Environmental stimuli such as UV, pathogen attack, and gravity can induce rapid changes in hydrogen peroxide (H₂O₂) levels, leading to a variety of physiological responses in plants. Catalase, which is involved in the degradation of H₂O₂ into water and oxygen, is the major H₂O₂-scavenging enzyme in all aerobic organisms. A close interaction exists between intracellular H₂O₂ and cytosolic calcium in response to biotic and abiotic stresses. Studies indicate that an increase in cytosolic calcium boosts the generation of H₂O₂. Here we report that calmodulin (CaM), a ubiquitous calcium-binding protein, binds to and activates some plant catalases in the presence of calcium, but calcium/CaM does not have any effect on bacterial, fungal, bovine, or human catalase. These results document that catalase/CaM can down-regulate H₂O₂ levels in plants by stimulating the catalytic activity of plant catalase. Furthermore, these results provide evidence indicating that calcium has dual functions in regulating H₂O₂ homeostasis, which in turn influences redox signaling in response to environmental signals in plants.

The absence of motility in higher plants has resulted in the acquisition of elaborate survival mechanisms to respond and adjust to a changing environment. The molecular and biochemical aspects of how plants perceive and respond to environmental signals are not clearly understood. Recent investigations have revealed that reactive oxygen species (ROS), such as hydrogen peroxide (H₂O₂) and superoxide (O₂⁻) are central components of the signal transduction cascade involved in plant adaptation to the changing environment. ROS plays two divergent roles: exacerbating damage or signaling the activation of defense responses under biotic and abiotic stresses (1–5). H₂O₂ is beginning to be accepted as a second messenger for signals generated by means of ROS because of its relatively long half-life and high permeability across membranes (1–5). To allow for these different roles, cellular levels of ROS must be stringently controlled. Catalase, which degrades H₂O₂ into water and oxygen, is one of the major antioxidant enzymes (6). It is one of the first enzymes to be purified and crystallized and has gained a lot of attention in recent years because of its link to cancer, diabetes, and aging in humans and animals (7, 8). In plants, catalase scavenge H₂O₂ generated during mitochondrial electron transport, β-oxidation of the fatty acids, and most importantly photorepiratory oxidation (6). Accumulating evidence indicates that catalase plays an important role in plant defense, aging, and senescence. Catalase activity is influenced by other factors such as salicylic acid and nitric oxide. Salicylic acid nonspecifically inhibits the activity of all catalases or protects them from inactivation, possibly depending on the redox status of the cell (9, 10). It was recently reported that nitric oxide inhibits the activity of tobacco catalase (11). However, the process by which catalase becomes activated/inactivated is not well understood.

Environmental signals trigger rapid and transient increases in cytosolic Ca²⁺. The cytosolic Ca²⁺ concentration can reach up to 100 μM from the resting level of 100–200 nM after stimulation in animal cells (12). Calcium signals modulate cellular processes by means of high-affinity, calcium-binding proteins such as calmodulin (CaM), which in turn regulates the activity of downstream target proteins (13–16). Recent studies suggest that changes in intracellular redox and calcium homeostasis are unifying consequences of biotic and abiotic stresses. Treatment with H₂O₂ can stimulate increases in cytosolic Ca²⁺ (17) by activating the calcium channel (18). Nonetheless, H₂O₂ production in the oxidative burst requires a continuous Ca²⁺ influx, which activates the plasma membrane-localized NADPH oxidase (19–21). Ca²⁺-binding EF hands are present in the gp91phox subunit of NADPH oxidase (22). Ca²⁺/CaM have been proposed to increase H₂O₂ generation through Ca²⁺/CaM-dependent NAD kinase that affects the concentration of available NADPH during assembly and activation of NADPH oxidase (23). The cloning and characterization of plant NAD kinase would further substantiate this process. We report herein that Ca²⁺/CaM binds to plant catalase and enhances its activity. However, Ca²⁺/CaM had no effect on bacterial, fungal, bovine, or human catalase. This finding suggests that calcium/CaM can down-regulate H₂O₂ levels in plants. Based on these results, it is proposed that calcium/CaM plays a critical role in controlling H₂O₂ homeostasis in plants.

Materials and Methods

Screening the cDNA Library. The Arabidopsis thaliana cDNA expression library (Uni-Zap) from the Arabidopsis Biological Resources Center (Columbus, OH) was screened by using 35S-labeled potato CaM (PCM6) as described (24). The positive cDNA clones were sequenced on both strands. DNA sequences were analyzed by using GCG version 10.0 software.

Construction of DNA Templates Coding Arabidopsis Catalase 3 (AtCat3) Proteins. The templates coding for wild type and mutants were produced by PCR amplification from the original cDNA with AtCat3-specific oligonucleotides containing appropriate restriction sites (NdeI at the 5’ end and BamHI at the 3’ end) for cloning into the downstream of [His]₆ tag into the pET-14b expression vector (Novagen). AtCat3 was expressed in Escherichia coli strain BL21(DE3) pLysS according to Studier et al. (25). The nucleotide sequence of the cloned fragments derived by PCR amplification was determined after cloning into the pET-14b vector with oligonucleotides designed for sequencing from both sides of the pET-14b cloning sites as primers.

CaM Binding Assay. The recombinant AtCat3 proteins were extracted and purified essentially as described (24). The amount of protein was estimated by the Bradford’s method with a protein assay kit (Bio-Rad). Proteins were separated by SDS/PAGE, electroblotted onto poly(vinylidene difluoride) membranes (Millipore), and incubated with 35S-labeled recombinant CaM (PCM6) plus 0.2 mM CaCl₂, 0.4 mM EGTA, 0.2 mM MgCl₂, and 10 mM Tris/HCl buffer, pH 8.0, for 90 min. Following incubation, the membrane was rinsed extensively with buffer and exposed to X-ray film (Konica). The film was developed and scanned with a densitometer.

This paper was submitted directly (Track II) to the PNAS office.

Abbreviations: ROS, reactive oxygen species; CaM, calmodulin; AtCat3, Arabidopsis catalase 3; IPTG, isopropyl β-D-thiogalactoside.

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0.2 mM MnCl$_2$, respectively, in 25 mM Tris-HCl (pH 7.5), 200 mM NaCl. The membranes were washed with the same buffer without $^{35}$S-labeled CaM and exposed to x-ray film overnight.

**Gel Mobility-Shift Assay.** The synthetic peptide was prepared by using an Applied Biosystems peptide synthesizer 431A in the Laboratory of Bioanalysis and Biotechnology, Washington State University. Recombinant CaM (PCaM6) was purified as described (24) and dialyzed thoroughly in 25 mM Tris-HCl (pH 7.0). Samples containing 240 pmol (4 µg) of PCaM6 and different amounts of purified synthetic peptides in 100 mM Tris-HCl, pH 7.2, and either 0.2 mM CaCl$_2$ or 0.4 mM EGTA in a total volume of 30 µl were incubated for 1 h at room temperature. The samples were analyzed by non-denaturing PAGE as described (24).

**Purification of Tobacco Catalase.** The catalase was purified from the leaves of tobacco plants (*Nicotiana tabacum* cv. Xanthi) following Durner and Klessig (10). Briefly, 1 kg tobacco leaves was homogenized in 2 liters of extraction buffer containing 50 mM Tris-HCl (pH 7.0), 100 mM NaCl, 10% glycerol, 10 mM DTT, 1 µM leupeptin, 10 µM antipain, and 1 mM PMSF. The homogenate was centrifuged at 9,000 g for 15 min. Ammonium sulfate was added to the supernatant by 22% saturation and centrifuged at 14,000 g for 25 min. Then the ammonium sulfate concentration of the supernatant was brought to 65% saturation after centrifugation at 14,000 g for another 25 min, the pellet was resuspended in the extraction buffer and fractionated with 0.8 vol of 0°C ethanol/chloroform (3:1, vol/vol) containing 1 mM PMSF. The upper aqueous layer was centrifuged at 47,000 g for 25 min. The supernatant was diluted 4-fold with 20 mM Tris-HCl (pH 7.0) and applied to a phenyl-Sepharose column (Amersham Pharmacia). After washing with the same buffer, the eluate was eluted with 50% (vol/vol) ethylene glycol in 20 mM Tris-HCl, pH 7.0. The catalase was precipitated by the addition of 3 vol of 50 mM Tris-HCl (pH 7.0), containing 90% (wt/vol) ammonium sulfate, and centrifuged at 47,000 g for 25 min. After suspension and buffer exchange (PD-10, Amersham Pharmacia) against 10 mM Tris-HCl (pH 7.0), the sample was applied to a hydroxyapatite column (Bio-Rad) equilibrated with the same buffer. The column was washed with 40 mM Tris-HCl (pH 7.0), and the catalase was eluted with 200 mM Tris-HCl (pH 7.0), 200 mM NaCl. After buffer exchange (PD-10) against 25 mM Tris-HCl (pH 7.0), 150 mM NaCl, and 1 mM CaCl$_2$, was added. The samples were applied to a CaM-Sepharose column (Amersham Pharmacia). By washing with the same buffer, the catalase was eluted from the column with the buffer containing 25 mM Tris-HCl (pH 7.0) and 2 mM EGTA. Fractions containing the catalase were pooled and thoroughly dialyzed against buffer containing 25 mM Tris-HCl (pH 7.0) and 10% glycerol.

**Electrophoretic Techniques.** SDS/PAGE was performed by using 10% gel for catalase and 15% gel for CaM. Gels were stained with Coomassie blue R-250 or with silver nitrate by using the Bio-Rad silver stain kit. Western blotting was performed by transferring proteins to the nitrocellulose membranes. Catalase was detected with an mAb against tobacco catalase, CaM was detected with a monoclonal anti-bovine CaM antibody (Santa Cruz Biotechnology) and an ECL detection kit from Pierce.

**Catalase Activity Assay.** Catalases from *Aspergillus niger*, bovine liver, and human erythrocytes were purchased from Sigma and treated as described (10). The catalytic activities of the catalases were measured spectrophotometrically by monitoring the changes of H$_2$O$_2$ at OD$_{240}$ at room temperature as described (10). The reaction mixtures for each catalase (0.1 µM) included calcium alone (0.2 mM CaCl$_2$), CaM alone (0.25 µM), both calcium and CaM (0.2 mM CaCl$_2$ + 0.25 µM CaM), and control (without CaCl$_2$ and CaM) in 50 mM Tris-HCl, pH 7.0. The reactions were started by adding 10 mM H$_2$O$_2$ from a 30% stock (Sigma). Catalase activity is expressed in units of mmol of H$_2$O$_2$ decomposed/mg per min.

**Results and Discussion**

The *Arabidopsis* cDNA library was screened by using the $^{35}$S-labeled CaM-binding screening approach as described (24). One of the positive clones showed an identical nucleotide sequence to *AtCat3* in the database (GenBank accession no. U43147). To prove that *AtCat3* encoded a CaM-binding protein, the full coding region of *AtCat3* was subcloned into the pET14b expression vector. The recombinant AtCat3 was induced by isopropyl-$\beta$-D-thiogalactoside (IPTG) (Fig. 1A) and was purified by CaM affinity chromatography to near homogeneity. Total bacterial extract was used for CaM-binding assay, and it was shown that $^{35}$S-labeled CaM binds to the *AtCat3* protein in the presence of Ca$^{2+}$ (Fig. 1A). After adding EGTA, no CaM binding was observed. Furthermore, $^{35}$S-labeled CaM did not bind to *AtCat3* when CaCl$_2$ was replaced by other divalent cations such as MgCl$_2$ or MnCl$_2$ (Fig. 1A), suggesting that CaM binding to *AtCat3* is Ca$^{2+}$-dependent.

To identify the CaM-binding area in *AtCat3*, two C-terminal deletion mutants were used for $^{35}$S-CaM-binding assays. In the presence of 0.2 mM Ca$^{2+}$, CaM binds to the wild type and mutant 1–451, whereas CaM did not bind to mutant 1–414 (Fig. 1B), suggesting that the CaM-binding region for *AtCat3* is restricted to amino acids 415–451. To further confirm CaM binding, *AtCat3* was subjected to SDS/PAGE and Coomassie staining or transferred onto a nitrocellulose membrane. The membrane was incubated with $^{35}$S-labeled 50 nM CaM in a buffer containing 0.4 mM EGTA, 0.2 mM CaCl$_2$, 0.2 mM MgCl$_2$, or 0.2 mM MnCl$_2$ (Left). Coomassie staining gel showing the induction of the *AtCat3* by IPTG. (Right) CaM-binding assay showing that CaM specifically binds to *AtCat3* in the presence of calcium. (B) Mapping of CaM binding region in *AtCat3*. (Left) Coomassie staining gel showing the total proteins from wild-type or deletion mutants of *AtCat3* after IPTG induction. (Right) CaM-binding assay showing that calcium/CaM binds to wild type and mutant 1–451, but not to mutant 1–414. (C) Gel mobility-shift assay showing CaM binding to the synthetic peptide (amino acids 415–451 in *AtCat3*) (Left) in the presence of 0.2 mM CaCl$_2$ and (Right) in the presence of 0.4 mM EGTA.

![Fig. 1](image)
binding to AtCat3, a synthetic peptide corresponding to amino acids 415–451 of AtCat3 was incubated with CaM. The formation of the peptide-CaM complex was assessed by non-denaturing PAGE. The 36-mer peptide is capable of forming a stable complex with CaM in the presence of Ca²⁺ (Fig. 1C). In the absence of peptide, a single CaM band was observed. As the peptide concentration increased, another band with low mobility appeared, representing the peptide-CaM complex. When the molar ratio between the peptide and CaM was 1:1, only the peptide-CaM complex band was detected, indicating that there is only one CaM-binding site in the peptide. No peptide-CaM complex was formed in the presence of EGTA.

Fig. 2. Comparison of the C-terminal portion of AtCat3 (amino acids 385–492) with other catalases from plants, animals, fungi, and bacteria. The CaM-binding region of AtCat3 is marked with a line above the sequences. The GenBank accession numbers are: AtCat1, U43340; AtCat2, X64271; AtCat3, U43147; Tobacco1, Z36975; Tobacco2, Z36978; Tobacco3, Z36977; Pumpkin1, D55645; Pumpkin2, D55646; Pumpkin3, D55647; Maize1, X60135; Maize2, Z54358; Maize3, L05934; human, X04076; bovine, P00432; Drosophila, X52286; A. niger, Z31138; Bacillus, M80796.
of AtCat3 predicted the existence of a basic amphiphilic amino acid sequence corresponding to the CaM-binding region, with the C-terminal 108-aa sequence comparison of 13 representative catalases showed high homology in this portion (about 78% similarity and 70% identity), particularly in the region corresponding to the CaM-binding site or closely juxtaposed regions in the CaM-binding region, most of them have a secondary structural feature, the basic amphiphilic α-helix (27), such as the auxin-regulated protein ZmSAUR1 (24) and the plant chimeric Ca\textsuperscript{2+}/CaM protein kinase (28). Note that there are often negatively charged amino acids in CaM-binding regions, but the net charge was neutralized CaM-binding proteins are not conserved in the CaM-binding region, as shown by the amino acid sequence comparisons of 37 catalases from bacteria, animals, and plants were compared by using the PILEUP program in the GCG10 package. The amino acid sequence comparison revealed very high homology in the N-terminal 384 aa (about 60% similarity and 50% identity). However, in the C-terminal 108 aa, where the CaM-binding region is located, there was very little similarity between AtCat3 and other nonplant catalases (about 21% homology and 14% identity). Nevertheless, all plant catalases showed high homology in this portion (about 78% homology and 70% identity), particularly in the region corresponding to the CaM-binding area in AtCat3. Fig. 2 shows the CaM-binding site or closely juxtaposed regions in other characterized CaM-binding proteins are not conserved in the CaM-binding region, most of them have a secondary structural feature, the basic amphiphilic α-helix (27), such as the auxin-regulated protein ZmSAUR1 (24) and the plant chimeric Ca\textsuperscript{2+}/CaM protein kinase (28). Note that there are often negatively charged amino acids in CaM-binding regions, but the net charge is positive (16). Based on the helical wheel projection using the GCG program, it was determined that the amino acids 438–451 in AtCat3 are capable of forming a basic amphiphilic α-helix. It is clear that one side of the helical wheel is hydrophobic and the other side is hydrophilic with net positive charges. Analysis of the amino acid sequences corresponding to the CaM-binding site of AtCat3 predicted the existence of a basic amphiphilic α-helix in some of other plant catalases. Interestingly, among three catalase isoforms in maize, tobacco, Arabidopsis, and pumpkin, there is at least one isoform with the amphiphilic α-helical structure in each species, e.g., tobacco1 (amino acids 431–444), maize3 (amino acids 442–455), and pumpkin1 (amino acids 438–451), as well as Arabidopsis AtCat1 (amino acids 438–451). Whether it is true for all plant species is not clear because of the limited catalase sequences for other plant species in GenBank. Based on the crystal structure data of bacterial and mammalian catalases, α-helices are the typical structures in the C-terminal portion of these catalases (29), but no basic amphiphilic α-helical structure exists in this portion according to helical wheel projection.

To test whether CaM binds to purified catalase from plant, tobacco catalase was purified from the leaves based on Durner and Klessig (10). A CaM-Sepharose column was used as the last step in the purification process. The summary of the purification steps of catalase using 1,000 g of tobacco leaves is presented in Table 1. Tobacco catalase was purified to near homogeneity as judged by SDS/PAGE and silver staining (Fig. 3, lane 1). Catalase identity was confirmed by Western blotting with a mAb against the tobacco catalase (Fig. 3, lane 3). CaM binding to purified tobacco catalase was demonstrated by the following approaches. (i) A CaM-Sepharose column was used for purification of tobacco catalase. The recovery rate for this step was 81% (Table 1), suggesting most of the catalase from tobacco leaves is a CaM-binding protein. (ii) \textsuperscript{35}S-labeled CaM was used to test the binding to the purified catalase in the presence of 0.2 mM CaCl\textsubscript{2} (Fig. 3, lane 5). Recombinant AtCat3 was used as a control (Fig. 3, lanes 2, 4, and 6). Addition of 0.4 mM EGTA abolished CaM binding, suggesting that CaM binds to the plant catalase in a Ca\textsuperscript{2+}-dependent manner. CaM-binding assays were also carried out to determine whether it binds to fungal, bovine, or human catalase. The results revealed that none of these nonplant catalases showed any CaM binding (data not shown), which is in agreement with the amino acid sequence comparisons (Fig. 2).

The CaM-binding site or closely juxtaposed regions in other characterized CaM-binding proteins often function as the autoinhibitory or pseudosubstrate domains. This region maintains the target proteins in an inactive state in the absence of Ca\textsuperscript{2+} signal, such as in plant chimeric Ca\textsuperscript{2+}/CaM-dependent protein kinase (30) and glutamate decarboxylase (31). To study the significance of CaM binding to plant catalases, catalytic activities of the recombinant AtCat3 and purified tobacco catalase were measured in the presence and absence of CaM.

### Table 1. Purification of catalase from tobacco leaves

<table>
<thead>
<tr>
<th>Purification steps</th>
<th>Total protein, mg</th>
<th>Total activity, unit*</th>
<th>Recovery rate, %</th>
<th>Specific activity, unit/mg</th>
<th>Purification factor*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Crude extract</td>
<td>3,640</td>
<td>362</td>
<td></td>
<td>0.099</td>
<td>B/A</td>
</tr>
<tr>
<td>Ethanol/chloroform</td>
<td>91.4</td>
<td>144</td>
<td>39.8</td>
<td>1.58</td>
<td>15.8</td>
</tr>
<tr>
<td>Phenyl-Sepharose</td>
<td>8.2</td>
<td>132</td>
<td>91.7</td>
<td>16.1</td>
<td>160</td>
</tr>
<tr>
<td>Hydroxyapatite</td>
<td>1.35</td>
<td>82.6</td>
<td>62.6</td>
<td>61.2</td>
<td>616.2</td>
</tr>
<tr>
<td>CaM-Sepharose</td>
<td>1.06</td>
<td>66.9</td>
<td>81.0</td>
<td>63.1</td>
<td>637.4</td>
</tr>
</tbody>
</table>

*Catalase activity in the absence of Ca\textsuperscript{2+}/CaM.

*Catalase activity in the presence of Ca\textsuperscript{2+}/CaM.

![Fig. 3. CaM binds to plant catalase. Lane 1, silver staining showing the purity of the catalase from tobacco leaves. Lane 3, Western blot showing that monoclonal tobacco catalase antibody can detect purified catalase. Lane 5, CaM-binding assay showing that \textsuperscript{35}S-CaM binds to purified catalase. Two-microgram recombinant AtCat3 was loaded as a control in each experiment (lanes 2, 4, and 6).](image-url)
of CaCl₂ and CaM. Similar experiments also were carried out by using other nonplant catalases. The E. coli-expressed recombinant AtCat3 showed no activity. Similar results were obtained in another laboratory (Zhixiang Chen, personal communication). This may have resulted from the failure to form the correct structure for recombinant catalase, because the active catalase is a tetramer made up of four identical or similar subunits (6, 10, 26). In contrast, the purified tobacco catalase showed a significant Ca²⁺/CaM-dependent change in activity during the purification steps (Table 1). The stimulatory effects of Ca²⁺/CaM are about 2.2-fold in each step, except the sample that was obtained after the CaM-Sepharose chromatography. It showed a higher stimulation of the catalase activity (2.5-fold) possibly because of the removal of non-Ca²⁺/CaM-binding catalase fraction (Table 1). The non-Ca²⁺/CaM-binding catalase fraction had a specific activity of 59.6 units (basal activity). However, Ca²⁺/CaM did not stimulate the activity of this fraction of catalase (data not shown). These results indicate that Ca²⁺/CaM is able to activate the tobacco catalase in the fraction that was obtained after CaM-Sepharose chromatography, which corresponds to the CaM-binding results. To compare the effects of Ca²⁺/CaM on the activity of catalases from different sources, the relative activity was used, because there is significant difference in the specific activity of catalases from different species. For example, A. niger catalase has the specific activity of 5.7 units; bovine liver catalase, 51.9 units; human erythrocytes, 39.5 units; tobacco catalase, 63.1 units (in the absence of Ca²⁺/CaM). Fig. 4A shows that CaCl₂ alone or CaM alone had no stimulatory effect on the tobacco catalase activity (about 40% of maximal activity). No difference in catalytic activity was observed in fungal, bovine, or human catalases in the presence or absence of Ca²⁺, CaM, and Ca²⁺/CaM (Fig. 4A). By replacing CaCl₂ with MgCl₂, there was no significant change in tobacco catalase activity in the presence or absence of CaM (data not shown). To further document the effect of Ca²⁺/CaM on plant catalase activity, 10 μM of the peptide corresponding to AtCat3 CaM-binding region (415–451) was added to each reaction mixture. Nonplant catalases were unaffected by the addition of the peptide. However, the stimulatory effect of Ca²⁺/CaM on tobacco catalase activity was abolished by the addition of the peptide (Fig. 4A). In addition, the inhibitory effect of the peptide on tobacco catalase activity depended on the concentration of the peptide (Fig. 4B). The catalase activity was inhibited by about 58%, which is close to the basal activity of catalase. However, the peptide had no effect on the bovine catalase activity at all concentrations tested (Fig. 4A). These results suggest that Ca²⁺/CaM has a stimulatory effect on the purified plant catalase, but has no effect on the nonplant catalases.

It is known that catalase is a predominant peroxisomal enzyme, but it also exists in the mitochondria and cytoplasm (32). For example, maize Cat3, which might be a CaM-binding catalase based on amino acid sequence comparison, is a mitochondrial protein (33). Our results (Figs. 1–4) demonstrate that CaM binds to plant catalase and regulates its activity. To demonstrate the presence of this regulatory activity in vivo, we studied whether CaM coexists with catalase in the peroxisomes of plants. Organelles from etiolated pumpkin cotyledon extracts were separated by sucrose density centrifugation. To remove the contamination of cytosolic proteins that might be present in the solution or merely associated with the peroxisomal membrane, the isolated peroxisomes were treated with proteinase K. In this way, the peroxisomal proteins shielded from protease by the peroxisomal membrane were not digested (34, 35). The identity of the peroxisomal fractions was proved by measuring the catalase activity (35). CaM is very conserved across kingdoms (13–16), hence we did Western analysis with an anti-human CaM antibody. As a control, the recombinant potato CaM PCM6 was used (Fig. 5, lane 3). Interestingly, a band in a size similar to PCM6 was present in the peroxisomal fractions, but the intensity was considerably lower compared to cytosolic CaM (Fig. 5). These results indicate that CaM and catalase are colocalized in...
the peroxisomes. CaM is a small acidic protein that is primarily expressed in the cytoplasm. However, in plants, CaM has been shown to be present in several organelles such as the nucleus and chloroplasts (36, 37) and extracellular matrix (38). Also, CaM-regulated proteins exist in extracellular matrix, nucleus, and chloroplasts (38–40). How CaM crosses the membrane is not clear. Recent studies indicate that posttranslational modifications play a role in the translocation of some CaM isoforms. For example, a petunia CaM-like protein, CaM53, is associated with plasma membrane when the protein is preylated. If preylation is inhibited, CaM53 is found predominantly in the nucleus (41).

To our knowledge, there is no report that discusses calcium concentration in the peroxisomes. Based on a recent modified model of the peroxisome biogenesis, the peroxisome originates from the endoplasmic reticular (ER) by means of preperoxisomal vesicles (42). It is known that the ER is one of the intracellular calcium pools. Thus, the calcium concentration in peroxisomes could be as high as in the ER. Measuring the peroxisomal calcium concentration and monitoring any link between the change of calcium concentration in cytoplasm and peroxisome should help in our understanding of how peroxisomal catalase is regulated by Ca2+/CaM. For example, the free calcium concentration could be measured in the transgenic plants with the reconstituted aequorin with a peroxisome-targeting signal. Aequorin is a calcium-sensitive luminescent protein, the luminescence of which directly reports the calcium change (43). Because peroxisome is a major organelle that removes the toxic ROS, mainly H2O2, it is reasonable to speculate that peroxisomal catalase activity remains high all of the time to degrade H2O2 in situ at a rapid rate. However, fluctuations in the cytosolic calcium could have a major effect on cytosolic catalase activity. It is important to stress that not all catalasas are CaM-binding proteins. Hence, further investigations are needed to address the significance of Ca2+/CaM in controlling catalase activity in different organelles.

Biotic and abiotic stresses trigger a Ca2+ influx, and the increased cytosolic Ca2+ stimulates the production of H2O2, which diffuses into surrounding cells as a messenger and induces the physiological response (19, 20). Our results suggest that increased cytosolic Ca2+ can reduce H2O2 levels by means of Ca2+/CaM-mediated stimulation of catalase activity (Fig. 4). We propose that increased cytosolic Ca2+ has dual roles in regulating H2O2 homeostasis (Fig. 6): (i) positive regulation generates H2O2 by directly activating NADPH oxidase, which has affinity to Ca2+ (22) and indirectly producing more NADPH by means of Ca2+/CaM-regulated NAD kinase (23); and (ii) negative regulation reduces H2O2 by stimulating catalase activity through Ca2+/CaM modulation (Fig. 4). Signal-induced changes in Ca2+ transients may differ in the frequency, duration, amplitude, and spatial localization of oscillations, and mode of spatial propagation, which leads to differential effects on the calcium target proteins (12, 44).

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