Cloning of *chrysanthemum* high-affinity nitrate transporter family (*CmNRT2*) and characterization of *CmNRT2.1*

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The family of NITRATE TRANSPORTER 2 (NRT2) proteins belongs to the high affinity transport system (HATS) proteins which act at low nitrate concentrations. The relevant gene content of the chrysanthemum genome was explored here by isolating the full length sequences of six distinct *CmNRT2* genes. One of these (*CmNRT2.1*) was investigated at the functional level. Its transcription level was inducible by low concentrations of both nitrate and ammonium. A yeast two hybrid assay showed that *CmNRT2.1* interacts with *CmNAR2*, while a BiFC assay demonstrated that the interaction occurs at the plasma membrane. *Arabidopsis thaliana* plants heterologously expressing *CmNRT2.1* displayed an enhanced rate of labeled nitrogen uptake, suggesting that *CmNRT2.1* represents a high affinity root nitrate transporter.

Plants have evolved both a high (HATS) and a low (LATS) affinity transport system, which act together to determine nitrate uptake from the soil. The genes involved in HATS and LATS belong to three distinct gene families, namely the nitrate transporter/peptide transporters (*NPFs*), the nitrate transporters (*NRT2s*) and the nitrate assimilation related genes (*NAR2s*). The NRT2 family is part of the major facilitator superfamily (MFS). The *Arabidopsis thaliana* (*A. thaliana*) genome harbors the seven *NRT2* genes, referred to as *AtNRT2.1* through *2.7*. NRT2 homologs are also known in other plant species: five have been identified in barley, four in rice and six in the green alga *Chlamydomonas reinhardtii*. NRT2 was initially identified as a putative high affinity transport gene in barley and tobacco before their presence was revealed in other species including *A. thaliana*. On the basis of their transcriptional response to the provision of nitrate, the *AtNRT2*s have been classified into three types: those which are inducible (*AtNRT2.1, 2.2 and 2.4*), those which are constitutively transcribed (*2.3, 2.6 and 2.7*), and the repressible gene *2.5*. *AtNRT2.1* encodes a high affinity nitrate transporter which functions at low external nitrate concentrations, acting as either a nitrate sensor or a signal transducer in the root. Although the *AtNRT2.1* product is an important contributor to the inducible component of high affinity nitrate transport, *AtNRT2.2* acts as an alternative when *AtNRT2.1* is disrupted.

The evidence which implied a functional interaction existed between NRT and NAR proteins was initially acquired in *Chlamydomonas reinhardtii*, but it has also been shown that both the *A. thaliana* mutants *atnrt2.1* and *atnar2.1* lack inducible HATS activity at low levels of nitrate. Direct evidence for the *AtNRT2.1/AtNAR2.1* interaction has also been generated via yeast two hybrid, *Xenopus* oocyte and blue native polyacrylamide gel electrophoresis technologies. The loss-of-function of both the rice gene *OsNAR2.1* and *AtNAR2.1* leads to a drastic reduction in plant growth.

Chrysanthemum (*Chrysanthemum morifolium*) is a leading ornamental species, particularly in China, where the heavy application of nitrogenous fertilizer is a common practice pursued by growers, despite its known deleterious effect on the environment. So far, only one *CmNRT2* (AB921547 renamed *CmNRT2.4* in this study) gene has been isolated; its product interacts with *CmNAR2*. Here, a description is given of the isolation of a further six *CmNRT2s* and a more detailed analysis of the function of *CmNRT2.1* has been given. In particular, its inducibility by both nitrate and ammonium has been characterized, and its interaction with *CmNAR2* further detailed.

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Finally, a demonstration is given that the heterologous expression of CmNRT2.1 in A. thaliana resulted in an increased rate of nitrate uptake into the root.

**Results**

**The CmNRT2s and their predicted products.** A peptide level sequence comparison between the set of AtNRT2s and the single previously known CmNRT2 (CmNRT2.4, AB921547) revealed the presence of two conserved regions, namely A1 [FGMRGRILW(N/T/A/I/G)(L/W)W] and A2 [(H/Y)FPFWG(S/G)(M/F/C)] (Fig. 1A,B). Based on these motifs, the two degenerate primers A1f and A2-2f (Table 1) were designed to amplify the 3′ end of the CmNRT2s using 3′ RACE. The outcome of the procedure was the successful identification of seven distinct 3′ untranslated regions (3′ UTRs), including that of CmNRT2.4 (Fig. 1C), which implied that the chrysanthemum genome housed seven CmNRT2 genes. The full length cDNAs of CmNRT2.1 through 2.3 and 2.5 through 2.7 were subsequently obtained by means of 5′ RACE-PCR, following Gu et al.19. The resulting predicted polypeptide sequences were then aligned (Fig. 1D, Table 2). The greatest extent of sequence identity obtained between CmNRT2.1/2.2 and 2.2/2.4 (94.3%), followed by the 2.1/2.4 pair (93.0%), but only a low level of identity (41.5%) existed between 2.1 and 2.7. The DNA sequences were used to determine the genes’ intron-exon structure and a schematic representation of the inferred structures is given as Fig. S1. Five of the genes (2.1, 2.2, 2.3, 2.4 and 2.6) share two conserved introns, whereas 2.5 harbors two introns sited in a different part of the gene, and 2.7 lacks any introns. A phylogenetic tree based on peptide sequences is shown in Fig. 2; this indicated that five of the gene products (CmNRT2.1, 2.2, 2.3, 2.4 and 2.6) cluster within a single clade with a highly supportive bootstrap value, while 2.5 and 2.7 are outliers. CmNRT2.4 has previously been shown to be a strong candidate as a nitrate uptake protein19, so the strong level of peptide similarity and gene structure between it and CmNRT2.1 was suggestive of the latter protein sharing a similar functionality.

**CmNRT2.1 is inducible by nitrate and ammonium.** The topological profiling of CmNRT2.1 transcription showed that the gene was constitutively transcribed throughout the plant except in the flower, with the root being the site where its transcript was most abundant (Fig. S2). When the plants were provided with various concentrations of either nitrate or ammonium, the abundance of CmNRT2.1 transcript in the root was enhanced. The nitrate response peaked after 4h irrespective of the nitrate concentration (Fig. 3A–C), and similarly, the provision of ammonium provoked a transcriptional peak after 4h (Fig. 3D).

**In vivo interaction between CmNRT2.1 and CmNAR2.** A BiFC analysis was conducted to characterize the interaction between CmNRT2.1 and CmNAR2, based on the transient expression of split YFP-labelled CmNRT2.1 and CmNAR2 in onion epidermal cells. The CmNRT2.1 and CmNAR2 fusion proteins were engineered to have complementary N terminal and C terminal YFP fragments. The epidermal cells expressing CmNAR2–cEYFP and CmNRT2.1–nEYFP showed strong YFP complementation. In contrast, cells transformed with either CmNRT2.1–nYFP/cEYFP, nYFP/CmNAR2.1–cYFP or nYFP/cYFP emitted no fluorescence (Fig. 4). The interpretation of these observations was that CmNRT2.1 can interact with CmNAR2 in vivo. As a follow-up experiment, a split-ubiquitin membrane two hybrid system experiment was performed. This confirmed that CmNRT2.1 and CmNAR2 interact, as shown by the ability of the yeast cells to grow on the quadruple dropout medium, while cells carrying either pPR3-N/pBT3-C-NAR2.1 or pPR3-N/pBT3-C-NRT2.1 were unable to grow (Fig. 5A). The three combinations used as a positive control all promoted growth on the double-dropout medium (Fig. 5B). When the colonies were re-streaked onto a plate containing X-Gal, only cells harboring both CmNRT2.1 and CmNAR2 were able to produce a signal (Fig. 5C).

**Transgenic A. thaliana lines expressing CmNRT2.1 showed an enhanced level of nitrate uptake into the root.** A. thaliana transgenic plants were generated which constitutively expressed CmNRT2.1. An analysis of their genomic DNA (Fig. 6A) and mRNA (Fig. 6B) demonstrated that the transgene was successfully incorporated and transcribed. Two independent transgene homozygous T3 selections (RT-2 and -19) were used to study the transgenic’s impact on plant growth and nitrate uptake. Both root and shoot fresh weight (FW) of the two lines were significantly higher than those of the wild type (WT) control and vector-transformed lines when the growing medium contained 0.25 mM nitrate. With respect to root FW, there was no significant difference in performance between the transgenics and the controls when the medium contained 10 mM nitrate, but shoot FW was enhanced in RT-2 (Fig. S3). When nitrate uptake was assessed using labeled nitrate, both transgenics out-performed both WT and the empty vector control (Fig. 6C), consistent with the suggestion that CmNRT2.1 provides an improved capacity to take up nitrate.

**Discussion**

NRT2s have been isolated to date largely using degenerate primers; presently, seven members are known in A. thaliana12, six in poplar20 and four in Lotus japonicus21. Here, the same approach has been used to identify and isolate six further CmNRT2s to add to the one isolated previously21. At the peptide level, AtNRT2.1, 2.2 and 2.4 share homology with one another, as do AtNRT2.3 and 2.6. Similarly, a group of four CmNRT2s (2.1, 2.2, 2.3 and 2.4) share closely related sequences (which aligned well with that of AtNRT2.6) (Table 2). It has been shown previously that CmNRT2.4 interacts with CmNAR2 to promote nitrate uptake18.

The extensive homology between CmNRT2.1 and 2.4, as well as the nitrate inducibility of CmNRT2.1 (Table 2) was suggestive of CmNRT2.1 being functionally similar to CmNRT2.4. The level of sequence homology between AtNRT2.1 and CmNRT2.1 was high (>76%). AtNRT2.1 is inducible by nitrate at a concentration of both 0.5 mM and 10 mM3, while the present experiments demonstrated that CmNRT2.1 was inducible by all three nitrate concentrations provided to the plants. CmNRT2.1 transcription showed to be induced by the 5 mM ammonium concentrations within 4h, thereafter was suppressed by 5 mM ammonium. OsNRT2.1 transcription is similarly suppressed by the presence in the medium of 5 mM ammonium22. CmNRT2.1 transcript was particularly...
Figure 1. Isolation and peptide-based alignment of CmNRT2s. (A,B) The two conserved regions identified by aligning the derived amino acid sequences from AtNRT2s and CmNRT2-4; (C) an alignment of the CmNRT2 3′ UTRs; (D) a peptide-based alignment of cloned CmNRT2s.
abundant in the root (Fig. S2), a feature which is also characteristic of AtNRT2.1. The rice NRT2 homologs OsNRT2.1, 2.2 and 2.3a are all induced by the presence of low concentrations of nitrate. In contrast to the transcriptional behavior of CmNRT2.4 and the A. thaliana genes AtNRT2.3, 2.6 and 2.7, CmNRT2.1 transcript was not detectable when the plants were deprived of either nitrate or ammonium (Fig. 3). Moreover, the increased folds of highest transcription level of CmNRT2.1 compared to the control is larger than that in CmNRT2.4 under 4 h nitrate exposure.

The yeast two hybrid and BiFC analyses confirmed that CmNRT2.1 and CmNAR2 interacted with one another in vivo, as do a number of the A. thaliana NRT2s with AtNAR2.1, rice NRT2s with OsNAR2.1 and barley NRT2s with HvNAR2.3. Further experiments will be needed to establish whether any of the CmNRT2s other than CmNRT2.1 and 2.4 are able to likewise interact with CmNAR2. The AtNRT2.1/NAR2.1 interaction has been shown to take place in the plasma membrane, forming a 150 kDa complex, thought to act as a high affinity nitrate transporter. The present data were consistent with the CmNRT2.1/NAR2 complex similarly localizing to the plasma membrane (Fig. 4). Transgenic A. thaliana plants constitutively expressing CmNRT2.4 display an enhanced rate of nitrate uptake compared to the WT and the empty vector controls, and the present experiments have shown that CmNRT2.1 activity can also contribute to nitrate uptake (Fig. 6), as expected given the high degree of sequence similarity (>93%) existing between CmNRT2.1 and 2.4 (Table 2). The conclusion is that CmNRT2.1 is a nitrate inducible gene, the product of which is a high affinity nitrate transporter. As such, it represents a suitable candidate for the engineering of nitrate uptake efficiency in chrysanthemum.

**Methods**

**Plant materials and growth conditions.** The experiments were based on the chrysanthemum cultivar ‘Nannongxuefeng’, maintained at the Nanjing Agricultural University Chrysanthemum Germplasm Resource Preserving Centre (Nanjing, China). Phenotypically uniform seedlings at the eight leaf stage were grown in a pH 6.5 medium containing 5 mM NH₄NO₃, 2.5 mM K₂SO₄, 1.5 mM MgSO₄·7H₂O, 1.33 mM NaH₂PO₄·2H₂O, 2.0 mM CaCl₂, 20 μM H₃BO₃, 9 μM MnCl₂·4H₂O, 0.77 μM ZnSO₄·7H₂O, 0.32 μM CuSO₄·5H₂O, 0.39 μM Na₂MoO₄·2H₂O and 20 μM FeNaEDTA, following Gu et al. Nitrification was inhibited by the inclusion of 7 μM dicyandiamide. The solution was refreshed every two days.
Figure 2. An unrooted phylogenetic tree of the CmNRT2 peptide sequences of chrysanthemum and the barley homologs HvNRT2.1 (U34198), HvNRT2.2 (U34290), HvNRT2.3 (AF091115), HvNRT2.4 (AF091116) and HvNRT2.5 (ABG20828), the rice homologs OsNRT2.1 (P0DKG9), OsNRT2.2 (P0DKH0), OsNRT2.3 (Q94JG1.1) and OsNRT2.4 (A2ZU80.2), the A. thaliana homologs AtNRT2.1 (AAC64170), AtNRT2.2 (AAC35884), AtNRT2.3 (BAB10099), AtNRT2.4 (BAB10098), AtNRT2.5 (AAF78499), AtNRT2.6 (CAB89321) and AtNRT2.7 (CAB87624) and the chrysanthemum homologs CmNRT2.1 (KT203959), CmNRT2.2 (KT203960), CmNRT2.3 (KT203961), CmNRT2.4 (DDBJ accession AB921547), CmNRT2.5 (KT203962), CmNRT2.6 (KT203963) and CmNRT2.7 (KT203964). The sequences were aligned using ClustalW software and the phylogeny constructed using the neighbor-joining method. Five CmNRT2 sequences (2.1, 2.2, 2.3, 2.4, and 2.6) clustered into a single clade supported by a bootstrap value of 100%.

Figure 3. The induction of CmNRT2.1 transcription in N starved roots exposed to (A) 0.5 mM nitrate, (B) 1 mM nitrate, (C) 5 mM nitrate, (D) 5 mM ammonium, as assayed by qRT-PCR. Error bars in (A–D) represent the SE (n = 3).
Isolation and sequencing of CmNRT2 full-length cDNAs. For the gene isolation experiment, seedlings were grown in the solution (as described in the part of plant materials and growing conditions) for four weeks, then starved of nitrogen by removing them to a nitrogen-free version of the same medium for one week. After exposing the seedlings to 5 mM KNO₃ for 4 h, RNA was extracted from the roots using the RNAiso reagent (TaKaRa, Tokyo, Japan), following the manufacturer’s protocol, then treated with RNase-free DNase I (TaKaRa, Tokyo, Japan). The concentration and the integrity of the extract were assessed following Gu et al. The first cDNA strand was synthesized using Reverse Transcriptase M-MLV (RNase H⁻) (TaKaRa, Tokyo, Japan), following the manufacturer’s protocol. Two degenerate primers A1f and A2-2f (sequences given in Table 1) were designed based on regions conserved between the CmNRT2.4 and the AtNRT2 sequences. The remainder of the cDNA sequence was acquired using RACE-PCR, following Liu et al. Open reading frames (ORFs) were identified from the resulting sequences using ORF finder software (www.ncbi.nlm.nih.gov). The genes’ deduced polypeptide sequences were used in a BLASTp search to identify homologs. An alignment of the seven derived CmNRT2s was performed using DNAman v 5.2.2 software (Lynnon Bio-Soft, Quebec, Canada). A phylogenetic analysis of the CmNRT2s and heterologous NRT2s was finally performed using MEGA 5 software.

Quantitative real-time PCR (qRT-PCR). Seedlings were deprived of nitrogen for one week after having been grown in the solution (as described in the part of plant materials and growing conditions) for four weeks. After providing a variable amount of nitrate (0.5, 1.0 or 5.0 mM KNO₃) or 5 mM NH₄Cl, the roots were harvested.
were used to detect the expression level of reference gene of CmPsaA-F (\textit{ura3::lacZ} \textit{ade2::ADE2} \textit{GAL4} CmNAR2 2.1-JX (Table 1), and the amplicon introduced into the pPR3-N (\textit{TRP1, AmpR}) plasmid. The medium at 30 °C overnight. When the OD546 of the cultures reached 1.0, the cultures were serially diluted (10£\textsuperscript{x}) to calculate relative changes in transcript abundance\textsuperscript{28}. Each derived relative transcript abundance was based on the mean of three biological replicates.

Protein-protein interaction assays. The existence of a CmNRT2.1/CmNAR2 interaction was tested using the mating-based split-ubiquitin system (Dualsystems Biotech, Schlieren, Switzerland) and a BiFC assay. The necessary fusion gene pSAT4A-CmNAR2-cEYFP-N1 has been described previously\textsuperscript{18}. For the BiFC analysis, the CmNRT2.1 ORF was amplified using the primer pair CmNRT2.1-Bi-S/CmNRT2.1-Bi-X (Table 1) and the amplicons of CmNRT2.1 and CmNAR2 were used to create the construct pSAT4A-CmNRT2.1-nEYFP-N1, pSAT4A-CmNAR2-cEYFP-N1 and pSAT4A-CmNRT2.1-nEYFP-N1. Two constructs were mixed with 1:1 gold particles (Bio-Rad, Hercules, CA) and transformed as described by Gu \textit{et al.}\textsuperscript{18}. Confocal laser microscopy was used to monitor the expression of YFP. For the yeast two hybrid analysis, full length \textit{CmNRT2.1} cDNA was first amplified using Phusion\textsuperscript{®} HS DNA polymerase (NEB, Ipswich, MA, US) based on the primer pair CmNRT2.1-1301-F/CmNRT2.1-1301-R (Table 1), digested with \textit{Bam}HI and \textit{Sac}I sites to produce the construct pCAMBIA1301-220 \textit{Bam}HI and \textit{SacI} sites to produce the construct p1301-220-CmNRT2.1. Either an empty pCAMBIA1301-220 vector or p1301-220-CmNRT2.1 was transformed into \textit{Agrobacterium tumefaciens} strain EHA105 using the freeze-thaw method\textsuperscript{29}. \textit{A. thaliana} Col-0 was transformed using the floral dip method\textsuperscript{18}. T\textsubscript{1} seedlings were raised on Murashige and Skoog (1962) (MS) medium containing 20 mg/L hygromycin and 25 mg/L ampicillin. Positive transformants were validated by observing GUS expression in the leaf\textsuperscript{29} and also by deploying a PCR assay on their genomic DNA, based on the primer pair CmNRT2.1-F/CmNRT2.1-R (Table 1).

Transgene construction and \textit{A. thaliana} transformation. The \textit{CmNRT2.1} ORF was amplified using the primer pair CmNRT2.1-1301-1-F/CmNRT2.1-1301-1-R (Table 1), digested with \textit{Bam}HI and \textit{SacI} and inserted into the pCAMBIA1301-220 \textit{Bam}HI and \textit{SacI} sites to produce the construct p1301-220-CmNRT2.1. Either an empty pCAMBIA1301-220 vector or p1301-220-CmNRT2.1 was transformed into \textit{Agrobacterium tumefaciens} strain EHA105 using the freeze-thaw method\textsuperscript{29}. \textit{A. thaliana} Col-0 was transformed using the floral dip method\textsuperscript{18}. T\textsubscript{1} seedlings were raised on Murashige and Skoog (1962) (MS) medium containing 20 mg/L hygromycin and 25 mg/L ampicillin. Positive transformants were validated by observing GUS expression in the leaf\textsuperscript{29} and also by deploying a PCR assay on their genomic DNA, based on the primer pair CmNRT2.1-F/CmNRT2.1-R (Table 1).
qRT-PCR was used to quantify CmNRT2.1 transcription using the primer pair CmNRT2.1s/CmNRT2.1x (Table 1); the relevant reference gene was AtUBQ (NM_116771.5) assayed using the primer pair AtUBQs/AtUBQx (Table 1).

The performance of transgenic A. thaliana expressing CmNRT2.1. Seedlings were raised on vertical MS agar plates two weeks, then transferred to a growth culture conditions medium containing either 0.25 mM nitrate (0.125 mM KNO3, 0.0625 mM Ca(NO3)2) or 10 mM nitrate (5 mM KNO3 and 2.5 mM Ca(NO3)2). The nitrates were replaced by the same molarity of chloride salts to produce a nitrogen deficient medium. Each of three replicated treatments comprised a set of 50 plants. Shoot and root FW was measured after 14 days. The uptake of labeled nitrate (15NO3) was assayed as described elsewhere. Briefly, the plants were exposed to 0.1 mM CaSO4 for 1 min, then to a complete nutrient solution containing 0.2 mM 15NO3-for 5 min and finally to 0.1 mM CaSO4 for 1 min. The root homogenate was dried overnight at 80°C. The content of labeled nitrate was analyzed using a PDZ Europa ANCA-MS device (Northwich, UK). The recorded measurements represent the mean of three biological repeats.

Statistical analysis. Statistical analysis was performed by the one-way analysis of variance (ANOVA) using SPSS 11.5 software (SPSS Inc., Champaign, IL), and Duncan's multiple range test was employed to detect differences between means.

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**Author Contributions**

C.G., E.C. and S.C. conceived the study and designed the experiments. C.G., A.S., X.Z., T.L., Y.C. and J.J. performed the experiments and C.G., A.S., H.W. and T.L. analysed the data, and S.C. wrote the manuscript. All authors read and approved the final manuscript.

**Additional Information**

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