The CNGCb and CNGCd genes from *Physcomitrella patens* moss encode for thermosensory calcium channels responding to fluidity changes in the plasma membrane

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**Abstract** Land plants need precise thermosensors to timely establish molecular defenses in anticipation of upcoming noxious heat waves. The plasma membrane-embedded cyclic nucleotide-gated calcium channels (CNGCs) can translate mild variations of membrane fluidity into an effective heat shock response, leading to the accumulation of heat shock proteins (HSP) that prevent heat damages in labile proteins and membranes. Here, we deleted by targeted mutagenesis the *CNGCd* gene in two *Physcomitrella patens* transgenic moss lines containing either the heat-inducible HSP-GUS reporter cassette or the constitutive UBI-Aequorin cassette. The stable CNGCd knockout mutation caused a hyperthermosensitive moss phenotype, in which the heat-induced entry of apoplastic Ca\(^{2+}\) and the cytosolic accumulation of GUS were triggered at lower temperatures than in wild type. The combined effects of an artificial membrane fluidizer and elevated temperatures suggested that the gene products of *CNGCd* and *CNGCb* are paralogous subunits of Ca\(^{2+}\) channels acting as a sensitive proteolipid thermocouple. Depending on the rate of temperature increase, the duration and intensity of the heat priming preconditions, terrestrial plants may thus acquire an array of HSP-based thermotolerance mechanisms against upcoming, otherwise lethal, extreme heat waves.

**Keywords** Plant heat shock response · Acquired thermotolerance · Ca\(^{2+}\) channels · *Physcomitrella patens* · Heat shock proteins

**Abbreviations**

CNGC Cyclic nucleotide-gated channel
HSR Heat shock response
HSPs Heat shock proteins
BA Benzyl alcohol
ET\(_{50}\) Physiological temperature at which half of the maximal activatory effect is observed
IT\(_{50}\) Excessive temperature at which half of the maximal inhibitory effect is observed

**Introduction**

On the break of a hot summer day, land plants in dry mid-latitude climates (Peel et al. 2007) need sensitive thermosensors to detect mild increments in the ambient temperature and trigger an appropriate heat shock response (HSR) in anticipation of upcoming extreme midday temperatures (Mittler et al. 2012). At the cellular level, the HSR is an appropriate accumulation of a complex network of protective heat shock proteins (HSPs), the most abundant of which are molecular chaperones that can prevent heat-induced misfolding and aggregation of labile proteins (Hinault et al. 2006; Sharma et al. 2010; Finka et al. 2011) and protect membranes from heat damage (Torok et al. 1997). At physiological temperatures, proteins and membranes are expected to be optimally stable and functional. At high excessive temperatures, the increased thermal movements of macromolecules may transiently dissociate protein complexes, unfold polypeptides, and decrease lipid order in membranes thus leading to the formation of inactive, potentially toxic, protein aggregates and hyperfluid membranes prone to disruption (Mittler and Blumwald 2010). Hence, it is of primary importance that plant cells can anticipate an upcoming noxious heat shock by detecting mild increments in the ambient temperature before labile macromolecular...
complexes are damaged and timely establish effective molecular defenses in the form of detoxifying enzymes, such as ascorbate peroxidase and of unfolding chaperones, such as Hsp101, Hsp90, Hsp70, and Hsp60, which can avert heat-induced protein aggregations (Finka et al. 2011; Mittler et al. 2012; Sharma et al. 2011), as well as small HSPs that can prevent protein aggregation (Veinger et al. 1998) and protect membranes against hyperfluidity (Toro et al. 1997; Horvath et al. 1998, 2012).

Because the massive accumulation of HSPs has a very high cellular cost, both in terms of energy and resources, the HSR needs to be finely tuned. Therefore, plant cells need precise thermosensors that can react to mild temperature increments, for examples when the sun rises, and send an appropriate timely signal to synthesize within 2–8 h protective HSPs to withstand the noxious heat at noon. During an abrupt continuous heat shock, as in the laboratory, the levels of HSP mRNAs in the plant cells strongly increase during the first 30 min and then rapidly decrease to reach ground levels within 1–2 h, despite the ongoing heat-inducing conditions (Saidi et al. 2005, 2009; Finka et al. 2011). Yet, when plants are treated as in nature with a gradual increase of temperatures during several consecutive hours, to slowly reach the noxious temperatures within 6 h (Larkindale and Vierling 2008), this produces more HSP mRNAs and better plant survival than a shorter and milder stepwise heat priming treatment (Larkindale and Vierling 2008). This suggests that during the gradual increments of the heat, plants may use a series of distinct heat sensors that become successively activated at different temperature thresholds and thus generate successive waves of HSP synthesis. Such a scenario might result in the observed effective continuous accumulation of protective HSPs, despite the transient nature of the heat activation process (Larkindale and Vierling 2008; Saidi et al. 2009). It is thus central to understand the molecular nature of the primary heat sensors and resolve how, despite the transient entry of Ca$^{2+}$ and mRNA accumulation within minutes, plant cells may increase their HSP load during hours and develop acquired thermotolerance lasting days (Cohen-Peer et al. 2010).

At physiological temperatures, the plant heat shock transcription factors (HSFs) are inactive. Like in animals and fungi, they are hypo-phosphorylated monomers, which may be found in association with cytosolic Hsp90s and Hsp70s (Hahn et al. 2011; Scharf et al. 2012). Under heat shock conditions, the higher plant non-heat-inducible HSFA1a and the heat-inducible HSFA2 become hyper-phosphorylated and may form heterocomplexes that rapidly enter the nucleus, where they may specifically activate the transcription of about a thousand HSP genes (Li et al. 2010; Busch et al. 2005). In Arabidopsis, the overexpression of SUMO1, which binds to the heat-inducible HSFA2, represses the transcription of reporters transgenes from Hsp17.4 and Hsp17.6 and Hsp101 promoters and decreases plant acquired thermotolerance (Cohen-Peer et al. 2010).

To specifically transcribe the thousand or so of heat-induced genes in plants (Finka et al. 2011; Mittler et al. 2012), the heat-activated plant HSFs must specifically bind to particular DNA motives, also called heat shock elements (HSEs), upstream to each of the 1,780 HSP genes in Arabidopsis (following 90 min at 38 °C) or 1,509 in wheat (following 60 min at 40 °C) (Finka et al. 2011; Qin et al. 2008).

Under conditions that are unchallenging for the HSP genes, histones forming nucleosomes are often considered as a transcriptional roadblock (Kumar and Wigge 2010). The binding of active HSFs to HSE, which needs to occur at nucleosome-free stretches of DNA (Petesch and Lis 2008; Petesch and Lis 2012) can activate chromatin remodeling complexes to modify nearby bound histones (Clapier and Cairns 2009), thereby unleashing the bound RNA polymerase to massively transcribe the HSP genes (Mittler et al. 2012; Petesch and Lis 2012).

In contrast to histones, which are likely to be the most downstream components of the heat shock signaling pathway to dissociate from the transcription start sites of HSR genes upon command form DNA-bound activated HSFs, there is strong evidence that the most upstream components are heat-sensitive membrane receptors consisting of the cyclic nucleotide-gated calcium channels (CNGCs) (Gao et al. 2012; Finka et al. 2012; Tunc-Ozdemir et al. 2012). A growing body of evidence has accumulated, pointing at a fluidity-based thermosensory mechanism in the plasma membrane of land plant cells (Saidi et al. 2011; Mittler et al. 2012), which can act as an effective early warning system during physiological warming, to trigger a timely buildup of HSP-based protections, in anticipation of upcoming damaging conditions. The plant plasma membrane apparently contains distinct populations of embedded thermosensors. Thus, a mild temperature increase can activate and depolarize one type of Ca$^{2+}$ channels, while the others still remain fully potent until exposed to higher temperatures (Saidi et al. 2005; Finka and Goloubinoff 2013). In confirmation, electrophysiology shows the presence of three distinct thermoresponsive Ca$^{2+}$ channels in the plasma membrane of moss protoplasts, with the distinct conductances of 75pS, 33pS, and 15pS (Finka et al. 2012).

A knockout of CNGC6 in Arabidopsis thaliana resulted in plants with vegetative tissues showing a decreased tolerance to heat stress (Gao et al. 2012). This appears analogous to the increased heat stress sensitivity that has been observed in the pollen of CNGC16 mutants (Tunc-Ozdemir et al. 2012). Among the land plants, the A. thaliana genome encodes for 20 closely related CNGC genes that clearly cluster with eight homologous CNGC genes in the genome of Physcomitrella patens and five in the genome of Selaginella moellendorfii. All have six conserved transmembrane helices, a putative cyclic nucleotide-binding
domain, a putative Ca\textsuperscript{2+}-calmodulin-binding domain, and a characteristic “phosphate-binding cassette” (Zelman et al. 2012; Wheeler and Brownlee 2008). Remarkably, no protein with similar enough sequences and reuniting similar putative domains can be found in the genome of liverwort Marchantia polymorpha and in algae (Zelman et al. 2012; Wheeler and Brownlee 2008; Finka et al. 2012), suggesting that the evolution of this class of cyclic nucleotide-gated Ca\textsuperscript{2+} channels in the plant kingdom is related to the adaptation of photosynthetic organisms to the harsh atmospheric conditions of the terrestrial environment.

Confirming the link between CNGCs and land plant heat sensing, a site-directed CNGCb knockout mutant in P. patens moss and the homozygote interruption mutants in the orthologous Arabidopsis genes AtCNGC2 and AtCNGC4 (see Supplementary Fig. S1) display dysregulated hyperthermosensitive responses and hyper-thermosensitive profile of acquired thermotolerance (Finka et al. 2012). Noticeably, electrophysiology shows that the CNGCb knockout mutant is devoid of the 75pS channel, whereas the remaining 33pS and 15pS Ca\textsuperscript{2+} channels have increased open probabilities, compared to the wild type (WT) moss membranes. Thus, while CNGCb is likely a major subunit component of the 75pS channel, it may be a minor component of other two channels and may be replaced by the other orthologous CNGCs, leading to the dysregulated thermoresponsive phenotypes of the 33pS and 15pS channels in the CNGCb mutant.

Here, we show that the P. patens CNGCd gene, which is a close ortholog of CNGCb, carries a similar thermosensory function. Whereas either the CNGCd or the CNGCb deletion caused a hyper-thermosensitive response at mild non-damaging temperatures, they did not affect the plant cell sensitivity to excessive damaging temperatures. In contrast, application of an artificial membrane fluidizer caused a hyper-thermosensitive response, both at mild and at excessively high temperatures, while it further exacerbated hyper-thermosensitive response in mutants implying a cumulative effect between increased membrane hyperfluidity and Ca\textsuperscript{2+} leakiness. Thus, the plant heat sensor acts as a proteolipid thermocouple (Horvath et al. 2012), responding to mild temperature-induced changes in the fluidity of the surrounding plasma membrane by sending a specific HSR to transcribe the HSP genes and thus establish acquired thermotolerance.

**Results**

The PpCNGCd loss-of-function moss mutant yields mildly chlorotic, but fertile gametophores

We used the reporter moss line Hsp-GUS (Saidi et al. 2005), referred here as the “wild type” strain (WT) and deleted most of the PpCNGCd gene by targeted gene replacement (Schaefer and Zryd 1997). Following antibiotic selection, ten stable transformants were identified and the genomic DNA was isolated. Southern blot analysis of one mutant, ΔCNGCd, confirmed that the CNGCd gene was disrupted by the stable integration of multiple copies of the selective marker (Fig. 1, lower left). In contrast to the growth-impaired ΔCNGCb mutant (Finka et al. 2012), the ΔCNGCd protonema grew at similar rates as the WT moss. Although the ΔCNGCd gametophores were slightly more chlorotic (Fig. 1, lower right), the transition to gametophores took place at about the same time as the ΔCNGCb and WT strains and ΔCNGCd apparently generated a similar number of capsules (Fig. 1, arrow heads) with viable spores.

CNGCd controls heat-induced Ca\textsuperscript{2+} entry and the expression of HSPs

Electrophysiology of WT and ΔCNGCb moss protoplasts has shown that mild temperature upshifts from 25 up to 32 °C

\[\text{Electrophysiology of WT and } \Delta \text{CNGCb moss protoplasts}\]

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![Image](https://example.com/fig1.png)  
**Fig. 1** The CNGCd loss-of-function moss mutant is mildly chlorotic but with fertile gametophores. Morphological appearance of 3-month-old moss gametophores of WT (upper left), ΔCNGCb (upper right), and ΔCNGCd (lower right) strains bearing spore capsules (arrows). *Inset:* schematic representation of the targeted genomic moss CNGCd locus in the WT strain and the CNGCd mutant (upper). The BglIII restriction sites (B, which are in the 5′ (white box) and the 3′ (black box) fragments of the CNGCd gene, are indicated. The southern blot from BglIII-digested genomic DNA from the WT CNGCd mutant strain probed with PCR-amplified segment corresponding to the 3′ (black arrow). The BglIII pattern shows that a 2,911-bp genomic fragment in the WT CNGCd gene has been mutated into a 3,520-bp-long 5′-35SptII-3′ fragment of the core region of the CNGCd locus, which became interrupted by a multiple copy insertion, creating also a single flanking region of about 3,100 bp (arrowheads).
caused the transient entry of apoplastic Ca\textsuperscript{2+} through three distinct temperature-sensitive Ca\textsuperscript{2+} channels in the moss cell plasma membrane (Saidi et al. 2009; Finka et al. 2012). Here, we used a less sensitive, but a noninvasive, whole tissue assay to measure temperature-dependent fluctuations of cytoplasmic Ca\textsuperscript{2+} in intact plant tissues. To this aim, we first introduced into the ΔCNGCd strain a stable UBI-AEQ cassette expressing an aequorin reporter from the strong constitutive ubiquitin promoter, which, in the presence of coelenterazine and Ca\textsuperscript{2+} ions, emits light thereby reporting fluctuations of cytoplasmic Ca\textsuperscript{2+} concentration (Plieth 2006; Saidi et al. 2009). Whereas a mild temperature upshift from 25 to 34 °C did not cause the entry of measurable amounts of apoplastic Ca\textsuperscript{2+} into the cytoplasm of the WT strain (Saidi et al. 2009; Finka et al. 2012), a significant Ca\textsuperscript{2+} signal was observed in the cytosol of both ΔCNGCb and ΔCNGCd strains (Fig. 2a). Confirming the hyper-thermoreponsive nature of both mutant strains, a stronger temperature upshift, from 25 to 36 °C, caused a massive Ca\textsuperscript{2+} entry in the cytoplasm of both ΔCNGCb and ΔCNGCd strains, and in contrast, a significant, yet much milder entry of apoplastic Ca\textsuperscript{2+} into the cytoplasm of the WT strain was observed.

These data confirm that both CNGCb and CNGCd are primary components of specific Ca\textsuperscript{2+} channels in the plasma membrane acting in plant cells as thermosensors. They apparently react to temperature-induced fluidity changes in the surrounding membranes, by allowing the transient entry of apoplastic Ca\textsuperscript{2+}, to trigger a specific HSR in the form of accumulating HSPs. Whereas the time range for thermally induced Ca\textsuperscript{2+} fluctuations is in the order of several minutes (Saidi et al. 2009; Finka et al. 2012), the time range for thermally induced HSP accumulation, leading to acquired thermotolerance is in the order of several hours. We therefore next addressed the effect of the gene deletions in the CNGCb and CNGCd thermosensors, on the heat induction profile of the moss HSPs, using the heat-inducible HSP-GUS expression cassette, as a general reporter for the conditional HSP expression in moss (Saidi et al. 2005, 2007, 2009; Finka et al. 2012).

Whereas in the HSP-GUS (WT) control strain, a 60-min abrupt temperature upshift from 25 to 32 °C did not induce any GUS, and half optimal GUS accumulation was observed at 34.8 °C, in the HSP-GUS ΔCNGCd strain, the same two thresholds were reached at 30 and 34 °C, respectively, and in the HSP-GUS CNGCb strain, at less than 28 and 31.5 °C, respectively (Fig. 2c). Thus, both CNGC mutants were hyper-thermoreponsive to a mild temperature rise, both in terms of Ca\textsuperscript{2+} entry within minutes and of HSP accumulation within hours. This indicates that heat-induced HSP expression at 36 °C is a direct consequence of a prior heat-induced entry of apoplastic Ca\textsuperscript{2+} through the heat-responsive CNGC channels in the plasma membrane.

The genome of P. patens encodes for six other putative CNGC genes, which are related to CNGCb and CNGCd (Finka et al. 2012). This raises the possibility that some of functional CNGC channels are heterocomplexes composed of more than two different subunits, in different ratios, each with a different threshold of thermo-responsiveness, and that when a given CNGC subunit is missing, it might be functionally but imperfectly replaced by one of its close orthologs.

![Fig. 2](image-url)
Artificial membrane fluidization renders moss hyper-thermoreponsive

When the WT Hsp-GUS strain was incubated with the membrane fluidizer benzyl alcohol (5 mM) and treated for 30 min at increasing temperatures (de Marco et al. 2005; Saidi et al. 2009; Horváth et al. 1998), this caused a general downshift of the temperature profile of Hsp-GUS accumulation, from ET50=34.8 °C without BA to 32.0 °C with BA (Fig. 3). Thus, in the low range of physiological warming temperatures in WT moss, the presence of the artificial membrane fluidizer recapitulated the hyper-thermoreponsive phenotype, which was observed without BA in the CNGCb or the CNGCd deletion mutants. Noticeably, the CNGCb/d deletion mutants and the WT strains showed the same degree of sensitivity to excessive temperatures, with IT50 values 42.1±0.1 °C (Fig. 3, Fig. S2). In contrast, the presence of BA downshifted the IT50 values of GUS overproduction from 42.1 to 41.3 °C in CNGCd, to 39.5 °C in CNGCb as compared to 39.1 °C in WT (Fig. S2B).

Discussion

As evidenced by their additive effects, the hyper-thermoreponsive phenotype that was caused by the BA treatment originated from an artificial change in the membrane fluidity, as compared to the CNGC knockout mutations that did not change membrane fluidity but rather changes the quality of the CNGC protein complexes that sense the fluidity state of surrounding plasma membrane. This corroborated previous observations in fully differentiated aging moss gametophores of the HSP-GUS strain, where the heat-induced GUS expression was dramatically reduced compared to young protonema tissues, in correlation with a significantly higher content of unsaturated lipids in the young membranes (Saidi et al. 2010). Together, our data strongly suggest that both the lipid part of the plasma membrane and the embedded CNGCb and CNGCd proteins act as two cooperating moieties of an effective thermocouple.

Our data suggest a mechanism where mild changes in the ambient temperature cause direct fluidity changes in the plasma membrane of the plant cells, which are sensed by the embedded CNGC channels, which consequently open. This allows the transient entry of apoplastic Ca2+ ions through the channels into the cytoplasm, to interact with the cytosolic C-terminal domains of the CNGCs that contain calmodulin-binding signatures (Fischer et al. 2013; Arazi et al. 2000). Thus, the heat shock signal may propagate from the plasma membrane, likely by way of specific Ca2+-dependent calmodulins and calmodulin kinases and phospholipases (Mishkind et al. 2009; Liu et al. 2008, 2006; reviewed in Saidi et al. 2011; Mittler et al. 2012), to activate the HSFs in the cytosol and cause HSF translocation to the nucleus to specifically transcribe protective heat shock proteins, leading to acquired thermostolerance.

It has been suggested that particular histones, such as H2A.Z, which are frequently but not exclusively found associated to HSP genes, may directly sense an increase of ambient temperature (from 11 to 17 °C), and like a molecular thermometer directly react to the temperature by dissociating from their repressory association with HSP genes, thereby allowing the active transcription of HSP genes, in particular Hsp70 (Kumar and Wigge 2010). Our finding that specific mutations in the CNGC proteins in the plasma membrane strongly affect the profile of plant cell response to temperature increments (Finka et al. 2012) and that the artificial prevention of apoplastic Ca2+ entry into the cytoplasm during the first minutes of heat shock totally prevent the HSR (Saidi et al. 2009) strongly argues against histones and histone-remodeling complexes being the primary thermosensors of the plant cells. Rather, our findings suggest that upon transiently increasing the membrane fluidity, higher temperatures trigger the transient opening of various populations of channels composed of CNGCb and CNGCd subunits, which in turn translate into a specific Ca2+-entry-dependent and calmodulin-dependent heat shock signal that likely activates a calmodulin-dependent kinase (Liu et al. 2008), specifically phosphorylating and thus activating the HSFs in the cytoplasm. Once hyper-phosphorylated, the activated HSFs may enter the nucleus and bind specifically only to the HSEs.

Interestingly, mammalian cells also carry plasma membrane-embedded Ca2+ channels acting similarly as cellular thermosensors responsible to the cellular HSR. Thus, the transient receptor potential vanilloids (TRPVs) are ion channels in the plasma membrane of various neural and

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![Fig. 3](image-url) The membrane fluidizer benzyl alcohol (BA) produces a hyper-thermosensitive response in Hsp-GUS WT. Heat induced GUS accumulation in the Hsp-GUS WT strain following 30-min exposure at the indicated temperatures and 16 h at 25 °C, without (plain lines) or in the presence of the medium of 5 mM benzyl alcohol (BA, dashed lines). Vertical dash-line and plain arrows show ET50 values and IT50 for BA-treated and non-treated moss, respectively.
epithelial cells that are distantly related to the plant CNGCs, both types being Ca$^{2+}$ channels composed of six transmembrane helices with a cytosolic calmodulin-binding domain that can apparently respond to fluidity changes, or to mild physiological warming, by sending a specific Ca$^{2+}$ entry-dependent signal to activate cytosolic HSF1. Upon warming, or the isothermal application of the TRPV agonist capsaicin, the TRPV-based thermosensory mechanism activates a typical HSR in the form of HSP accumulation. Moreover, the heat/fever-induced HSR is inhibited by TRPV1 siRNA, as well as by the TRPV1 antagonist capsazepine, or more generally by the external application of a Ca$^{2+}$ chelator (Bromberg et al. 2013).

It is traditionally thought that the primary heat sensors in eukaryotic cells, including in plants, were some unidentified thermolabile cytosolic proteins that presumably denature under a noxious heat shock, thereby recruiting the cytosolic Hsp90 and Hsp70 and derepressing the inactive HSFs form their inhibitory association with these chaperones (Voellmy and Boellmann 2007). Yet, this mechanism is counterintuitive as it implies a dilatory induction of the cellular defenses, only once damages in proteins have occurred. In contrast, we bring here strong evidence pointing at a fluidity-based thermosensory mechanism in the plant plasma membrane of moss plant that can act as an early warning system under a physiological warming, to trigger the timely buildup of HSP-based cellular protections, in anticipation of upcoming heat damaging conditions.

### Material and methods

#### Plasmids

**PpCNGCd-KO vector**

The 5′ targeting fragment of PpCNGCd was amplified using ApaI (GGGCGGCTTTCGGAAGATATCCATCTTCTC) and XhoI (CTCGAGCACACAAACAACAAATTAGAAATCTC) and 3′ targeting fragment was amplified by SpeI (ACTAGTGTGGGGCTGCTAGGGATCGA) and SacII (CCGGAGAATTGTGTGCTGG AAGTGTGGCGT) using Taq polymerase (Promega, Madison, WI). Resulting products were first subcloned in pGEM-T-easy vector (Promega Madison, WI) and digested with Apal/Xho or SpeI/SacII and subcloned in two steps into corresponding sites of pBSMDII (Finka et al. 2008, 2012), to generate the PpCNGCd-KO vector. To minimize the possibility of super integration of CNGCb-KO cassette into HSP-GUS locus, about 10 μg of CNGCd-KO transgene was amplified by PCR employing standard T3 and T7 primers, followed by enzymatic digestion with Apal and SacII restriction endonucleases.

Plant material, moss transformation, chemicals, GUS, and aequorin experimentations

The moss *P. patens* transgenic lines were grown on moss solid medium and transferred when stated to liquid minimal medium as previously described (Saidi et al. 2009; Finka et al. 2012). To generate stable transgenic moss *CNGCd* lines in the HSP-GUS background, a polyethylene glycol-mediated transformation of moss protoplasts was performed using PCR-amplified *CNGCb*-KO cassette followed by G418 antibiotic selection (Schaefer and Zryd 1997). The aequorin-expressing *CNGCd* mutant was created by introduction of pBUAzeo (Finka et al. 2012) into the previously generated *CNGCd* mutant line by super integration of the pBUAzeo into the artificial Hsp-GUS locus.

The coelenterazine hcp, EGTA, and 4-methylumbelliferone glucoronide were purchased from Sigma (St Louis, MO, USA). GUS-specific activities, electrophysiological experimentations, and the concentration of cytosolic calcium using reconstituted aequorin system were described previously (Saidi et al. 2009; Finka et al. 2012).

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