Importance of the $\alpha$C-helix in the cyclic nucleotide binding domain for the stable channel regulation and function of cyclic nucleotide gated ion channels in Arabidopsis

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Abstract

The involvement of cyclic nucleotide gated ion channels (CNGCs) in the signal transduction of animal light and odorant perception is well documented. Although plant CNGCs have recently been revealed to mediate multiple stress responses and developmental pathways, studies that aim to elucidate their structural and regulatory properties are still very much in their infancy. The structure–function relationship of plant CNGCs was investigated here by using the chimeric Arabidopsis AtCNGC11/12 gene that induces multiple defence responses in the Arabidopsis mutant constitutive expresser of PR genes 22 (cpr22) for the identification of functionally essential residues. A genetic screen for mutants that suppress cpr22-conferring phenotypes identified over 20 novel mutant alleles in AtCNGC11/12. One of these mutants, suppressor S58 possesses a single amino acid substitution, arginine 557 to cysteine, in the $\alpha$C-helix of the cyclic nucleotide-binding domain (CNBD). The suppressor S58 lost all cpr22 related phenotypes, such as spontaneous cell death formation under ambient temperature conditions. However, these phenotypes were recovered at 16°C suggesting that the stability of channel function is affected by temperature. In silico modelling and site-directed mutagenesis analyses suggest that arginine 557 in the $\alpha$C-helix of the CNBD is important for channel regulation, but not for basic function. Furthermore, another suppressor mutant, S136 that lacks the entire $\alpha$C-helix due to a premature stop codon, lost channel function completely. Our data presented here indicate that the $\alpha$C-helix is functionally important in plant CNGCs.

Key words: Calcium ion channel, CNGC, cpr22, cyclic nucleotide gated ion channel, environment, programmed cell death, temperature.

Introduction

For plant survival, cationic nutrients play essential roles in a wide variety of physiological aspects during growth and development. They also play important roles for signal transduction. For example, $Ca^{2+}$ and $K^+$ fluxes have been reported to be important for abiotic stress as well as biotic stress responses. Uptake and distribution of cationic nutrients mainly relies on membrane-localized cation transporter proteins. Based on genomic sequence data, Arabidopsis contains over 150 cation transport proteins (Mäser et al., 2001). Among them cyclic nucleotide-gated ion channels (CNGCs) form a large gene family consisting of 20 members that have been implicated in defence responses, development, and ion homeostasis in plants (Talke et al., 2003; Kaplan et al., 2007; Chin et al., 2009).

CNGCs were first discovered in retinal photoreceptors and olfactory sensory neurons and so far, six CNGC channel genes have been found in mammalian genomes (Zagotta and Siegelbaum, 1996; Zufall et al., 1994).
The structure of CNGCs is similar to that of the voltage-gated outward rectifying K⁺-selective ion channel (Shaker) proteins, including a cytoplasmic N-terminus, six membrane spanning regions (S1–S6), a pore domain located between S5 and S6, and a cytoplasmic C-terminus (Zagotta and Siegelbaum, 1996). However, CNGCs are opened by the direct binding of cyclic nucleotides (CN), such as cAMP and cGMP (Fesenko et al., 1985). The CN-binding domain (CNBD) of CNGCs is located at the cytoplasmic C-terminus and exhibits significant sequence similarity to that of protein kinase A, protein kinase G, and the catabolite activator protein of E. coli (Bridges et al., 2005). The cytoplasmic C-terminus contains a CNBD that is connected to the pore domain by a C-linker region. Important functional features of CNGCs were extensively studied in animal systems and it has been suggested that the subunit composition of the respective channel complex is an important determinant for functional features such as ligand sensitivity, selectivity, and gating. (Kaupp and Seifert, 2002).

The first plant CNGC, HvCBT1 ( Hordeum vulgare calmodulin (CaM)-binding transporter), was identified as a CaM-binding protein in barley (Schuurink et al., 2002). Subsequently, several CNGCs were identified from *Arabidopsis* and *Nicotiana tabacum* (Arazi et al., 1999; Köhler et al., 1999). A precise analysis of the CaM binding site has been reported using the tobacco CNGC, NtCBP4 as well as *Arabidopsis* AtCNGC1 and AtCNGC2 and it has been suggested to be located at the αC-helix of the CNBD in these plant CNGCs (Arazi et al., 2000; Köhler and Neuhaus, 2000). However, only a handful of studies on the structure–function analysis of plant CNGCs have so far been published (Hua et al., 2003; Bridges et al., 2005; Kaplan et al., 2007; Baxter et al., 2008).

The *Arabidopsis* mutant constitutive expreser of PR genes 22 (cpr22), which contains a novel chimeric CNGC, AtCNGC11/12, shows environmentally sensitive defence responses, such as heightened salicylic acid (SA) accumulation and a hypersensitive response (HR)-like programmed cell death (Yoshioka et al., 2001; Moeder and Yoshioka, 2009). It has been reported that the expression of AtCNGC11/12 and its channel activity is attributable for the cpr22 phenotype (Yoshioka et al., 2006; Baxter et al., 2008). Interestingly, all SA dependent phenotypes, are suppressed under high humidity conditions and enhanced by low humidity (Yoshioka et al., 2001). This type of environmental sensitivity has been reported for various pathogen resistance mutants as well as on defence responses in wild-type plants (Moeder and Yoshioka, 2009).

The importance of the interaction between the C-linker domain and the CNBD domain for basic channel function was reported previously by Baxter et al. (2008). In addition, it was shown that intragenic mutants of cpr22 are useful tools to study the structure-function relationship of CNGCs (Baxter et al., 2008). The use of intragenic mutants of cpr22 were used here for the structure–function study and to reveal the importance of the αC-helix of the CNBD for stable channel function in planta.

**Materials and methods**

**Plant growth conditions**

*Arabidopsis thaliana* plants were grown on Pro-Mix soil (Premier Horticulture Inc., Red Hill, PA, USA) in a growth chamber under ambient humidity as described by Silva et al. (1999). *Nicotiana benthamiana* plants were grown on the same soil in a greenhouse under a 14:10 h light:dark regimen at 25 °C (day) and 20 °C (night).

**Suppressor screening and identification of the S58 mutant**

Suppressor no. 58 (S58) was identified in the same screen that was described in Baxter et al. (2008).

**Trypan blue staining**

Leaf samples were taken from 3–4-week-old plants grown on soil. Trypan blue staining was performed as described previously (Yoshioka et al., 2001).

**RNA extraction, RT-PCR, and Northern hybridization**

Small-scale RNA extraction was carried out using the TRIzol reagent (Invitrogen, Carlsbad, MO, CA), according to the manufacturer’s instructions. Reverse transcriptase (RT)-PCR was performed using cDNA generated by SuperScript™ II Reverse Transcriptase (Invitrogen, Carlsbad, MO, CA) according to the manufacturer’s instructions. For the detection of CNGC gene expression in *N. benthamiana*, CNGC, and actin gene expression in yeast, and β-tubulin gene expression in *Arabidopsis*, the same sets of primers described by Baxter et al. (2008) were used. For the detection of PR-1 by RT-PCR, PR-1F: 5'-GCTCTTTGA-GGTGCTCTTTGTTTTTT and PR-1R: 5'-CACGCTTTATTTGTATATT- TATT-3' were used. Northern hybridization of the PR-1 gene was performed as previously described (Yoshioka et al., 2006). For the RT-PCR, 25 cycles for each analysis were applied.

**Pathogen infection**

Infection with *Hyaloperonospora arabidopsidis* isolate Emw1 was performed as described previously (Yoshioka et al., 2001).

**Plasmid construction**

The yeast expression vector plasmid pYES2-empty vector (Invitrogen), pYES2-AtCNGC12, pYES2-AtCNGC11/12, and pYES2-AtCNGC11/12:ES19K (S73) were constructed as previously described (Yoshioka et al., 2006; Baxter et al., 2008). For pYES2-AtCNGC11/12:R557C (S58), total RNA was extracted from S58 and cDNA was generated as described above. cDNA of AtCNGC11/12:R557C was then amplified by RT-PCR using primers: BamH-ATG3: 5'-GGGATCCCATGAATCTTCAGAG-GAGAAA-3' and cDNA14-R1: 5'-CAGATGCTCTTACGCTT- TGC-3', and then cloned into pGEM-T Easy (Promega, Madison, WI). In the case of pYES2-AtCNGC11/12:Q543X (S136), AtCNGC11/12 cDNA was used as template for PCR using primers: BamH-ATG3: 5'-GGGATCCCATGAATCTTCAGAG-GAGAAA-3' and Reverse 136 stop mutation R: 5'-TTATCTTT-GAAAGACATTTA-3'. Both cDNA clones were then subcloned into pYES2 at the BamHI and NotI restriction enzyme sites. AtCNGC-R557C (S58) cDNA minus stop was generated by RT-PCR using forward 5'-CTCTAGACATGAATCTTTGAGAG-GAGAAA-3' and reverse 5'-AGTCTAGATGCTCTAGGCTT- GC-3' primers and subsequently cloned into pGEM-T Easy (Promega, Madison, WI). This clone was then subcloned into the *Xba*I site in pMPB3 (Yoshioka et al., 2006; Urquhart et al., 2007). For site-directed mutagenesis, *AtCNGC-R557C* (S58) cDNA was excised using *Xba*I and subsequently subcloned into pBluescript (Stratagene, La Jolla, CA). The point mutation R557I (C1627T)
was introduced into *AtCNGC11/12* cDNA in pBluescript (Baxter et al., 2008) using the QuickChange site-directed mutagenesis kit (Stratagene) according to the manufacturer’s instructions. The *AtCNGC11/12:*R557I cDNA was then subcloned into pMBP3 for transient assays in *N. benthamiana*. All constructed plasmids were sequenced for fidelity.

**Agrobacterium-mediated transient expression**

*Agrobacterium*-mediated transient expression in *N. benthamiana* was performed as described by Urquhart et al. (2007). The expression of these genes was confirmed by RT-PCR (see RNA extraction and RT-PCR).

**Functional complementation in yeast**

*K*⁺ yeast mutant complementation: The *K*⁺-uptake deficient yeast mutant strain CY162 (MAT α, ura3-52, trk1,2) was transformed with the empty plasmid pYES2, pYES2-AtCNGC12, pYES2-AtCNGC11I2, pYES2-AtCNGC11I2:R557C, pYES2-AtCNGC12:R557I and pYES2-AtCNGC11I2:Q543X following the lithium acetate transformation protocol (Ausubel et al., 1987). Yeast transformants were grown in synthetic minimal media supplemented with 0.1 mM KCl (Leng et al., 1999) and hygromycin (5 mg l⁻¹) (Mercier et al., 2004) to suppress background growth. Growth rates were monitored by determining the OD₆₀₀ of the growing yeast cultures at 20 h and 40 h.

**Ca²⁺ yeast mutant complementation**: A wild-type strain of *S. cerevisiae* W303-1A (Wallis et al., 1989) and the Ca²⁺ channel mutant (*cch1*:TRP1 null mutant) strain K927 (Locke et al., 2000) were provided by Dr H Iida (Tokyo Gakugei University). K927 was transformed with pYES2 empty vector, pYES2-AtCNGC12, pYES2-AtCNGC11I2, pYES2-AtCNGC11I2:R557C, pYES2-AtCNGC12:R557I, and pYES2-AtCNGC11I2:Q543X following the lithium acetate transformation protocol (Ausubel et al., 1987).

To test for complementation of the *cch1* mutation, yeast transformants were grown to logarithmic phase in synthetic minimal media and were diluted to 10⁶ cells ml⁻¹ and exposed to 20 μM α-mating factor in modified synthetic minimal media containing 100 μM CaCl₂ (Muller et al., 2001). 100 μl aliquots of cells were harvested by centrifugation and resuspended in 10 mg ml⁻¹ Trypan blue solution at 4, 8, and 12 h. Yeast viability was measured by assessing the ratio of stained to unstained cells under brightfield microscopy. A minimum of 200 cells were counted for each transformant.

Ca²⁺ uptake was measured by the method described by Kurusu et al. (2004) with slight modifications. K927 transformants were grown to logarithmic phase in synthetic minimal media and diluted to 10⁶ cells ml⁻¹. Cultures were then preincubated for 120 min at 30 °C in 10 mM MES-TRIS buffer (pH 6.0) containing 100 mM glucose. 4⁵CaCl₂ was added to a final concentration of 72 kBq ml⁻¹. 100 μl aliquots were harvested at 10, 20, and 30 min by centrifugation and washed five times with washing solution (20 mM MgCl₂, 0.1 mM LaCl₃). Radioactivity in yeast cells was measured using a liquid scintillation counter. All experiments have been conducted with three biological repeats with three technical repeats.

Green fluorescence protein (GFP) visualization by confocal microscopy

*Agrobacterium*-mediated transient expression in *N. benthamiana* was performed as described in Urquhart et al. (2007) at 16 °C and 22 °C. Plants were shifted to 16 °C 12 h after infiltration, and protein stability in either condition was confirmed by GFP expression at 30 h. Small sections of the infiltrated area were excised and used for confocal microscopy. Confocal fluorescence images were acquired using a Leica TCS SP5 confocal system with AOBS® (HCS PL APO CS 40× immersion oil objective; NA, 1.25) with the AOTF for the argon laser (488 nm) set at 35% and the detection window at 500–600 nm (Leica Microsystems Inc., Wetzlar, Germany).

**Computational modelling and sequence alignment**

The tertiary structure modelling of *AtCNGC11/12* was conducted as described previously (Baxter et al., 2008) using the crystalized structure of the cytoplasmic C-terminus of invertebrate CNGC, SpH (Flynn et al., 2007, PDB no. 2PTM). The protein fold recognition server (Phyre; Kelley and Sternberg, 2009) was used to model the protein coordinates with estimated precision of 100%. All the images were generated using PyMOL (DeLano, 2002).

The sequence alignments of the CNBD amino acid sequences of all 20 *Arabidopsis* CNGCs were aligned using ClustalW (Thompson et al., 1994). The accession numbers of all protein sequences used for the alignment are indicated in the figure legends.

**Results**

**The intragenic suppressor S58 loses cpr22 related phenotypes under ambient conditions**

The suppressor screen of *cpr22* was reported previously (Baxter et al., 2008). Through this screen more than 20 novel mutant alleles in *AtCNGC11/12* have been discovered so far. One of these mutants, suppressor S58 was found to be morphologically identical to the wild type (wt) plants (Fig. 1a). The homozygosity of the *cpr22* allele in S58 was confirmed by PCR-based marker analysis (data not shown). In contrast to homozygous *cpr22*, S58 is neither lethal, nor shows HR-like spontaneous lesion formation and constitutive *PR-1* gene expression (Fig. 1a, b). Pathogen resistance was evaluated using the oomycete pathogen, *Hyaloperonospora arabidopsidis*, isolate Emw1, which is virulent for the ecotype Wassilewskija (Ws, the background ecotype of *cpr22*). *cpr22* showed enhanced resistance to *Emw1* (Yoshioka et al., 2006). As predicted, suppressor S58 lost *cpr22*-mediated enhanced resistance to this pathogen (Fig. 1c; see Supplementary Table S1 at *JXB* online). Taken together; it is concluded that suppressor S58 lost all tested *cpr22*-related phenotypes under ambient temperature condition (22 °C). Sequence analysis of the *cpr22* gene (*AtCNGC11/12*) revealed one nucleotide substitution, C to T in the cyclic nucleotide binding domain (CNBD) that caused an amino acid substitution, Arginine 557 to Cysteine (T in the cyclic nucleotide binding domain (CNBD) that caused an amino acid substitution, Arginine 557 to Cysteine (C) revealed one nucleotide substitution, C to T in the cyclic nucleotide binding domain (CNBD) that caused an amino acid substitution, Arginine 557 to Cysteine (C) revealed one nucleotide substitution, C to T in the cyclic nucleotide binding domain (CNBD) that caused an amino acid substitution, Arginine 557 to Cysteine (C) revealed one nucleotide substitution, C to T in the cyclic nucleotide binding domain (CNBD) that caused an amino acid substitution, Arginine 557 to Cysteine (C) revealed one nucleotide substitution, C to T in the cyclic nucleotide binding domain (CNBD) that caused an amino acid substitution, Arginine 557 to Cysteine (C) revealed one nucleotide substitution, C to T in the cyclic nucleotide binding domain (CNBD) that caused an amino acid substitution, Arginine 557 to Cysteine. This indicated the semi-dominant nature of this mutation (22:1 (wild type like):2 (heterozygous-like):1 (lethal).

**Green fluorescence protein (GFP) visualization by confocal microscopy**

*Agrobacterium*-mediated transient expression in *N. benthamiana* was performed as described in Urquhart et al. (2007) at 16 °C and 22 °C. Plants were shifted to 16 °C 12 h after infiltration, and protein stability in either condition was confirmed by GFP expression at 30 h. Small sections of the infiltrated area were excised and used for confocal microscopy. Confocal fluorescence images were acquired using a Leica TCS SP5 confocal system with AOBS® (HCS PL APO CS 40× immersion oil objective; NA, 1.25)
The S58 mutation (R557C) does not affect channel function in yeast

Growth enhancement of mutant yeast has been previously demonstrated upon expression of various plant CNGCs (Köhler et al., 1999; Leng et al., 1999; Ali et al., 2006). More recently, it was suggested that AtCNGC11, 12, and AtCNGC11/12 can function as K⁺ channels as well as Ca²⁺ channels when expressed in the same systems (Yoshioka et al., 2006; Urquhart et al., 2007). Therefore, it is possible that the loss of cpr22-related phenotypes in S58 can be attributed to either a loss in basic ion channel function, or a loss in the constitutive active character of AtCNGC11/12 while still maintaining basic channel function. To address this point, a heterologous expression system using K⁺ and Ca²⁺ uptake-deficient yeast mutants was used to evaluate the channel function of AtCNGC11/12:R557C. The trk1,trk2 K⁺ uptake-deficient yeast mutant, CY162 (Leng et al., 1999), transformed with AtCNGC11/12 or AtCNGC11/12:R557C was tested in low external K⁺ and in the presence of the cationic antibiotic hygromycin. As shown in Fig. 2a, AtCNGC11/12 and AtCNGC12 were able to complement the trk1,trk2 phenotype compared to the empty vector control. Interestingly, the mutants carrying AtCNGC11/12:R557C were also able to complement the trk1,trk2 phenotype with the same efficiency as AtCNGC11/12, whereas another CNBD mutant, S73 (E519K), which affects ion channel function (Baxter et al., 2008) did not, suggesting that the R557C mutation does not affect its basic ion channel function. To explore if this is also the case for Ca²⁺ channel function, a similar complementation analysis was conducted using the Ca²⁺-uptake yeast mutant strain K927 (chl::TRP1; Locke et al., 2000). CCH1 has been implicated in Ca²⁺ influx in response to mating pheromones (Fischer et al., 1997). Previously, it has been shown that Ca²⁺ is important for cell death induction by the expression of AtCNGC11/12 (Urquhart et al., 2007). Therefore, it is possible that only Ca²⁺ channel function is affected by the S58 mutation. However as shown in Fig. 2c by Trypan blue staining, AtCNGC11/12, AtCNGC12, and AtCNGC11/12:R557C rescued this yeast phenotype to comparable levels, indicating that AtCNGC11/12:R557C is functional as a Ca²⁺ ion channel. These data were quantitatively confirmed in Fig. 2b. In addition, ⁴⁵Ca²⁺ uptake analysis was conducted using the same yeast mutant strain. Although the ⁴⁵Ca²⁺ uptake did not reach the same level as wild-type yeast, K927 mutant yeast with AtCNGC11/12 or AtCNGC11/12:R557C had a higher uptake rate of ⁴⁵Ca²⁺ than K927 carrying the empty vector (see Supplementary Fig. S1 at JXB online). The same result was observed in four independent experiments. No significant difference between AtCNGC11/12 and AtCNGC11/12:R557C was observed, suggesting that the R557 (S58) mutation does not impair the function of AtCNGC11/12. This result is consistent with the mating pheromone analyses using the same yeast mutant strain (Fig. 2b). Taken together, it was concluded that the mutation R557C does not affect basic ion channel function but rather affects the regulation of AtCNGC11/12.

AtCNGC11/12:R557C recovers the activity to induce cell death under low temperature

As mentioned above, cpr22-related phenotypes show temperature sensitivity. Relatively high temperatures (>28 °C) can suppress its phenotypes and on the other hand relatively lower temperature (<16 °C) enhanced them (Mosher et al., 2010). Since the R557C mutation seems to suppress the constitutive active character of AtCNGC11/12 but not basic ion channel function, the question was asked if lower temperature can restore this constitutive active character (channel activity). Strikingly, a temperature shift from 22 °C to 16 °C induced cpr22-like morphology, such as curly leaves 4 d after shift in S58. Chlorosis on leaves, which indicates cell death development, could also be observed (Fig. 3a). Trypan blue staining as well as electrolyte leakage analyses further confirmed cell death development in S58 (Fig. 3a, b). PR-1 gene expression also recovered after the

Fig. 1. Characterization of the suppressor mutant S58. (a) Morphological phenotypes and spontaneous HR cell death formation of wild type (Ws-wt), cpr22, and suppressor 58 (S58). A cpr22 homozygous plant is shown in the white square. Approximately 4-week-old plants were used. (b) Northern blot analysis for PR-1 gene expression in Ws-wt, cpr22, and S58. The samples were taken from approximately 4-week-old plants. Ethidium bromide staining of ribosomal RNA (rRNA; lower panel) served as a loading control. (c) Growth of _Hyaloperonospora arabidopsidis_, isolate Emw1 in Ws-wt, cpr22, and S58. Plants were infected by spraying a conidiospore suspension of 10⁶ ml⁻¹ on 7-d-old plants. The Trypan blue analysis 8 d after infection was done to visualize pathogen growth.
shift from 22 °C to 16 °C (Fig. 3c), further supporting the temperature sensitivity of this mutant.

To characterize the temperature sensitivity of S58 further (AtCNGC11/12:R557C), Agrobacterium-mediated transient expression analysis was conducted in Nicotiana benthamiana. This is an established system to analyse HR cell death development and has been used to show that AtCNGC11/12 induces cell death in a synchronized manner (Urquhart et al., 2007). As shown in Fig. 4a, cell death was induced
by the transient expression of \textit{AtCNGC11/12} but not \textit{AtCNGC11/12:R557C} at 22 °C (ambient temperature) indicating the inactivity of \textit{AtCNGC11/12:R557C} in planta under this condition. On the other hand, \textit{AtCNGC11/12:R557C} recovered cell death induction in plants that were shifted from 22 °C to 16 °C. This was also confirmed quantitatively by electrolyte leakage analysis (Fig. 4b). Transcript and protein levels were not significantly altered by the temperature shift as depicted by RT-PCR (Fig. 4c) and GFP fluorescence (Fig. 4d). Note that while we have observed enhanced cell death in \textit{cpr22} when it was shifted from 22 °C to 16 °C (Fig. 3a), there is no significant difference between 22 °C and 16 °C when \textit{AtCNGC11/12} was transiently expressed in \textit{N. benthamiana}. This is probably due to the sampling timing. The samples for Fig. 4b were taken at 80 h after \textit{Agrobacterium} infiltration. At this time point, cell death development induced by transient expression of \textit{AtCNGC11/12} is almost completed and also it is very strong and uniform compared to the mutant itself due to the constitutive CaMV35S promoter.

3D computational modelling of \textit{AtCNGC:R557C}

In order to predict the role of R557 in channel structure, a computational analysis of the three-dimensional structure of the cytoplasmic C-terminal region was conducted. Previously, the crystal structure of the cytoplasmic C-terminal region of a hyperpolarization-activated cyclic nucleotide-modulated channel, HCN2, has been used as a template (Zagotta et al., 2003, PDB ID: 1Q50; Baxter et al., 2008). The recently published crystal structure of another HCN, SpIH was used here (crystallized with cAMP; Flynn et al., 2007; PDB ID: 2PTM) that possesses higher structural similarity than HCN2 to \textit{AtCNGC11/12} (H Abdel-Hamid and D Shahinas, unpublished data). As shown in Fig. 5a, R557 is located in the middle of the \(\alpha\)-helix in the CNBD. The importance of this \(\alpha\)-helix for cNMP binding has been reported in animal CNGCs (Rehmann et al., 2007). According to the crystal structure of the CNBD of HCN2 in the presence of cAMP, cAMP binds in the anti-configuration between the \(\beta\)-roll and the \(\alpha\)-helix of the CNBD. In our model, the side chain of R557 faces away from the interior of the cNMP binding pocket formed by the \(\beta\)-roll of the CNBD and therefore, likely does not bind to cNMPs directly (Fig. 5a). Hydrophobic interactions between the \(\alpha\)-helix and the base of cNMPs were also postulated to stabilize cNMP binding (Rehmann et al., 2007), further suggesting that the hydrophilic R557 does not directly interact with cNMPs.

Fig. 4. Temperature sensitivity of cell death induction by transient expression of \textit{AtCNGC11/12}, empty vector, S58 (\textit{AtCNGC11/12:R557C}), and S73 (\textit{AtCNGC11/12:E519K}) in \textit{Nicotiana benthamiana}. (a) Induction of cell death in \textit{N. benthamiana} 24, 48, and 80 h after \textit{Agrobacterium} infiltration, either shifted from 22 °C to 16 °C at 12 h after \textit{Agrobacterium} infiltration (lower panels) or not shifted (upper panels). Cell death induction was observed in the leaf area expressing S58, but not empty vector (EV) or S73 after the temperature shift. Cell death induced by \textit{AtCNGC11/12} was enhanced by temperature shift. Red circles indicate HR development. (b) Quantitative analysis of cell death in \textit{N. benthamiana} by electrolyte leakage of leaf discs. S58 expression induced cell death after the temperature shift, but not empty vector (EV) or S73. Samples were taken 80 h after \textit{Agrobacterium} infiltration. (c) RT-PCR analysis of leaf discs from \textit{N. benthamiana} leaves expressing \textit{AtCNGC11/12}, S58 or empty vector (EV). The temperature shift did not significantly affect gene expression of \textit{AtCNGC11/12} or \textit{AtCNGC11/12:R557C} in \textit{N. benthamiana} leaf discs. \textit{actin} served as a loading control. Samples were taken 24 h after \textit{Agrobacterium}-infiltration (12 h after the shift). (d) The expression of \textit{AtCNGC11/12}:GFP and \textit{AtCNGC11/12:R557C}:GFP was not altered by the temperature shift. The samples were taken 30 h after \textit{Agrobacterium} infiltration. The fluorescence of the GFP-fusion proteins was monitored by confocal microscopy.
A sequence comparison revealed that R557 is conserved in 17 out of 20 *Arabidopsis* CNGCs (Fig. 5b). Only three CNGCs, AtCNGC1, 3, and 11 have a lysine (K) or glutamine (Q) that mediate ionic interactions similarly to arginine (R) in this position. All of them are hydrophilic (hydropathy index: -4.5). Cysteine is slightly polar due to its -SH group but does not mediate any ionic interactions (hydropathy index: +2.5). To address whether a more hydrophobic residue with less polarity than cysteine such as isoleucine (hydropathy index: +4.5) can completely disrupt channel function, AtCNGC11/12:R557I was created by site-directed mutagenesis and it was transiently expressed in *N. benthamiana* to assess cell death induction. As shown in Fig. 6a, cell death is not induced by AtCNGC11/12:R557I at 22°C, but is recovered at 16°C similar to AtCNGC11/12:R557C. The expression of all constructs was confirmed.

**Fig. 5.** The location of R557 and Q543 in tertiary structure and amino acid sequence alignment. (a) Ribbon diagram of the cytoplasmic C-terminal region of AtCNGC11/12 (AtCNGC12) (left panel) and close-up of the indicated area of the left panel (right panel). R557 is located in the αC-helix and Q543 is located in the αB-helix of the CNBD. cAMP is indicated by pink colour. (b) Alignment of the area of R557 with 20 *Arabidopsis* CNGCs and tobacco NtCBP4. NCBI:AF079872, AtCNGC3:CAB40128, AtCNGC11:AAD20357, AtCNGC10:AAF73128, AtCNGC13:AAL27505, AtCNGC1:AAK43954, AtCNGC12:Ad23055, AtCNGC12 (Ws ecotype):EU541495, AtCNGC6:AAC63666, AtCNGC9:CAB79774, AtCNGC5:TS2573, AtCNGC7:AG12662, AtCNGC8:NP_173408, AtCNGC15:AAD29827, AtCNGC14:AAD23886, AtCNGC17:CAB4129, AtCNGC16:CAB4139, AtCNGC18:CAC01828, AtCNGC19:BAB02061, AtCNGC20::BAB02062, AtCNGC2:CAC01740, AtCNGC4:T52574. The black box indicates the position of R557. The red box indicates the CaM binding domain and bold characters highlight the critical four amino acids for the CaM binding suggested by Arazi et al. (2000). The location of Q543 (S136) is indicated by a black dot.
by RT-PCR (Fig. 6b). Therefore, even by altering R557 to isoleucine complete suppression of channel function at 16 °C could not be obtained.

The αC and αB helices in the CNBD are crucial for channel function

The αC-helix in the CNBD was suggested to play an important role for animal CNGC function/channel gating (Rehmann et al., 2007). However, so far there is almost no report about the importance of the αC-helix in plant CNGCs except one report showing that the CaM binding domain is located in the αC-helix (Arazi et al., 2000). To study the role of the αC-helix in channel function further, another intragenic suppressor mutant, S136, was used that has a premature stop codon at Q543 (C to T point mutation) in the CNBD. Our structural model revealed Q543 to be located at the middle of the αB helix, indicating that S136 does not have the entire αC-helix and only a partial αB helix (Fig. 5a). S136 did not display cpr22 phenotypes under 22 °C or 16 °C (Fig. 7a). It also lost constitutive expression of PR-1 and enhanced pathogen resistance (Fig. 7b, c). Furthermore, heterologous expression in the previously mentioned trkl, trk2 K+ and cehl Ca2+ yeast mutants revealed that AtCNGC11/12:Q543X does not have channel function (Fig. 7d; data not shown). Taken together, these data suggest that the αC-helix and possibly the αB-helix of the CNBD is functionally essential for CNGC function in plants.

Discussion

Through this study it has been found that R557 in the αC-helix of the CNBD plays an important role in stable channel regulation. Substitution of R557C in the Arabidopsis mutant cpr22 impaired cell death formation and other cpr22-related phenotypes suggesting a disruption of channel function. However, the cpr22-related phenotypes as well as cell death caused by transient expression of AtCNGC11/12:R557C could still be observed under slightly lower temperatures (16 °C) suggesting an alteration in channel regulation rather than channel function. Yeast complementation analysis demonstrated that AtCNGC11/12:R557C maintains its basic channel function for both K+ and Ca2+ conductance, further supporting this notion. R557 is highly conserved in the CNGC family and all residues in this position share similar polarity and hydrophobicity. It has been examined if a change to an even more hydrophobic and less polar residue (R557I) would have a stronger effect that completely disrupts basic channel function. However, this channel remained to be conditional just like AtCNGC11/12:R557C, further strengthening the notion that R557 is important for the regulation of channel activity rather than basic channel function.

In animal systems, it has been reported that cNMPs bind within the pocket formed by the αC-helix and the β-barrel composed of the eight β sheets in the CNBD (Weber and Steitz, 1987; Rehmann et al., 2007). The αC-helix was suggested to function as the lid of this pocket that stabilizes the cNMP binding by forming hydrophobic interactions with the bound cNMP (Rehmann et al., 2007). However, because R557 is hydrophilic, it does not seem to participate directly in cNMP binding. Our computational modelling also supported this notion.

Regarding the role of the αC-helix in plant CNGCs, a 19–20 amino acid sequence of this region was suggested to be the CaM binding domain in AtCNGC1 and AtCNGC2 by Köhler and Neuhaus (2000) using yeast two hybrid analysis. Arazi et al. (2000) biochemically demonstrated that a 23 amino acid sequence overlapping with this 19–20 amino acids is the CaM binding domain in the tobacco CNGC, NtCBP4. Furthermore, they reported that the four additional amino acids (W R T/S W) which are located just outside of the 19–20 amino acid sequence are crucial for efficient binding. As shown in the alignment in Fig. 5b, R557 is located in this crucial sequence (indicated by bold characters). Considering the fact that R557C altered its regulation, not the basic channel function itself, it can be hypothesized that R557C displays an alteration in its...
binding affinity to CaM which causes an alteration to the regulation of its function. Considering the importance of CaM in regulating this channel group, our finding is significant. Biochemical studies that aim to identify the CaM interactions with AtCNGC11/12 (AtCNGC12) are underway.

The experimental results involving of suppressor S136 was unexpected, since the deletion of the part of the CNBD that includes the CaM binding site has been reported to enhance the AtCNGC1 channel function in yeast (Ali et al., 2006). Ali et al. (2006) demonstrated that two deletion constructs of AtCNGC1 showed enhanced K⁺ channel function in a yeast complementation analysis. They hypothesized that the binding of the negative regulator CaM does not occur in these deletion constructs thereby promoting channel activity. Based on our computational modelling, the deletion of S136 is positioned between these two constructs (data not shown). However, we did not see any channel function using both K⁺ and Ca²⁺ yeast mutants in the case of S136, suggesting the importance of the αC-helix and possibly the αB-helix for its basic channel function. This discrepancy could be due either to differences in regulatory properties of different AtCNGC subunits or to unknown structural changes caused by the different deletion positions used in both studies. As of now, the answer remains unknown and further analyses will be required to understand these differences.

In this study, it has been demonstrated that R557 in the αC-helix of the CNBD of CNGCs plays an important role for channel regulation and that the αC-helix and possibly αB-helix are crucial for plant CNGC channel function. It has been proposed that the αC-helix binds to both cNMP and CaM as a mechanism to regulate CNGC channel gating (Arazi et al., 2000). Our data here demonstrates the importance of the αC-helix in a plant CNGC for the first time. Further precise structural and biochemical studies to elucidate the role of the αC-helix are underway.

Supplementary data

Supplementary data are available at JXB online.

**Supplementary Fig. S1.** Ca²⁺-uptake analysis using Calcium-45 AtCNGC11/12 and S58 (AtCNGC11/12:R557C) partially complemented the Ca²⁺-uptake deficient mutant K927, whereas the empty vector did not.

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**Supplementary Fig. S1.** Ca²⁺-uptake analysis using Calcium-45 AtCNGC11/12 and S58 (AtCNGC11/12:R557C) partially complemented the Ca²⁺-uptake deficient mutant K927, whereas the empty vector did not.
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