nitrate into the medium assay, and during the next 60 s the velocity of acridine absorbance increase was considerably lower (Figures 2A–C and S5A–C). Nevertheless, in the membranes obtained from N-deprived plant roots, the overall rate of nitrate transport was slower when compared to the vesicles obtained from NO$_3^-$-treated plants. The results of direct determination of nitrate accumulation within tonoplast vesicles measured by high-performance liquid chromatography clearly confirmed the output of assays employing pH-sensitive probes (Figure 2D). The highest and the lowest rate of nitrate accumulation occurred in membranes isolated from NO$_3^-$-induced and N-deprived plants, respectively. Nitrate accumulation within the membranes was also concentration-dependent and saturable, showing the features of Michaelis–Menten kinetics with Km ~6±1.6 mM and ~12±1.9 mM for tonoplast obtained from NO$_3^-$-treated and N-deprived plants, respectively (Figure 3B).

Uptake of Cl$^-$ and SO$_4^{2-}$ into Tonoplast Vesicles

To determine whether the observed nitrate antiport with its characteristics is selective only for nitrate ions we used potassium chlorate and potassium sulfate instead of potassium nitrate in the assay. As shown in Figure 4, the influx of Cl$^-$ and SO$_4^{2-}$ into tonoplast vesicles isolated from plants grown under different nitrate regimes could also occur through a ΔpH-dependent antiport system. However, the kinetic profiles of NO$_3^-$, Cl$^-$ and SO$_4^{2-}$ antiport were clearly different. Similarly to nitrate antiport, sulfate H$^+$-coupled influx into membranes was concentration dependent, and saturable at 50–100 mM SO$_4^{2-}$ concentration (Figures 4A and 4B). The overall kinetics of sulfate transport in all tonoplast membranes exhibited highly comparable features with the same apparent Vmax and Km of ~20±2.8 mM (Figures 4A and 4B). In comparison, H$^+$-coupled Cl$^-$ transport into the vesicles was concentration dependent only in the lower range of external Cl$^-$ concentration (1–5 mM), showing the features of Michaelis–Menten kinetics with the same apparent Km of ~3±2.1 mM in all membranes (Figures 4C and 4D). Contrary
The Properties of $\text{NO}_3^-/\text{H}^+$ Tonoplast Antiport

to sulfate antiport activity, the $V_{\text{max}}$ of chloride transport was different under varying nitrate regimes, showing the highest value (the highest capacity for $\text{Cl}^-$ influx) in tonoplast isolated from plants grown under constant $\text{NO}_3^-$ supply and the same lowest value in membranes obtained from N-deprived or $\text{NO}_3^-$-induced plants (Figure 4D). Hence, it may be assumed that the $\text{NO}_3^-$, $\text{Cl}^-$ or $\text{SO}_4^{2-}$ antiport in tonoplast membranes is mediated by different transport proteins or by a protein with different affinity and selectivity to these ions.

The Effect of Cytosolic Fraction, Protein Kinase and Phosphatase Inhibitors and EGTA on the Proton and Nitrate Transport across Tonoplast Membranes

The increased nitrate antiport across root tonoplast membranes isolated from $\text{NO}_3^-$-induced plants could be attributed to $\text{NO}_3^-$-dependent post-translational modification of nitrate transport protein localized to the tonoplast membranes. Phosphorylation/dephosphorylation events catalyzed by protein kinases and protein phosphatases are the most common reverse modulations of protein activities in plant cells. Therefore, we have investigated whether protein kinases or phosphatases could be involved in the vacuolar nitrate antiport stimulation in $\text{NO}_3^-$-induced plants. For this purpose, the effect of soluble cytosolic fractions isolated from the roots of plants grown under different nitrate regimes on the $\text{NO}_3^-/\text{H}^+$ transport activity in tonoplast obtained from the roots of $\text{NO}_3^-$-grown plants was studied. The soluble fraction was introduced into the reaction media at the beginning of 5-min incubation of membranes in the reaction mixture containing acridine orange. The addition of cytosolic fractions obtained from $\text{NO}_3^-$-grown or $\text{NO}_3^-$-induced plants to the assay resulted in over 30% and over 65% stimulation of nitrate antiport activity, respectively (Figure 5A). In contrast, the soluble fraction obtained from N-deprived plants did not affect $\text{H}^+/\text{NO}_3^-$ transport significantly (Figure 5A). Due to its higher stimulatory effect, the cytosolic fraction obtained from $\text{NO}_3^-$-grown plants was used in the further studies on the posttranslational modulation of nitrate antiport activity. Specific inhibitors of protein kinases (staurosporine) and phosphatases (okadaic acid or cantharidin), as well as the calcium chelator EGTA, were added to the reaction media along with the cytosolic fraction in order to determine which component of the fraction affects nitrate antiport activity. Specific inhibitors of protein kinases (staurosporine) and phosphatases (okadaic acid or cantharidin), as well as the calcium chelator EGTA, were added to the reaction media along with the cytosolic fraction in order to determine which component of the fraction affects nitrate antiport activity. Specific inhibitors of protein kinases (staurosporine) and phosphatases (okadaic acid or cantharidin), as well as the calcium chelator EGTA, were added to the reaction media along with the cytosolic fraction in order to determine which component of the fraction affects nitrate antiport activity. Specific inhibitors of protein kinases (staurosporine) and phosphatases (okadaic acid or cantharidin), as well as the calcium chelator EGTA, were added to the reaction media along with the cytosolic fraction in order to determine which component of the fraction affects nitrate antiport activity. Specific inhibitors of protein kinases (staurosporine) and phosphatases (okadaic acid or cantharidin), as well as the calcium chelator EGTA, were added to the reaction media along with the cytosolic fraction in order to determine which component of the fraction affects nitrate antiport activity.

Figure 2. The activity of proton-coupled nitrate transport in the tonoplast membranes isolated from cucumbers grown under different nitrate supply. 50 $\mu$g of tonoplast membrane protein was incubated with 20 mM Tris-Mes, 0.25 M sucrose, 1 mM DTT and acridine orange until a stable baseline was reached (3–5 min). The increase in acridine orange absorbance in membranes isolated from N-deprived (A), $\text{NO}_3^-$-grown (B) and $\text{NO}_3^-$-induced (C) plants was initiated by the addition of different concentrations of KNO$_3$ into the reaction media (indicated by the arrows) and monitored during the following 3 min. Presented values are representative for the results obtained in three to four independent experiments with each experiment done in triplicate. D. The rate of nitrate accumulation in tonoplast vesicles isolated from N-deprived (white bars), $\text{NO}_3^-$-grown (light grey bars) and $\text{NO}_3^-$-induced (dark bars) plants determined by HPLC. The reaction mixture for HPLC assay was deprived of pH-sensitive probe. Values are the means ±SE ($n=5–6$ measurements from 4–6 independent tonoplast preparations). Asterisks indicate a significant difference ($P<0.05$) between proton-coupled nitrate transport activities in tonoplast membranes isolated from different plants.

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![Diagram](image)

**Figure 3. The kinetics of proton-coupled nitrate transport across the tonoplast membranes isolated from cucumbers grown under different nitrate supply.** The effect of nitrate concentration on changes in acridine orange absorbance (A) or nitrate accumulation (B) in ΔpH-energized tonoplast vesicles isolated from N-deprived (circles), NO$_3^-$-grown (squares) and NO$_3^-$-induced (triangles) plants. The Km and R$^2$ values were calculated using GraphPrism Software. The $\frac{1}{1}$Km values are indicated by red arrows. V represents the $\Delta A_{649}$ s$^{-1}$mg$^{-1}$ protein (A) or the $\mu$mol NO$_3^-$ s$^{-1}$mg$^{-1}$ protein (B). doi:10.1371/journal.pone.0073972.g003

suggest that Ca-dependent, staurosporine-sensitive protein kinase could be involved in the regulation of the vacuolar nitrate accumulation processes.

Since sufficient ΔpH across the tonoplast membrane is required for nitrate antiport activity, the effect of soluble cytosolic fractions on V-ATPase-mediated proton transport was also studied to establish whether the two closely linked activities (proton pumping into the vacuole and nitrate/proton exchange) undergo a similar regulation mode. Contrary to nitrate antiport activity in the tonoplast, none of the soluble fractions obtained from plants grown under different nitrate regimes affected proton pumping by V-ATPase (Figure S7).

**CsCLCa, CsCLCc and CsCLCg Expression in Cucumber Roots under Different Nitrate Regimes**

Four of the seven *Arabidopsis* CLCs have been localized to the tonoplast membrane: CLCa, CLCb, CLCc and CLCg. Only three homologs of tonoplast AtCLCs have been found in cucumber and were designated CLCa, CLCc and CLCg based on the homology to *Arabidopsis* proteins (Bogusz and Kłobus, unpublished; Lauge and Kłobus, unpublished). The genes encoding three cucumber CLCs were differentially expressed in roots (Figures 6A and 6B). The CLCa expression was considerably stimulated under constant (over 8-fold) and temporary (over 5-fold) nitrate provision when compared to the level of transcript of the gene under nitrogen deficiency (Figure 6A). Similarly, CLCg mRNA was the most abundant in roots of cucumbers grown under constant nitrate supply, but contrary to CLCa, the expression of CLCg in NO$_3^-$-induced and N-deprived plants was comparable (Figure 6A). In contrast, CLCa transcript significantly increased only upon short-term nitrate provision (Figure 6A).

**Discussion**

It is well evidenced that plants can develop considerably high levels of nitrate within vacuoles using proton-coupled nitrate transporters. Recent years have brought growing information regarding the molecular features and properties of the nitrate antiport system operating at the tonoplast. Active nitrate accumulation within vacuoles allows for storage of large amounts of nitrate inside the cell that can be remobilized during nitrate starvation. Vacular nitrate may also serve as an osmoticum at low light conditions to compensate for the shortage of carbohydrates resulting from decreased photosynthesis [48]. Although nitrate antiporter activity has been demonstrated at the tonoplast and protein candidates for nitrate antiporters have been proposed, the mechanisms of the regulation of active nitrate accumulation within vacuoles, with some exceptions, have not been revealed yet. Using tonoplast membranes isolated from cucumber roots grown in nitrate, we previously demonstrated that tonoplast nitrate antiport utilizes energy from the ATP-generated transmembrane proton gradient [28]. In this study we focus on elucidation of the regulation of the tonoplast H$^+$/NO$_3^-$ antiport activity in response to external nitrogen (NO$_3^-$) nitrate availability.

**H$^+$/NO$_3^-$ Antiport Activity Depends on the Availability of External Nitrate**

The H$^+$-coupled NO$_3^-$ antiport is closely connected to ΔpH generation across the tonoplast membrane catalyzed by V-ATPase. The assays using acridine orange as a ΔpH-sensitive dye showed that both V-ATPase-mediated and NO$_3^-$-induced proton movement across the tonoplast were significantly dependent on the availability of nitrate in the external solution (Figures 1 and 2). Nitrate is a commonly used V-ATPase inhibitor since it uncouples V-ATPase-mediated H$^+$ pumping from ATP hydrolysis. Nevertheless, nitrogen deprivation of plants resulted in the significant decrease of H$^+$-pumping and hydrolytic activity of the tonoplast proton pump (Figures 1A–C). This is not surprising since nitrogen is regarded as a major micronutrient required for the synthesis of crucial metabolites within plant cells, including amino acids, peptides, proteins, nucleic acids, etc. Hence, the biosynthesis of a large multimeric tonoplast proton pump should strictly depend on the nitrogen status of plant cells. Indeed, immunodetection of subunit a of tonoplast V-ATPase in membranes isolated from plants grown under different nitrogen regimes clearly confirmed that the lower V-ATPase activity in N-deprived plants could be a consequence of the decreased biosynthesis of proton pump subunits during N starvation (Figure 1E). However, though the level of V-ATPase protein was also decreased in NO$_3^-$-induced plants, the proton transport activity of this pump was significantly increased, suggesting the change in the coupling of proton transport and ATP hydrolysis of V-ATPase upon short-term nitrate provision. It has previously been reported that the efficiency of coupling may be increased (by Cu ions, transmembrane ΔpH decrease) or decreased (by Cd ions), affecting vacuolar acidification [31], [49], [50]. Hence, short-term (24 h) nitrate provision might have resulted in the change of cytoplasmic and...
vacuolar pH (decrease of ΔpH between cytosolic and vacuolar compartment), leading to the increased coupling efficiency of tonoplast V-ATPase and enhanced vacuolar acidification. Similar, nitrate-dependent changes in coupling efficiency of tonoplast V-ATPase were reported earlier in tobacco plants grown under varied nitrate nutrition [51]. High nitrate concentrations (10 mM, 20 mM) in the nutrition media caused higher rates of ATP hydrolysis and relative rates in proton transport activity when compared to the effect of 2 mM nitrate or ammonium as a nitrogen source [51]. As a result, relative coupling ratios were higher at the higher NO3− concentrations [51].

The different rate of V-ATPase-dependent proton gradient generation could not provide the same level of proton motive force required for tonoplast antiport activities; hence an artificial ΔpH was imposed on the membranes to eliminate the effect of the transmembrane gradient on H+-coupled nitrate transport. Interestingly, nitrate antiport activity was detectable in plants cultivated in all nutritional conditions, though it was considerably lower in N-deprived plants (Figures 2 and S5). Hence, the decreased V-ATPase synthesis in N-deprived plants may result from the decreased activities of the proteins utilizing V-ATPase-catalyzed proton motive force formation. Such co-regulation of the proton pump and antiport activities at the tonoplast was previously demonstrated in mutants unable to synthesize CAX metal/H+ antiporters. The CAXI knock-out mutants not only exhibit a 50% reduction in vacuolar H+/Ca2+ antiport activity but also a 40% decrease in V-ATPase activity [52]. Thus both activities, ATP-dependent proton pumping and H+-coupling across tonoplast membranes, appear to be closely related and influence each other.

Interestingly, similar correlation between proton pumping and nitrate transport activities was found at plasma membranes. As already demonstrated in maize and cucumber, the higher level of nitrate uptake by root [10], [11], [53] and leaf cells [53] was related with a significant increase in the biosynthesis and activity of PM H+-ATPase. Therefore, it has been suggested that nitrate may be the signal triggering the expression of the members of PM H+-ATPase subfamily resulting in the enhanced proton-pumping activity under increased nitrate inflow [11].

The properties of active nitrate transport into the tonoplast were strongly dependent on the nitrogen (NO3−) availability. Despite the lack of nitrogen in nutrient solution, the active nitrate antiport still operated at tonoplast membranes, suggesting that a constitutive, H+-coupled transport system with low capacity and affinity to NO3− ions is present in the vacuolar membrane of plant cells (Figures 2 and S5). The H+/NO3− antiport activity in plants treated with nitrate displayed significantly higher affinity and capacity for NO3− ions, indicating that the existing nitrate antiport system was modified or an additional nitrate antiport system(s) with distinct kinetic properties operates in the vacuolar membranes of plants grown under nitrate availability. Interestingly, the overall nitrate antiport activity at the tonoplast was the highest in NO3−-inducible plants, suggesting that vacuoles were much more capable of active accumulation of nitrate ions after 24 hours of nitrate treatment than after 7 days of nitrate nutrition or

Figure 4. The properties of H+-coupled transport of sulphate and chloride in tonoplast vesicles isolated from the roots of cucumbers grown under different nitrate supply. The time-course of acidine orange absorbance change after the addition of different concentration of K2SO4 (A) or KCl (C) to the reaction media containing ΔpH-energized tonoplast vesicles obtained from NO3−-grown (grey bars), NO3−-induced (dark bars) or N-deprived (white bars) plants. The effect of sulphate (B) or chloride (D) concentration on the acidine orange absorbance ΔpH-energized, tonoplast obtained from NO3−-grown (squares), NO3−-induced (triangles), or N-deprived (circles) plants. Values are the means ±SE (n = 5–6 measurements from 4–6 independent tonoplast preparations). Asterisks indicate a significant difference (P<0.05) between H+-coupled SO42− and Cl− transport in tonoplast isolated from different plants. The Km and R² values were calculated using GraphPrism Software. The −1/Km values are indicated by red arrows. V represents the ΔA495/×min−1 ×mg−1 protein.

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deprivation. Such increased NO$_3^-$ activity might have caused the decrease of ΔpH across the tonoplast membrane followed by the increase in the coupling efficiency of tonoplast V-ATPase (Figures 1A–C). The striking differences in nitrate antiport activity of tonoplast membranes isolated from roots of cucumbers grown under various nitrate availability may result from the morphological and developmental differences between differentially grown plants. Since only plants constantly supplied with nitrate developed the first leaf, the nitrate trafficking and its direction within the tissues of NO$_3^-$-induced and NO$_3^-$-grown plants were probably quite distinct. The leaf developed in NO$_3^-$-grown plants could serve as a sink for nitrate, since leaves play a significant role in nitrogen metabolism. It is known that of the whole absorbed nitrogen, only some is assimilated into amino acids in roots, whereas the majority seems to be translocated to the leaves in the transpiration stream. Consequently, most amino acid synthesis occurs in leaves [54]. This may explain why the tonoplast H$^+$/NO$_3^-$ antiport activity in roots of NO$_3^-$-grown plants was considerably lower from the relevant activity observed in the NO$_3^-$-induced, leaf-deprived plants. The nitrogen (NO$_3^-$)-induced plants were morphologically similar to N-deprived plants, so after the transfer into nitrate-enriched medium following 6-day long N deprivation, they could accumulate higher nitrate levels in root cell vacuoles for storage and further remobilization during NO$_3^-$-induced increase of plant growth and development.

**The H$^+$/NO$_3^-$ Antiport may be Subjected to Phosphorylation in Response to NO$_3^-$ (N) Induction**

In plants, reversible protein phosphorylation seems to be the most common mechanism of post-translational modification of protein activity, ensuring rapid and reversible changes in the ability of plants to respond to specific stimuli. Nitrogen constitutes the major macronutrient limiting plant growth and development; thus plants had to develop mechanisms ensuring the rapid response and adaptation to frequent changes in the external nitrogen status. We have therefore investigated whether the increased nitrate antiport at tonoplast membranes isolated from N-
Figure 6. CsCLCs expression in roots of cucumber grown under different nitrate supply. The level of CsCLCa, CsCLC and CsCLCg transcript was analyzed by Real-time PCR (A) and PCR (B) using reverse transcribed total RNA isolated from plant roots. For the real-time PCR assay, total RNA was prepared from NO₃⁻-grown (grey bars), NO₃⁻-induced (dark bars) and N-deprived (white bars) plants. The cDNA synthesized from RNA isolated from NO₃⁻-grown plants was also used for the PCR assay (B). The gene encoding clathrin adaptor complex subunit (CACS) was used as internal control. Values are the means ± SE (n = 5–6 measurements from 3 independent experiments). Asterisks indicate a significant difference (P < 0.05) between the expression of CsCLCs under different availability of nitrate in nutrient solution (A). Presented picture (B) is representative for the results obtained in three to four independent experiments.

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induced plants may result from the posttranslational modification of antiprotomers occurring in response to 24-hour-long NO₃⁻ induction. Soluble cytosolic fractions were used as the source of protein kinases or protein phosphatases that could be activated or synthesized under different NO₃⁻ availability. The results show that the soluble fraction isolated from NO₃⁻-treated plants affected H⁺/NO₃⁻ antiport activity but not V-ATPase activity in membranes isolated from roots of NO₃⁻-grown plants (Figures 5A and S7), indicating that the stimulation of both tonoplast activities, V-ATPase-mediated proton transport and H⁺-coupled NO₃⁻ antiport, occurs via different regulation modes. Contrary to this, the plasma membrane proton pump was significantly stimulated by the cytosolic fraction isolated from the roots of NaCl-treated plants due to the increased phosphorylation of the membrane protein [55]. It should not be surprising, though, that these two proteins of strikingly different structure, composition and origin are regulated via distinct molecular mechanisms. Since phosphatase inhibitors (okadaic acid or cantharidin) enhanced and protein kinase inhibitors or EGTA completely abolished the stimulatory effect of the soluble fraction on tonoplast H⁺/NO₃⁻ antiport activity through the binding of calcium, and thus decreasing the concentration of free calcium ions in the reaction media.

Summing up, the presented data indicate that the stimulation of H⁺/NO₃⁻ antiport activity at tonoplast membranes of cucumber roots in response to short-term nitrate supply may occur through the phosphorylation of the membrane antiporter involving Ca-dependent, staurosporine-sensitive protein kinase. It remains to be elucidated, however, whether the protein kinase inducing the active nitrate accumulation within the vacuole belongs to the CIPK family of protein kinases activated by calcineurin B-like protein in a calcium-dependent manner. Since nitrate was the only source of nitrogen for plants in our experiments, it also remains to be elucidated whether the observed effect of different nitrate availability on active nitrate transport into the vacuole is nitrate specific, or results from alterations in the nitrogen status of plant cells.

Proteins that could be Responsible for the H⁺/NO₃⁻ Antiport at Tonoplast Membranes

In this work, we present evidence for the presence of distinct, NO₃⁻-responsive antiport systems specific for nitrate in the tonoplast of cucumber root cells. For the last decades much attention has been given to identification of proteins that could mediate nitrate traffic and distribution within plant cells. In comparison to plasma membrane-localized NO₃⁻ transporters, knowledge of the molecular identity and functions of vacuolar nitrate transporters is still very limited and mainly restricted to the chloride channel (CLC) family. Four of seven CLC proteins in A. thaliana were reported in tonoplast membranes: CLC-a and CLC-g [65]. All Arabidopsis CLC genes are ubiquitously expressed throughout the plant with AtCLCa showing the highest expression level [66]. Using electrophysiological and genetic approaches, it was demonstrated that AtCLCa functions as a 2NO₃⁻/1H⁺ exchanger that is able to accumulate nitrate in the vacuole [37]. AtCLCa transcript levels were also shown to be up-regulated by nitrate in roots and shoots [67]. CLC-c might be another contributor of nitrate transport across the tonoplast, since the atclc-c mutant accumulated lower nitrate concentrations when grown at a range of external nitrate levels [68]. Contrary to AtCLCa-a, the expression of AtCLC-c was down-regulated in the presence of nitrate [68]. CLC-b, the closest relative of CLC-a, was also shown to be a tonoplast H⁺-coupled NO₃⁻ transporter; however, the clc-b knockout mutants displayed the same phenotype and accumulated as much nitrate and chloride as the wild type plants [38]. Hence, it seems that only CLC-a is critically involved in the nitrate storage in the vacuole of plant cells. The evidence for posttranslational modification of plant CLC proteins is scarce. It
was only demonstrated that ATP reversibly inhibits AtCLCa by interacting with the C-terminal domain [69]. We have recently identified all but one gene (CLC6) encoding tonoplast-localized AtCLCs homologs in the cucumber genome (Bogusz and Kłobus, unpublished). Only CsCLCe expression was increased upon short-term nitrate provision (Figure 6A), indicating that the enhanced nitrate accumulation in NO$_3^-$-induced plants may result from nitrate-induced antiporter phosphorylation and increased CsCLCe synthesis in response to nitrate supply. Still, the CsCLCs transcript was the most abundant under nitrogen (NO$_3^-$) provision even if its expression was significantly lower when compared to the mRNA level upon constant NO$_3^-$ supply (Figure 6A). Nevertheless, it remains to be established whether CLC6 or CLCe undergoes phosphorylation in response to nitrate or nitrogen to clarify whether these proteins are responsible for the phosphorylation-mediated enhanced nitrate accumulation in root cell vacuoles in response to short-term nitrogen (NO$_3^-$) provision.

Phosphorylation-coupled nitrate transport has been well evidenced for the NRT1.1 transporter. Since the protein kinase C recognition motifs have also been found in the N- and C-terminal domains of the HvNRT2.1 transporter [20], the phosphorylation events may be common for both NRT1 and NRT2 transporters. Until now, among all characterized NRT proteins only few has been shown to be localized to vacuolar membrane: AtNR2.7 and PTR2-like transporters from A. thaliana (PTR4 and PTR6), barley (Hv PTR2) and Hak← actes (HvPTR4) [39], [70–74]. Nevertheless, only AtNR2.7 is able to transport nitrate; this NRT2-like transporter is responsible for nitrate accumulation in the vacuoles of reproductive organs and seeds of A. thaliana [39]. All the functionally characterized NRT1-like transporters with confirmed ability to transport nitrate localize to the plasma membrane [reviewed by [18]]. The screening of the whole-genome shotgun reads (GenBank) revealed lack of the AtNR2.7 homolog in cucumber genome (Lange and Kłobus, unpublished); hence we assume that either CLC protein or the NRT1-like transporter (a low affinity transporter) or both systems are responsible for the H$^+/NO_3^-$ antiport at the tonoplast in cucumber root cells. We have recently identified 13 homologs of AtNRT1 genes in cucumber that are differentially regulated by nitrate provision [75]. However, the subcellular localization and regulation of the CsNRT1 as well as the CsCLCs proteins remain to be established.

Supporting Information

Figure S1 Effect of MgATP (A), V-ATPase inhibitors: KN03 or bafilomycin and gramicidin or NH$_4$Cl (B) on the proton flux assay in tonoplast vesicles. A. To induce V-ATPase-mediated proton influx into tonoplast vesicles, 3 mM MgATP (+MgATP) or 3 mM MgSO$_4$ (-MgATP) were added to the reaction media. The proton gradient generation occurred only in the presence of ATP and was immediately inhibited by bafilomycin, a specific inhibitor of tonoplast proton pump V-ATPase. Protonophore gramicidin (5 μM) or NH$_4$Cl (5 mM) caused an increased proton flux in acridine orange absorbance at 495 nm indicating that the observed changes in the absorbance of the probe result from changes in proton gradient generation (by MgATP) and recovery (by gramicidin or NH$_4$Cl). B. Bafilomycin (500 nmol) and KN03 (50 mM) prevent V-ATPase-mediated proton translocation when added to the reaction along with MgATP, MgATP or MgSO$_4$ were added to the reactions containing vacuolar membrane vesicles (50 μg), acridine orange (10 μM), 20 mM TRIS-MES (pH 7.2), 0.25 M sucrose, 50 mM KCl, and 1 mM DTT that had been preincubated for 5 min at room temperature. The quenching of acridine orange absorbance at 495 nm was monitored as described in Materials and methods.

Figure S2 Mechanism for the use of ΔpH sensing acridine orange to measure MgATP-dependent proton fluxes in isolated tonoplast vesicles. Acridine orange is a weak hydrophobic permeant probe containing a basic group as a weak base [76], [77]. In the unprotonated form, acridine orange has the capacity for free movement across the membrane while protonation of the basic amine group prevents free transmembrane movement of the probe [77]. During 5 min-long incubation of tonoplast vesicles with acridine orange (pH 7.5), the unprotonated probe (green circles) freely moves across the membranes until the balance between the interior and exterior of the vesicles is achieved. The addition of MgATP into tonoplast vesicles initiates the V-ATPase-mediated proton influx into tonoplast vesicles and acidification of vesicle lumen [77]. Under these conditions the protonation of the probe occurs (red circles) which renders acridine orange positively charged. As a result, the protonated form of the probe accumulates in the interior of the tonoplast membranes as it cannot freely move outside the lumen [77]. The pH-dependent accumulation of acridine orange within tonoplast membranes results in a change in the absorption spectrum and is detectable as the decrease of probe absorbance at 495 nm [76], [77]. Following the inhibition of V-ATPase (by bafilomycin) and the addition of NO$_3^-$ ions into the reaction media, NO$_3^-$-mediated proton efflux from the tonoplast membranes results in the decrease of the proton gradient and unprotonation of the acridine orange accumulated within vesicles. As a result, the corresponding edlux of the probe from the membranes into external solution occurs coupled to the increase of the probe absorbance at 495 nm [77]. OA, acridine orange.

Figure S3 Mechanism for the use of ΔpH sensing acridine orange to measure ΔpH-dependent proton fluxes in isolated tonoplast vesicles. Tonoplast vesicles suspended in pH 6.0 (to impose transmembrane ΔpH) or 8.0 (to maintain equal pH in the interior and exterior of the vesicles) were introduced into the reaction media of pH 8.0 and incubated with acridine orange for 5 min. Protonated probe (red circles) accumulated within the acidic interior of the tonoplast membranes. Following the addition of NO$_3^-$ ions to the reaction, the NO$_3^-$-induced proton efflux from the tonoplast membranes resulted in unprotonation of the acridine orange (green circles), change in the absorption spectra and corresponding efflux of the probe outside tonoplast lumen detectable as the increase in the absorbance of the probe at 495 nm (Figures 2A–C). In vesicles without transmembrane ΔpH, the addition of NO$_3^-$ ions did not induce proton efflux from the vesicle lumen due to the lack of proton motive force. The lack of vesicle lumen acidification rendered the acridine orange unprotonated and prevented accumulation of the probe inside vesicles. Hence, the absorbance of the probe at 495 nm did not change (Figure S6 D–F). Similar assays were performed with quinacrine using fluorescence change as ΔpH indicator (Figures S5 and S6 A–C). OA, acridine orange.

Figure S4 The quality and integrity of RNA isolated from cucumber roots. Total RNA was extracted from the roots of cucumber plants grown in constant (NO$_3^-$-grown) and temporary 24-hour-long (NO$_3^-$-induced) nitrate supply or from plants cultivated without nitrogen source (N-deprived). Following 5 min-long denaturation at 65°C and 2 min-long cooling on ice,
RNA samples were subjected to electrophoresis using 2100 Bioanalyzer (Agilent) and RNA kits provided by the manufacturer. L = RNA ladder (DOC)

Figure S5 Characteristics of NO₃⁻–dependent increase in quinacrine fluorescence. A pH gradient (inside-acid) was imposed in tonoplast vesicles isolated from roots of N-deprived (A), NO₃⁻–-grown (B) and NO₃⁻––induced (C) cucumbers. The increase of quinacrine fluorescence was initiated by the addition of KNO₃ into the reaction media (indicated by the black arrows) and monitored during the following 3 min. Presented values are representative for the results obtained in three to four independent experiments with each experiment done in triplicate. D. The effect of nitrate concentration on the changes in quinacrine fluorescence in ΔpH=energized tonoplast vesicles isolated from N-deprived (circles), NO₃⁻–-grown (squares) and NO₃⁻––induced (triangles) plants. The Km and Vᵢ values were calculated using GraphPrism Software. The -1/Km values are indicated by red arrows. V represents the ΔF/min×1⁻¹×mg⁻¹ protein. (DOC)

Figure S6 The effect of NO₃⁻ on the changes in quinacrine fluorescence (A–C) and acridine orange absorbance (D–F) in tonoplast membranes without transmembrane ΔpH. Tonoplast vesicles were isolated from roots of N-deprived (A, D), NO₃⁻––grown (B, E) and NO₃⁻––induced (C, F) cucumbers. Following 5-min-long incubation of vesicles with pH-sensitive probes, the different concentrations of KNO₃ were added into the reaction media (indicated by the arrows) and the acridine orange absorbance or quinacrine fluorescence were monitored during the following 3 min. Present- ed values are representative for the results obtained in three to four independent experiments with each experiment done in triplicate. (DOC)

Author Contributions
Conceived and designed the experiments: MM GK. Performed the experiments: MM AW AP. Analyzed the data: MM. Contributed reagents/materials/analysis tools: MM AW AP. Wrote the paper: MM.
33. Maeshima M (2000) Vacuolar H+/pyrophosphatase. Biochim Biophys Acta 1465: 37–51.
34. Nakanihi Y, Maeshima M (1998) Molecular cloning of vacuolar H+/pyrophosphatase and its developmental expression in growing hypocotyl of mung bean. Plant Physiol 116: 589–597.
35. Kabala K, Klobus G (2001). Characterization of the tonoplast proton pumps in *Cuscuta sativa* L. root cells. Acta Physiol Plant 23: 53–63.
36. Krebs M, Beyhl D, Gortlich E, Al-Rashidi KA, Marten I et al. (2010) *Arabidopsis* V-ATPase activity at the tonoplast is required for efficient nutrient storage but not for sodium accumulation. Proc Natl Acad Sci, U S A 107(17): 3251–6.
37. De Angelis A, Monachello D, Ephritikhine G, Frachisse JM, Thomine S et al. (2006) The nitrate/proton antiporter AtCLC:mediates nitrate accumulation in plant vacuoles. Nature 442: 939–942.
38. Von der Fecht-Bartenbach J, Boguer M, Dynowski M, Ladewig U (2010) CLC+/mediated NO3−/H+ exchange across the tonoplast of *Arabidopsis* vacuoles. Plant Cell Physiol 51(6): 960–968.
39. Chopin F, Orsel M, Dorbe M, Chardon F, Truong H et al. (2007) *The Arabidopsis AtNRT2.7 nitrate transporter controls nitrate content in seeds*. Plant Cell 19: 1590–1602.
40. Migocka M, Papierniak A, Koszata E, Klobus G (2011) Comparative study of the active cationic efflux systems operating at the plasma membrane and tonoplast of cucumber root cell. J Exp Bot 62(14): 4905–4916.
41. Gallagher SR, Leonard RT (1982) Effect of vanadate, molybdate and azide on membrane-associated ATPase and soluble phosphatase activities of corn roots. Plant Physiol 70: 1335–40.
42. Anes BN (1966) Assay of inorganic phosphate, total phosphate and phosphatases. Method Enzymol 8: 115–8.
43. Thayer JR, Huffaker RC (1980) Determination of nitrate and nitrite by high-pressure liquid chromatography: Comparison with other methods for nitrate determination. Anal Biochem 102(1): 110–119.
44. Bradford MM (1976) A rapid and sensitive method for quantitation of microgram quantities of protein utilizing the principles of protein dye binding. Anal Biochem 72: 248–54.
45. Huang S, Li R, Zhang Z, Li L, Gao X et al. (2009) The genome of the cucumber, *Cuscuta sativa* L. Nat Genet 41: 1273–1281.
46. Salamov AA, Solovyev VV (2000) Ab initio gene finding in *Cucumis sativus* L. Nat Genet 41: 1275–1281.
47. Lomsadze A, Ter-Hovhannisyan V, Chernoff YO, Borodovsky M (2005) *Gene identification in novel eukaryotic genomes by self-training algorithm*. Nucl DNA. Genome Res 10: 516–522.
48. Blom-Zandstra M, Lampe JEM (1985) *The role of nitrate in the osmoregulation of Crassulacean Acids Res 33: 6494–6506*.
49. Huo S, Li Y, Song Y, Wang W, Li Y et al. (2007) *The calcium sensor CBL19 mediates salt tolerance by regulating ion homeostasis in Arabidopsis*. Plant J 52(3): 473–84.
50. Xu J, Li HD, Chen LQ, Wang Y, Liu L et al. (2006) A protein kinase, interacting with two calcium-binding-like proteins, regulates K+ transporter AKT1 in *Arabidopsis*. Cell 125: 1347–1360.
51. Kim BG, Waadt R, Cheong YH, Pandey GK, Dominguez-Solís JR et al. (2007) *The calcium sensor CBL19 mediates salt tolerance by regulating ion homeostasis in Arabidopsis*. Plant J 52(3): 473–84.
52. Cheng NH, Pittman JK, Barkla BJ, Shigaki T, Hirschi KD (2003) *The CBL4/CIPK6 calcium sensor/protein kinase complex*. Cell 121(1): 84–92.
53. Andrews M (1986) *The partitioning of nitrate assimilation between root and shoot of higher plants*. Plant Cell Environ 9: 511–519.
54. Knobus G, Janicka-Russak M (2008) *Modulation by cytosolic components of proton pump activities in Cuscuta sativa* roots during salt stress. Plant Physiol 121(1): 84–92.
55. Tsay FY, Schreoder JL, Feldmann KA, Crawford NM (1993) The herbicide sensitivity gene CHL1 of *Arabidopsis* encodes a nitrate-inducible nitrate transporter. Cell 72: 705–713.
56. Krouk G, Crawford NM, Coruzzi GM, Tsay FY (2010) *Nitrate signaling: adaptation to fluctuating environments*.Curr Opin Plant Biol 13(3): 263–272.
57. Held K, Pascaud F, Eckert C, Gajdusanovcic P, Hashimoto K et al. (2006) *Calcium-dependent modulation and plasma membrane targeting of the AKT2 potassium channel by the CBLA/CIPK6 calcium sensor/protein kinase complex*. Cell Res 21(7): 1116–1130.
58. Li L, Kim BG, Cheong YH, Pandey GK, Luyn S (2006) *A Ca2+ signaling pathway regulates a K+ channel for low-K response in Arabidopsis*. Proc Natl Acad Sci, USA 103: 12625–12630.
59. Xu J, Li HD, Chen LQ, Wang Y, Liu L et al. (2006) A protein kinase, interacting with two calcium-binding-like proteins, regulates K+ transporter AKT1 in *Arabidopsis*. Cell 125: 1347–1360.
60. Li L, Zhang J, Wei J, Wang H, Wang Y et al. (2009) *Functions and mechanisms of the CBL-CIPK signaling system in plant response to abiotic stress*. Prog in Nat Sci 19: 667–676.
61. Zhou JJ, Wei JJ, Wang C, Wu J, Ratnamurtha D et al. (2010) *Arabidopsis Calcium-Dependent Protein Kinase CIPK10 functions in abscisic acid- and Ca2+-mediated stomatal regulation in response to drought stress*. Plant Physiol 154: 1232–1243.
62. Lo Q, Tang R, Liu H, Gao X, Li Y et al. (2009) *Cloning and molecular analyses of the Arabidopsis thaliana chloride channel gene family*. Plant Sci 176: 650–661.
63. De Angelis A, Monachello D, Ephritikhine G, Frachisse JM, Thomine S et al. (2009) *CLC-mediated anion transport in plant cells*. Philos Trans R Soc Lond B Biol Sci 364(1514): 195–201.
64. Greener D, Larin C, Bouchez D, Frachisse JM, Lealveve F et al. (2009) *Disruption of putative anion channel gene AtCLC-a in Arabidopsis suggests a role in the regulation of nitrate content*. Plant J 21: 259–267.
65. Harada H, Kuronovri T, Hayama T, Shinozaki K, Leigh RA (2004) *Quantitative trait loci analysis of nitrate storage in Arabidopsis leading to an investigation of the contribution of the anion channel gene, AtCLC-a, to variation in nitrate levels*. J Exp Bot 55(405): 2005–14.
66. De Angelis A, Moran O, Wege S, Filleur S, Ephritikhine G et al. (2009) *ATP binding to the C-terminus of the Arabidopsis thaliana nitrate/proton antiporter, AtCLC-a, regulates nitrate transport into plant vacuoles*. J Biol Chem 284(19): 26526–26.
67. Paungfoo-Lonhienne C, Lonhienne G, Teng TS, Robinson N, Christie M et al. (2006) *Plants can use protein as a nitrogen source without assistance from other organisms*. Proc Natl Acad Sci, U S A 103(11): 4524–9.
68. Weichert A, Brinkmann C, Komarova NV, Dietrich D, Thor K et al. (2012) *AtPTr4 and AtPTr6 are differentially expressed, tonoplast-localized members of the peptide transporter/nitrate transporter 1 (PTR/NRT1) family*. Planta 225(2): 311–23.
69. Carter C, Pan S, Zouhar J, Avila EL, Girké T et al. (2004) *The vegetative vacuolar proteome of Arabidopsis thaliana reveals predicted and unexpected proteins*. Plant Cell 16(12): 3285–303.
70. Dunkley TP, Hester S, Sadhoff FP, Kiyomizu H, Weimar T et al. (2006) *Mapping the Arabidopsis organelle proteome*. Proc Natl Acad Sci, U S A 103(17): 5186–9.
71. Endler A, Meyer S, Schellbirt S, Schneider T, Weschke W et al. (2006) *Identification of a vacuolar sucrose transporter in barley and Arabidopsis mesophyll cells* by a tonoplast proteomic Approach. Plant Physiol 141(1): 196–207.
72. Migocka M, Warzybok A, Klobus G (2012) *The genomic organization and transcriptional pattern of genes encoding nitrate transporters 1 (NRT1) in cucumber*. Plant Soil 364: 245–260.
73. Rasi-Caldogno F, Pugliarello MC, De Michelis MI (1985) *Electrogenic transport of nitrate in barley and Arabidopsis*. Cell Res 21(7): 1116–1130.
74. Rasi-Caldogno F, Pugliarello MC, De Michelis MI (1985) Electrogenic transport of nitrate in barley and Arabidopsis. Cell Res 21(7): 1116–1130.
75. Migocka M, Warzybok A, Klobus G (2012) *The genomic organization and transcriptional pattern of genes encoding nitrate transporters 1 (NRT1) in cucumber*. Plant Soil 364: 245–260.