Expression analysis of calmodulin and calmodulin-like genes from rice, Oryza sativa L.

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Abstract

Background: In plants, a large family of calmodulin (CaM) and CaM-like (CML) proteins transduce the increase in cytosolic Ca\(^{2+}\) concentrations by binding to and altering the activities of target proteins, and thereby affecting the physiological responses to a vast array of stimuli. Here, transcript expression analysis of Cam and CML gene family members in rice (Oryza sativa L.) was extensively examined.

Results: Cam and CML genes in rice exhibited differential expression patterns in tissues/organs. Under osmotic stress and salt stress, expression of OsCam1-1, OsCML4, 5, 8, and 11 was induced with different kinetics and magnitude. OsCML4 and 8 mRNA levels significantly increased by 3 h after treatment and remained elevated for at least 24 h while expression of OsCam1-1, OsCML5 and 11 was up-regulated as early as 1–3 h before rapidly returning to normal levels. Several cis-acting elements in response to abiotic stresses, including DREs (important promoter elements responsive to drought, high salt, and cold stress), were detected in the 5' upstream regions of these genes. The observed induction of the GUS activity of transgenic rice plants via the OsCam1-1 promoter appeared to be biphasic and dependent on the severity of salt stress.

Conclusions: Large OsCam and OsCML gene family members likely play differential roles as signal transducers in regulating various developmental processes and represent important nodes in the signal transduction and transcriptional regulation networks in abiotic stress responses mediated by the complex Ca\(^{2+}\) signals in plants, which are rich in both spatial and temporal information.

Keywords: Calcium signaling, Calmodulin, CaM, CML, Rice, Oryza sativa

Background

Transient changes in the cytosolic Ca\(^{2+}\) concentration ([Ca\(^{2+}\)\(_{\text{cyt}}\)] of a different magnitude and specialized character due to the activities of Ca\(^{2+}\)-ATPases and Ca\(^{2+}\)-channels in the cellular membrane of eukaryotes are utilized as a second messenger in generating physiological responses to extracellular stimuli. The use of Ca\(^{2+}\) signals has been implicated in generating responses to a wide variety of environmental changes in plants [1]. The changes in [Ca\(^{2+}\)\(_{\text{cyt}}\]) are not only transient, but also vary spatially and temporally with different organelles acting as distinct compartments [2], therefore, the diverse array of different changes in the [Ca\(^{2+}\)\(_{\text{cyt}}\]) must be discriminated so as to elicit the correct subsequent cellular response, a task performed by various Ca\(^{2+}\)-modulated proteins. For the majority of these proteins, the Ca\(^{2+}\)-binding sites are composed of a characteristic helix-loop-helix motif called an EF-hand [3], which binds Ca\(^{2+}\) with high affinity, resulting in conformation changes that modulate their activity or their ability to interact with other proteins.

Plants possess a complex network of Ca\(^{2+}\) signal transduction in mediating responses to various biotic and abiotic environmental stimuli. In plants, genes encoding EF-hand-containing proteins have been extensively annotated in Arabidopsis thaliana (L.) Heynh [4] and rice (Oryza sativa L.) [5]. Three groups of EF-hand-containing Ca\(^{2+}\) sensor proteins have been identified, which include Ca\(^{2+}\)-dependent protein kinase (CPK), calcineurin B-like protein (CBL), and calmodulin (CaM) [6]. These proteins can be considered as sensor responders or sensor relays [7], CPKs, and CBLs together with their respective CBL-interacting protein kinases (CIPKs), are sensor responders
as they possess kinase activity either in their molecules or in their high affinity interacting partners as a responder function. By contrast, CaMs and CMLs, which have no other identifiable functional domain other than EF-hand motifs, are considered sensor relays. A large family of Cam and Cam-like (CML) genes has been extensively identified from the two model plants, A. thaliana [8] and O. sativa [5]. Although the existence of these proteins in a single plant species is believed to be important to correctly perceive and discriminate the Ca$^{2+}$ signals from different stimuli, and thus aid in eliciting the correct subsequent response, the molecular mechanisms and physiological significance of most of these proteins have not been established. Nevertheless, accumulating evidence suggests that each of the different Cam and CML genes may have distinct and significant functions [9].

Currently, the information on Cam and CML genes in O. sativa is quite limited despite the fact it is a model plant for monocot and especially cereal plants. Regarding expression studies, reports on only a few members of the OsCam and OsCML gene family, are available [10,11]. Here, the transcript expression levels of several members of the OsCam and OsCML gene family were examined in O. sativa that had experienced abiotic stresses. Their 5’ upstream sequences were examined to identify putative stress-responsive cis-acting elements. The putative promoter of the OsCam1-1 gene was further examined using three transgenic rice lines. Based on their expression patterns, the OsCam and OsCML genes that are likely to represent important nodes in the signal transduction and transcriptional regulation networks of Ca$^{2+}$ signaling in response to different abiotic stresses will be revealed, and subject to confirmation, this is likely facilitate the efforts in characterizing the mechanisms of abiotic stress Ca$^{2+}$ signalling in plants.

**Results**

**Differential expression of OsCam and OsCML gene family members in various organs**

The rice (Oryza sativa L.) genome has been reported to encode for a large family of five Cam genes and 32 genes encoding CML calcium sensor proteins [5]. To determine which of these OsCam and OsCML genes may have tissue- and/or stress-specific expression patterns, which could suggest their functions, their expression levels in different rice tissues and organs were investigated using publicly available microarray data of global gene expression from the experiment GSE 6893 from the O.sativa IR64 cultivar [12]. The available expression data of all OsCam and OsCML genes plus OsEF1a were retrieved and the results are shown as heat map in Figure 1A, and are grouped according to the previously reported clades found by phylogenetic analysis of their amino acid sequences. It is clearly noticeable that all the OsCam genes (group 1) are highly expressed in almost all of the organs examined. In contrast, several highly conserved OsCML (groups 2–5) genes tend to be highly expressed in some organs/tissues. Interestingly, one member of each subgroup in group 6 is found expressed at a higher level compared with the other members in their subgroup (OsCML30 in group 6a; OsCML27 in group 6b; OsCML29 in group 6c; OsCML31 in group 6e) except group 6C, in which all members are highly expressed in most organs examined.

Transcript expression levels of ten genes, one Cam (OsCam1-1) and nine highly conserved CML genes (OsCML1, 3, 4 and 5 in group 2, OsCML8 and 13 in group 3, OsCML7 and 9 in group 4, and OsCML11 in group 5) were closely examined in different rice organs and tissues (Figure 1B). Our preliminary results indicated that OsCam1-1 gene is the only OsCam gene which expression is induced by osmotic stress and salt stress, therefore, only OsCam1-1 among the five OsCam genes was selected for this study. Based on the microarray data analysis, ubiquitous transcript expression at relatively high levels of OsCam1-1, OsCML7 and 13 was observed. The other three genes, OsCML1, 4, and 8 had similar expression patterns with higher levels in the leaf and the root and lower levels in the young inflorescence (Figure 1A). Their expression levels were then increasing upon maturation of the inflorescence until reaching similar levels to those of the leaf and the root in the early developing seed. Expression levels of OsCML4, and 8 were maintained in the seed until dormancy and desiccation while that of OsCML1 later decreased upon seed maturation (Figure 1B). For the other four genes: OsCML3, 5, 9, and 11, overall, they exhibited relatively lower expression levels compared to the aforementioned genes.

To examine whether these genes are expressed in the three-week old Khao Dawk Mali 105 (KDML105) rice cultivar seedlings and to optimize the PCR conditions, RT-PCR was performed using the oligonucleotide primers specific for each gene (Table 1). Following separation of the PCR products in agarose-TBE gels and visualization by uv-transillumination after ethidium-bromide staining, bands of the expected size based on the OsCam1-1 and OsCML sequences (Table 1) were specifically detected (data not shown). Thus, transcripts of OsCam1-1 and all nine OsCML genes examined are expressed in leaves of the three-week old ‘KDML105’ rice seedlings, and can be specifically amplified by RT-PCR under these conditions.

The transcript expression levels of these genes in the leaf blades/sheaths of the three-week old ‘KDML105’ rice seedlings, was evaluated by rt-RT-PCR using the transcript expression level of OsCML5 as reference (Figure 2), OsCam1-1 was found to have the highest expression level (~150-fold higher than OsCML5) in the leaf tissue.
of the three-week old rice seedlings, followed by OsCML4, 8 (~120-fold) (Figure 2A). OsCML1, 7, and 13 also exhibited a relatively high expression level (>30-fold), while OsCML3, 5, 9, and 11 expressed at a relatively lower level.

The transcript expression levels of OsCam1-1 and all nine OsCML genes assayed in the leaf blades/sheaths, roots, flowers, and seeds of the KDML105 rice cultivar during the grain filling period were determined and compared with their respective levels in the leaf tissue of the three-week old rice seedlings (Figure 2B). The expression profiles of these genes may be divided into three groups when only large differences (≥5-fold) among organs were considered. The first group, which has the highest expression levels in the root includes OsCML4, and 5. The second group with the highest expression levels in the flower and the seed were OsCML3, and 11. Finally, OsCam1-1 and the five other OsCML genes (OsCML1, 7, 8, 9, and 13) had more or less similar expression levels among these organs and comprised the third group.

Expression of OsCam and OsCML genes under abiotic stress
To investigate whether OsCam1-1 and the nine selected OsCML genes may possibly be involved in mediating responses to abiotic stress, the publicly available rice microarray data was retrieved and examined to acquire data for the IR64 rice cultivar, whilst the transcript and expression levels in the ‘KDML105’ rice were evaluated after different abiotic stresses by rt-RT-PCR.

The results of the DNA microarray data of OsCam1-1 and the nine OsCML genes in the 7-day-old ‘IR64’ rice seedlings grown under 3 h of dehydration (drought) or salt stress (200 mM NaCl) [12] retrieved via the rice oligonucleotide array database are shown as a heat map in Figure 3. The transcript expression levels of OsCam1-1, and especially of OsCML4, 5, 8, and 11 were significantly increased under dehydration, whilst that of OsCML13 was slightly down-regulated. Under salt stress, significantly increased transcript expression levels of OsCam1-1, and especially of OsCML4, 8, 9, and 11 were observed, whilst that for OsCML13 was slightly
Table 1 Oligonucleotide primers used in this study

| Primer name       | Sequence               | Position* | Amplicon size (bp)b | Annealing temp. (°C) |
|------------------|------------------------|-----------|--------------------|---------------------|
| OsCam1-1-F       | 5'-ACCGTGCAATTGCGTAGT-3' | 499-518   | 177                | 58.3                |
| OsCam1-1-R       | 5'-GAAGCTTACAGATCATCC-3' | 656-675   |                    |                     |
| OsCML1-F         | 5'-CCGAAAGTGCGTAGTCCGT-3' | 543-562   | 184                | 58.3                |
| OsCML1-R         | 5'-ACTACGGAACCGCTGTGA-3' | 707-726   |                    |                     |
| OsCML3-F         | 5'-TACACAGCTTCCCAAG-3'  | 410-429   | 190                | 57.3                |
| OsCML3-R         | 5'-CATCGGAGAGCTAAGCAAC-3' | 440-460   |                    |                     |
| OsCML4-F         | 5'-GCAGGCTGACTACGATGAA-3' | 402-421   | 193                | 56.3                |
| OsCML4-R         | 5'-TACCATAGCTGAACTCCCA-3' | 575-594   |                    |                     |
| OsCML5-F         | 5'-ATGATGCTCTCCGACCAATA-3' | 481-500   | 180                | 57.3                |
| OsCML5-R         | 5'-CAGAGGCCAATTAATGAC-3' | 641-660   |                    |                     |
| OsCML7-F         | 5'-TACAGGCTTGCGCCTAAAT-3' | 429-448   | 193                | 57.3                |
| OsCML7-R         | 5'-GTTCAATACACACCAGGAT-3' | 602-621   |                    |                     |
| OsCML8-F         | 5'-AGATGATGAGAGGATAGGA-3' | 539-555   | 185                | 56.3                |
| OsCML8-R         | 5'-AAAATGAAGGGTGATGATA-3' | 701-720   |                    |                     |
| OsCML9-F         | 5'-TACAGGCTTGCTCAAGG-3'  | 430-449   | 170                | 58.3                |
| OsCML9-R         | 5'-GATTGCTTGCAATCTACGCG-3' | 579-600   |                    |                     |
| OsCML11-F        | 5'-CAATATCTTCCTTCTGAAAT-3' | 621-640   | 183                | 56.3                |
| OsCML11-R        | 5'-ATTCCACAGGCTCAGC-3'  | 784-803   |                    |                     |
| OsCML13-F        | 5'-ATGGAATATGTTGATGTTGAG-3' | 437-456   | 193                | 58.3                |
| OsCML13-R        | 5'-GCTGTTGCTTCTGTTCTGCA-3' | 610-629   |                    |                     |
| OxF10a-F         | 5'-ATGGATGGAGGACGCTCTC-3' | 1192-1209 | 127                | 58.3                |
| OxF10b-R         | 5'-TCACCTTGACGCGACGTTG-3' | 1301-1318 |                    |                     |

*Position of the primers from the GenBank sequence given in the materials and methods, where position 1 is the predicted open reading frame start codon and numbered 5’ to 3’ on the sense strand.

bExpected amplicon size based upon the primer positions on the GenBank sequence.

down-regulated. Note that although numerically the transcript expression level of OsCML3 was increased following both dehydration and salt stress, this was not statistically significant due to the large variance in the control samples, and so remains to be clarified.

With respect to the three-week-old O.sativa ‘KDML105’ seedlings, rt-RT-PCR revealed that under osmotic stress (20% PEG), OsCML4 and OsCML8 exhibited a nine- and four-fold increases in their transcript expression levels, respectively, and that this started from 3 h after osmotic stress and were maintained at these elevated expression levels until at least 48 h after treatment (Figure 3B). However, whilst significantly elevated transcript expression levels of OsCML11 were also detected at 3 h after osmotic stress (13-fold), in contrast these expression levels then declined to six-fold higher and control levels at 6 h and 12 h onwards after osmotic stress, respectively. Expression of OsCam1-1, and OsCML5 showed an earlier response, being up-regulated 5-6-fold at 1 h after osmotic stress treatment compared with the control, but whilst OsCam1-1 expression levels remained slightly up-regulated at 3 to 48 h after osmotic stress that for OsCML5 returned to their normal levels after 3 to 6 h. OsCML7 and OsCML9 exhibited a slight increase in their transcript expression level at early time points (1–3 and 3–6 h after treatment, respectively), while no significant changes in the transcript expression levels of OsCML1, 3 and 13 were observed at all time points examined.

Similar to osmotic stress, OsCML4 and OsCML8 also showed significant prolonged increases in their transcript expression levels under salt stress (150 mM NaCl), starting from 3 h until at least 24 h after treatment (Figure 3C). Compared to the respective control, the OsCML4 transcript expression levels were maximal at a four-fold elevated level at 6 and 12 h after treatment while OsCML8 reached the highest levels (12-fold) at 12 and 24 h after treatment. Early up-regulated transcript expression of OsCam1-1 and OsCML5 at 1 h after treatment (five- and eight-fold, respectively) and of OsCML11 at 3 h after treatment (six-fold) was also observed. In contrast, the transcript expression levels of OsCML3 and OsCML13 were not significantly altered in response to salt stress at all time points examined in this study, whilst OsCML7 and OsCML9 were only slightly and transiently increased at 1 h and 6 to 12 h after salt stress, respectively. However, unlike under osmotic stress,
OsCML1 exhibited increase in its expression at 1 h after salt stress.

**Examination of the OsCam1-1 and OsCML promoters**

In order to examine the promoters of these genes and their cis-acting elements, the sequences upstream of their coding regions were retrieved from GenBank and bioinformatically analyzed using the PLACE [13,14] and the PlantPAN [15] software and databases. Several characteristic elements were located; including motifs involved in responses to dehydration, abscisic acid (ABA), salt stress, and cold stress (Figure 4). Putative DREs (drought-responsive elements) [16] were found in the 5' flanking (upstream) region of five of the loci (OsCam1-1 at -1064, OsCML4 at -892, OsCML5 at -954, and OsCML11 at -561 and -565). In addition, the 5’ upstream sequences from eight of these genes (OsCam1-1, OsCML1, 3, 4, 5, 7, 8 and 13) contained putative ABREs (ABA-responsive elements) [17], while LTREs (low temperature responsive elements) [18] were found in seven of these genes (OsCam1-1, OsCML3, 4, 5, 7, and 11).

To investigate inducibility of the OsCam1-1 promoter by salt-stress, (0 (control), 100, 150 or 300 mM NaCl), the β-glucuronidase (GUS) activity of three independent lines of the homozygous T3 transgenic KDML105 rice cultivars harbouring the OsCam1-1::gus construct was examined fluorometrically in two-week-old seedlings. All three independent transgenic lines exhibited similar trends of GUS activity.
Figure 3 (See legend on next page.)
activity to each other under the different NaCl treatments (Figure 5). The plants treated with 100 mM NaCl did not exhibit any statistically significant difference in GUS activity in all three independent lines (p < 0.05) compared to their respective controls at 0 and the untreated plants, whereas at 150 mM NaCl all transgenic lines had exhibited a significantly higher GUS activity from 4 h after treatment in lines 1 and 3, and after 1 to 8 h in line 2. For the plants treated with 300 mM NaCl, a biphasic induction of GUS activity was observed in all three independent transgenic lines with an initial induction and highest induction of GUS activity at 1 h after treatment, which then decreased to the control level by 2 h and was followed by the second phase of induction at 4 h after treatment. This second phase was either transient (line 2) or maintained for up to 8 h (line 1) or 24 h (line 3) after treatment.

Discussion

Whilst the OsCaM group contains five members that share a highest degree of amino acid sequence identity (≥ 97%) to known typical CaMs from other plants, previous report phylogenetically classified the highly conserved OsCML proteins into four groups [5]. The OsCML proteins are small proteins of 145 to 250 amino acid residues with an approximately 44% to 85% amino acid identity to typical plant CaMs. A large family of six Cam and 50 CML genes have also been annotated in the A. thaliana genome [8], suggesting the existence of an extensive set of CaM and CML proteins in each plant species.

Even though rice is considered an important crop and a model for other monocots and especially cereal crops, almost all of OsCam and OsCML genes have not been characterized. From the publicly available microarray data, all OsCam genes were found highly expressed in almost all organs examined (Figure 1A). One of the defining characteristics of CaMs in plants is the presence of multiple CaM isoforms. Even though they are ubiquitously expressed, the different isoforms can display differential responses to individual stimuli in any given plant tissue [5,11,19-22], suggesting that each of the OsCam genes may have distinct physiological functions depending on where and how the expression of each gene is up-regulated in addition to the different biochemical properties that might be affected by the slight differences in their primary structures [21,23].

In contrast, several highly conserved OsCML genes (groups 2–5) genes tend to be highly expressed in some

**Figure 3** OsCam and OsCML transcript expression in response to stress. (A) Microarray based transcript expression profiles under dehydration (drought) and salt stress shown as a heat map. Transcript expression levels examined in the KDML105 rice cultivar under osmotic stress (B) and salt stress (C) showing relative transcript expression levels standardized to that of OsEF1α and expressed relative to the levels on day 0 of the treatment. Rice seedlings were grown in a CRD with three replicates, and for each replicate, ten seedlings were pooled for RNA extraction. The PCR reaction of the same cDNA preparation was performed in triplicate for technical replication. Data are shown as the mean ± 1 SD and means with a different lowercase letter are significantly different (p < 0.05).

**Figure 4** Cis-acting elements involved in response to abiotic stress in the OsCam1-1 and OsCML promoters. The 5’ upstream regions are represented by lines and the putative ABRE, DRE and LTRE sequences are shown with a striped rectangle, a close rectangle, and an open rectangle, respectively.
organs/tissues and their levels of expression are modulated during different stages of development (Figure 1A), indicating that they are developmentally regulated. It is conceivable that CaM proteins have many targets and are ubiquitously involved in numerous cellular processes while the highly conserved CML proteins have more specialized targets and are involved in more specific processes of the cell. Interestingly, each subgroup of the highly diverged group of OsCML (group 6a to 6e) genes had one member whose expression is found at a noticeably higher level than the other members. Members in the same subgroups have the same number and configuration of EF-hand motifs. It is speculated that these highly expressed genes may be the predominant gene representing each subgroup that has essential functions in the cell.

When the microarray data was closely examined for the transcript expression levels of the OsCam1-1 and nine OsCML genes (Figure 1B), ubiquitous expression at relatively high levels of OsCam1-1, OsCML7, and I3 was observed, which suggests that they may have important functions during the regular growth and development processes of rice plants. The increasing transcript expression levels of OsCML1, 4, and 8 with maturation of the inflorescence until they reached similar levels as those in the leaf and root tissues in the early developing seed, suggest that they may have important roles during the maturation of the inflorescence and the early seed development, especially OsCML1, which expression level peaked during those stages.

The transcript expression levels in the ‘KDML105’ rice in the leaf blades/sheaths of the three-week old ‘KDML105’ rice seedlings, as evaluated by rt-RT-PCR (Figure 2A) agree well with the microarray data in the IR64 cultivar (Figure 1A). When the transcript expression levels of OsCam1-1 and OsCMLs were examined in different organs (Figure 2B), their profiles could be divided into three groups as those with the highest expression level in (i) the root, or (ii) in the flower and the seed, or (iii) those genes with similar expression levels among the organs examined. Relatively high transcript levels of OsCML3 and OsCML11 in the flower and the seed, and OsCML4, and 5 in the root suggest their functional significance in those respective organs. The OsCam1-1 and the other five OsCML genes (OsCML1, 7, 8, 9 and 13) had more or less similar expression levels among the different organs examined. From the microarray data of the IR64 rice cultivar, four of these genes (OsCam1-1, OsCML7, 9 and I3), were expressed at more or less constant levels in different developmental stages. However, expression patterns among different organs/tissues that are not consistent with those analyzed from the microarray database may also result from variation in the rice varieties examined.
Up-regulated expression of a gene in response to a stress signal may reflect the function of the corresponding gene product, especially in signal cascades. In a large gene family, investigating expression patterns of their members could point to genes or isoforms that potentially function under the conditions of interest. In this study, the transcript expression analysis of OsCam1-1 and nine OsCML genes in the KDML105 rice cultivar by rt-RT-PCR revealed that expression levels of OsCam1-1, OsCML4, 5, 8, and 11 were increased under osmotic stress (20% (w/v) PEG) (Figure 3B), which is consistent with the GSE6901 data for the IR64 cultivar from the DNA microarray database (Figure 3A) [12] and suggests that these genes may function in the mechanisms of Ca\(^{2+}\)-mediated responses to osmotic stress. Similarly, expression of these genes was also found to increase in the KDML105 rice cultivar under salt stress (150 mM NaCl) along with OsCML1 (Figure 3C), suggesting that OsCam1-1, OsCML1, 4, 5, 8, and 11 may function in the mechanisms of Ca\(^{2+}\)-mediated responses to salt stress. The up-regulated transcript expression levels under salt stress of OsCML4, 8 and 11 were also consistent with the GSE6901 data for the IR64 cultivar from the DNA microarray database (Figure 3A). Whether the discrepancy in the transcript expression patterns of OsCam1-1, OsCML1, and 5 is due to the different timings of expression being monitored or differences in the rice varieties examined remains to be evaluated but, overall, up-regulation of almost the same set of the highly conserved OsCML genes under osmotic and salt stresses was observed, confirming that conclusion that genes responsive to osmotic stress overlap to a high extent with those that are responsive to salt stress [24].

Several of these genes not only exhibited up-regulated transcript expression levels by osmotic stress and salt stress, but also exhibited different patterns of up-regulation in terms of timing and levels of expression within a tissue type and exhibited differential expression in different tissues/organs. Together, differential temporal and spatial expression patterns of these OsCam1-1 and OsCML genes suggest that each individual gene product may possess specific roles during Ca\(^{2+}\)-mediated responses to osmotic stress and salt stress. Comparison of the transcript expression patterns of these genes under osmotic or salt stress in the KDML105 rice cultivar, as determined by rt-RT-PCR in this study, with the publicly available RNA-Seq data and the other microarray data sets [25], revealed that OsCML4 and OsCML8 consistently exhibited higher expression levels under osmotic and salt stresses suggesting their significant functions in Ca\(^{2+}\)-mediated responses to these stimuli. In addition, the early induction within 1 h of OsCam1-1 and OsCML5 under osmotic and salt stresses, and of OsCML1 under salt stress suggests their importance in conveying the stress signals early in the transduction cascades of Ca\(^{2+}\) signaling. However, caution must be taken in interpreting these results because the changes in levels of transcripts generally do not coincide with the changes in the levels of the proteins they encode and, given the time it takes to express a protein, may not necessarily reflect their involvement in the responses of the cell to stress. Nonetheless, a gene of which transcript expression is early induced would be given special attention as a candidate for further investigation into its possible involvement in response to a particular stress.

Several cis-acting elements in response to abiotic stresses in the 5' upstream regions of the OsCam1-1 and OsCML genes were detected (Figure 4). DREs, important promoter elements that are responsive to drought, high salt, and cold [16], were located in the putative promoters of OsCam1-1, OsCML4, 5, 7, and 11. Consistent with this is that the transcript expression level of all these genes was shown by rt-RT-PCR to increase in the KDML105 rice cultivar under osmotic and salt stresses with OsCML7 exhibiting a slight increase in its expression level and the others significantly much higher level of up-regulation. DREs specifically interact with the transcription factors DREBs and regulate expression of many stress-inducible genes. In rice, the DRE binding protein 1 (OsDREB1) functions in the cold stress response, whereas OsDREB2 functions in the heat and osmotic stress responses [26]. The results here indicate that these DREs are potentially responsible for the osmotic stress-induced expression of these genes and interesting candidates for further characterization.

In the 5' upstream region of the OsCam1-1 promoter, the putative DRE motif (ACCGAC) was located at −1062. Induction of the GUS activity level in the three independent OsCam1-gus transgenic rice plants (Figure 5) suggested that induction of OsCam1-1 expression under salt stress is, at least, partly due to the activity of its promoter. Its induction appears to be dependent on the severity of the salt stress (concentration of NaCl), with no significant induction in the plants treated with 100 mM NaCl while a monophasic and biphasic induction were observed in the plants treated with 150 mM and 300 mM NaCl, respectively, in all three independent transgenic lines. A biphasic induction of OsCam1-1 expression by salt stress in the KDML105 rice cultivar, as determined by rt-RT-PCR has been reported previously [27], whilst heat shock (HS) induced biphasic [Ca\(^{2+}\)]\(_{cyt}\) signal in rice root cells and the HS-induced expression of OsCam1-1 strongly oscillated [11]. The complex responses of the OsCam1-1 gene to salt stress suggest that OsCam1-1 is a significant player in the Ca\(^{2+}\) signal transduction network under salt stress. However, caution must be taken in interpreting these results, especially on timings of the induction because of the possible
differences in mRNA and protein stability between the *gus* gene and the *OsCam1-1* gene.

**Conclusions**
Differential expression patterns in tissues/organs were reported among different *Cam* and *CML* genes in rice. Under osmotic and salt stresses, expression of several genes including *OsCam1-1*, *OsCML4*, 5, 8, and 11 was induced with different kinetics and magnitude. In agreement with their inducibility, several cis-acting elements in response to abiotic stresses including DREs, important promoter elements responsive to drought, high salt, and cold, were detected in the 5′ upstream regions of these genes. Induction in the GUS activity of the transgenic rice expressing *gus* gene under the control of the *OsCam1-1* promoter, which contains a putative DRE motif (ACCGAC) at −1062 was observed. Its induction appeared to be biphasic and dependent on the severity of salt stress. These results suggest that these *OsCam* and *OsCML* genes play differential roles as sensor relays in regulating developmental processes and Ca2+-mediated responses to abiotic stress.

**Methods**

**Materials**
TRI REAGENT was purchased from Molecular Research Center, Inc. (Cincinnati, OH, USA). The iScript cDNA synthesis kit and the SsoFast EvaGreen Supermix were purchased from Bio-Rad (Hercules, CA, USA). 4-methylumbelliferyl β-D-glucuronide (4-MUG) and 4-methylumbelliferone (4-MU) were from Sigma (St. Louis, MO, USA). Synthetic oligonucleotides for real-time reverse transcription polymerase chain reaction (rt-RT-PCR) were obtained from 1st Base (Singapore). Seeds of *Oryza sativa* L. cultivar Khao Dawk Mali 105 (KDMML105) were provided by the Rice Research Center (Pathumthani, Thailand).

**Plant growing and stress treatments**

*O. sativa* 'KDMML105' seedlings were hydroponically grown in nutrient solution under a 12-h light/12-h dark photoperiod. After three weeks, the plants were treated with 150 mM NaCl for 1, 3, 6, 12, 24 or 48 h by adding the same medium except supplemented with 150 mM NaCl or 20% (w/v) PEG4000 into the chamber containing the rice seedlings with minimal touch contact and disturbance. Rice seedlings were grown with a completely randomized design (CRD) with three replicates, and for each replicate, ten seedlings were pooled for RNA extraction.

**Real-time reverse transcription polymerase chain reaction (rt-RT-PCR)**

Leaf blades/sheaths of three-week old seedlings; or leaf blades/sheaths, roots, flowers and seeds during the grain filling period were collected and immediately frozen in liquid nitrogen and stored at −80°C. Rice tissues were ground in liquid nitrogen to a fine powder using a chilled mortar and pestle. Total RNA from the frozen tissues was extracted using the TRI REAGENT with 0.2 mL chloroform and precipitated by mixing with isopropanol. Subsequently, cDNA was synthesized by RT-PCR using the iScript cDNA synthesis kit according to the manufacturer’s instruction. Then, rt-PCR was performed in a final volume of 20 μL, which contained a 2-μL aliquot of the first strand cDNA reaction, 0.05 mM of each of the gene-specific primers (Table 1), and 1x SsoFast EvaGreen Supermix. The reaction included an initial 8 min at 95°C, followed by 40 cycles of 95°C for 30 s; 60°C for 30 s and 72°C for 45 s (where the annealing temperature, Ta, was as in Table 1). The specific oligonucleotide primers for the second stage of the rt-RT-PCR (rt-PCR of the cDNA), shown in Table 1, were selected using the Primer3 algorithm [28] from the *OsCam1-1* and *OsCML* cDNA sequences (see sequence analysis below).

The *O. sativa* elongation factor-1α (*OsEF1α*) was assumed to be independent of the abiotic stresses in its expression levels (a housekeeping gene), and so amplified as an internal control using the specific primers *OsEF1α*-F/R (Table 1), designed as above against the corresponding sequence (GenBank accession code AK105030). The level of transcript expression of the *OsCam1-1* and respective *OsCML* genes expression was then standardized against that for the *OsEF1α* gene transcript expression level from the same cDNA template as an internal control, and then expressed with reference to the standardized transcript expression level on day 0 of the treatment. Rt-RT-PCR reactions of each cDNA preparation were performed in triplicate for technical replication. Data were compared using the analysis of variance (ANOVA), and then the means were compared with Duncan’s multiple range test (DMRT) accepting significance at the p < 0.05 level.

**Determination of β-glucuronidase (GUS) activity**

The homozygous seeds of three independent *OsCam1-1*: *gus* transgenic rice lines [10] were hydroponically grown in 0.25 × strength NB medium [29] under a 16-h light/8-h dark photoperiod and treated with salt stress by adding, with minimal contact and disturbance, the medium containing 0 (control), 100, 150 or 300 mM NaCl into the chamber containing the rice seedlings to be treated. The leaf tissues of two-week-old rice seedlings were collected and frozen in liquid nitrogen at 0, 1, 2, 4, 8, 24 and 48 h after treatment. Rice seedlings were grown in a CRD with four replicates, and for each replicate, two seedlings were pooled for RNA extraction. The collected tissues were ground with a chilled mortar and a pestle, and homogenized in protein extraction buffer (50 mM Sodium phosphate, pH 7.0, 10 mM EDTA, 10 mM β-
Mercaptoethanol, 0.1% Sodium n-lauroylsarcosine, 0.1% Triton X-100). The β-glucuronidase enzyme (GUS) reaction assay was performed in protein extraction buffer containing 1 mM 4-MUG. For each reaction the substrate solution (200 μL) was pre-incubated at 37°C for 5 min, and then the reaction was initiated by the addition of 1 μL of the test plant extract and incubated at 37°C. After 10, 20, 30 and 40 min of incubation time 10 μL aliquots of the reaction mixture was transferred to 100 μL of stop solution (0.2 M Na2CO3). The standard curve was generated using five concentrations (10 nM to 100 nM) of 4-MU. The fluorescence intensity of the samples was measured by setting the excitation at 365 nm and the emission at 455 nm. The fluorescence intensity of the samples was generated using five concentrations (10 nM to 100 nM) of 4-MU. The fluorescence intensity of the samples was compared with the means were compared with REGWQ accepting significance at the p < 0.05 level.

Sequence analysis and microarray data retrieval

To determine which OsCML genes may have altered expression patterns in response to stress, their expression levels in different rice organs and tissues were compared. DNA microarray data of the OsCam and OsCML genes were retrieved from the publicly available microarray data of global gene expression from the experiment GSE 6893, which examined rice gene expression in various rice tissues/organs and stages of reproductive development from the IR64 rice cultivar or GSE 6901, which examined gene expression of 7-day-old rice seedlings grown under 3 h of dehydration (drought) or salt stress (200 mM NaCl) [12] via the rice oligonucleotide array database [25] available at (http://www.ricearray.org/).

For PCR design, nucleotide sequences from O. sativa were retrieved from of the Rice Genome Annotation Project database [31] and GenBank at the National Center for Biotechnology Information (http://www.ncbi.nlm.nih.gov/). Sequences of the following genes (loci): OsCam1-1 (LOC_Os03g20370), OsCML1 (LOC_Os01g59330), OsCML3 (LOC_Os12g03816), OsCML4 (LOC_Os03g53200), OsCML5 (LOC_Os12g41110), OsCML7 (LOC_Os08g02420), OsCML8 (LOC_Os10g25010), OsCML9 (LOC_Os05g41200), OsCML11 (LOC_Os01g32120) and OsCML13 (LOC_Os07g42660) were obtained.

To identify cis-acting regulatory DNA elements within the promoters of the OsCam1-1 and OsCML genes, their 5’ upstream sequences were analyzed using a Database of Plant Cis-acting Regulatory DNA Elements (PLACE) (http://www.dna.aaffrc.go.jp/PLACE/) and the Plant Promoter Analysis Navigator (PlantPAN) (http://plantpan.mbc.nctu.edu.tw/).

Competing interests

The authors declare that they have no competing interests.

Authors’ contributions

AC carried out the laboratory work along with KL and SP; and prepared figures and tables. KL and TB participated in database searches and data analyses. TL and TB performed data interpretation and drafted the manuscript. All authors read and approved the final manuscript.

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