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Role of myristoylation in modulating PCaP1 interaction with calmodulin

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ABSTRACT

Plasma membrane-associated Cation-binding Protein 1 (PCaP1) belongs to the plant-unique DREPP protein family with largely unknown biological functions but ascertained roles in plant development and calcium (Ca^{2+}) signaling. PCaP1 is anchored to the plasma membrane via N-myristovlation and a polybasic cluster, and its Nterminal region can bind Ca²⁺/calmodulin (CaM). However, the molecular determinants of PCaP1-Ca²⁺-CaM interaction and the functional impact of myristoylation in the complex formation and Ca2+ sensitivity of CaM remained to be elucidated. Herein, we investigated the direct interaction between Arabidopsis PCaP1 (AtPCaP1) and CaM1 (AtCaM1) using both myristoylated and non-myristoylated peptides corresponding to the N-terminal region of AtPCaP1. ITC analysis showed that AtCaM1 forms a high affinity 1:1 complex with AtPCaP1 peptides and the interaction is strictly Ca^{2+} -dependent. Spectroscopic and kinetic Ca^{2+} binding studies showed that the myristoylated peptide dramatically increased the Ca²⁺-binding affinity of AtCaM1 and slowed the Ca²⁺ dissociation rates from both the C- and N-lobes, thus suggesting that the myristoylation modulates the mechanism of AtPCaP1 recognition by AtCaM1. Furthermore, NMR and CD spectroscopy revealed that the structure of both the N- and C-lobes of Ca²⁺-AtCaM1 changes markedly in the presence of the myristoylated AtPCaP1 peptide, which assumes a helical structure in the final complex. Overall, our results indicate that AtPCaP1 biological function is strictly related to the presence of multiple ligands, i.e., the myristoyl moiety, Ca²⁺ ions and AtCaM1 and only a full characterization of their equilibria will allow for a complete molecular understanding of the putative role of PCaP1 as signal protein.

1. Introduction

Calcium (Ca²⁺) is a crucial intracellular messenger in all eukaryotic organisms; in plants, Ca²⁺ signaling is involved in many events, such as cellular and developmental processes, response to biotic and abiotic stress, symbioses with rhizobial bacteria and mycorrhizal fungi, and immunity (Yang et al., 2003; Kudla et al., 2010, 2018; DeFalco et al., 2010; Oldroyd et al., 2008; Köster et al., 2022). The free cytosolic Ca²⁺, which is kept at a low concentration (~50–200 nM) in the resting state, rapidly increases upon stimuli perception due to the entry of Ca²⁺ from external and internal stores in a spatially and temporally controlled manner (Pirayesh et al., 2021; Lee et al., 2021). Afterward, Ca²⁺-binding sensor proteins transduce the Ca²⁺ signals into the downstream

responses by reversibly binding Ca^{2+} and interacting with various targets (Pirayesh et al., 2021; Xu et al., 2022). Different Ca^{2+} sensors exist in plants, such as calmodulins (CaM) and calmodulin-like-proteins (CMLs) (Astegno et al., 2016, 2017; Ogunrinde et al., 2017; La Verde et al., 2018a, 2018b; Dobney et al., 2009; Trande et al., 2019; Leba et al., 2012; Cho, 2016; Vallone et al., 2016; Vandelle et al., 2018), Ca^{2+} -dependent protein kinases (CDPKs) and calcineurin B-like proteins (CBLs) (DeFalco et al., 2010; Kudla et al., 2018; Edel et al., 2015), suggesting a high level of sub-functionalization and specificity in plants with respect to the animal counterpart.

CaM is one of the most conserved regulatory proteins in all eukaryotes. Seven CaM genes, encoding for four isoforms that share 90–100% primary sequence identity, were identified in the *Arabidopsis thaliana*

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genome (McCormack et al., 2003, 2005). CaM has a dumbbell-like structure consisting of two domains (the N- and C-lobes) connected by a flexible helix (McCormack et al., 2003; Perochon et al., 2011). Each globular domain contains two EF-hands that reversibly bind Ca^{2+} in a cooperative manner (La Verde et al., 2018b; Gifford et al., 2007). Binding of Ca^{2+} ions to CaM induces significant conformational changes resulting in the exposure of large hydrophobic grooves that usually mediate the target recognition (Astegno et al., 2016; Tidow et al., 2013; Gifford et al., 2013).

The Cation binding Protein 1 (PCaP1), also known as Microtubule-Destabilizing Protein 25 (MDP25), belongs to plant-specific Developmentally Regulated Plasma membrane Polypeptide (DREPP) family whose members interact with the plasma membrane (PM) and are differentially regulated during plant development (Vosolsobě et al., 2017). PCaP1 regulates many processes in plant life, such as microtubule organization, stomatal closure, root hydrotropism, and viral cell-to-cell movement (Yang et al., 2022; Nagata et al., 2016; Li et al., 2011; Qin et al., 2012; Vijayapalani et al., 2012). Moreover, PCaP1 was demonstrated to be involved in elicitor-induced plant immune response by either specific endogenous Damage-Associated Molecular Patterns (DAMPs), i.e., oligogalacturonides (OGs), and Pathogen-Associated Molecular Patterns, i.e., flagellin (flg22) (Giovannoni et al., 2021). These findings are in line with previous studies on elicitor-induced phosphoproteome changes of Arabidopsis thaliana that revealed PCaP1 as phosphoregulated in response to both OGs and flg22 (Mattei et al., 2016; Rayapuram et al., 2014).

PCaP1 from A. thaliana (AtPCaP1) is a hydrophilic, Ca²⁺-binding protein (Ide et al., 2007) composed of 225 residues with no obvious functional domain. It is anchored to the PM through both N-myristoylation at Gly2 and a relatively strong polybasic amino acid cluster in the N-terminal region that contribute to the interaction with phosphatidylinositol phosphates (PtdInsPs) (Vosolsobě et al., 2017; Kato et al., 2010; Nagasaki et al., 2008). Recently, AtPCaP1 was found to bind uranyl which induces protein oligomerization (Vallet et al., 2023). Furthermore, AtPCaP1 interacts with CaM, and the presence of Ca²⁺-CaM impairs the interaction of AtPCaP1 with PtdInsPs in a competitive manner (Kato et al., 2010; Nagasaki et al., 2008). The involvement of the N-terminal region in recognition of CaM and interaction with PtdInsPs was recently demonstrated also for the Arabidopsis isoform PCaP2, which is highly homologous to AtPCaP1 (25% overall primary sequence identity; 70% identity of the N-terminal region) (Kato et al., 2013), thus supporting a functional role of the AtPCaPs-CaM interaction in the transduction of the Ca²⁺ signals (Giovannoni et al., 2021; Ide et al., 2007; Kato et al., 2010, 2013; Tanaka-Takada et al., 2019). However, the molecular determinants of AtPCaP-CaM interaction and the role of myristoylation in the complex formation remain to be elucidated.

Here, we identified the specific CaM-binding site within the N-terminus of AtPCaP1 and investigated the *in vitro* properties of the interaction between myristoylated (myr) and non-myristoylated (non-myr) peptides corresponding to this site and the Arabidopsis CaM1 (AtCaM1) to clarify which roles AtCaM1 might play in AtPCaP1 regulation. The results obtained suggested that the N-terminal myristoylated region is directly involved in the AtPCaP1-AtCaM1 interaction, providing new insights into the interplay between AtCaM1 and AtPCaP1 in Ca²⁺mediated signaling processes in plants.

2. Materials and methods

2.1. Proteins production and peptide synthesis

AtCaM1 and its mutants B12Q-AtCaM1 (E32Q-E68Q) and B34Q-AtCaM1 (E105Q-E141Q) were generated and purified as described in (Astegno et al., 2016; Gut et al., 2009). Uniformly labeled ¹⁵N or ¹⁵N-¹³C-AtCaM1 samples for nuclear magnetic resonance (NMR) experiments were prepared using M9 minimal medium supplemented with

 $[^{13}C]$ -glucose and/or $[^{15}N]H_4Cl$ at a final concentration of 4 g L^{-1} and 1 g L^{-1} , respectively.

Synthetic gene (GenScript USA Inc.) corresponding to the complete cDNA of AtPCaP1 (UniProt Q96262) with a tag of six His at the C-terminus was cloned into pET21b expression vector and the recombinant plasmid was transformed into E. coli Rosetta (DE3) expression host cells. The culture was grown at 37 $^\circ C$ to OD_{600} of 0.6 and induced with 0.5 mM IPTG (isopropyl-β-d-thiogalactoside) at 24 °C for 16 h. The cells were harvested by centrifugation and resuspended in 20 mM Tris-HCl pH 7.5, 500 mM KCl, 20 mM imidazole, 10% glycerol, 1 mM DTT in the presence of protease inhibitor EDTA free. After sonication and centrifugation, the supernatant was applied to a Ni-chelating column equilibrated with 20 mM Tris-HCl pH 7.5, 500 mM KCl, 20 mM imidazole, 10% glycerol, 1 mM DTT. Protein elution was performed using a linear gradient from 20 to 500 mM imidazole. Purified protein fractions were pooled and extensively dialyzed in 50 mM Tris-HCl pH 7.5, 150 mM KCl, 0.5 mM DTT buffer to remove imidazole. The homogeneity and purity of the protein were verified by SDS-PAGE.

The myristoylated and non-myristoylated AtPCaP1 peptides (res: 2-GYWNSKVVPKFKKLFEKNSAKKAAA-26) were synthesized by Gen-Script USA Inc. The concentration of each peptide was determined using their predicted molar extinction coefficient at 280 nm.

2.2. Isothermal titration calorimetry

Isothermal titration calorimetry (ITC) experiments were performed on a MicroCal PEAQ-ITC instrument (Malvern Ltd., Malvern, UK) at 25 °C following protocols described in (Pedretti et al., 2020). AtCaM1, full-length AtPCaP1 and the AtPCaP1 peptides (non-myr-AtPCaP1p and myr-AtPCaP1p) were solubilized in the same buffer containing 50 mM KCl, 50 mM Tris (pH 7.5) in the presence of 5 mM CaCl₂ or 5 mM EGTA, filtered and degassed. The pH of the solutions was carefully checked to exclude any pH-variation during ITC experiments. For AtCaM1 - AtP-CaP1 peptides interaction, 200 µL of 25 µM AtCaM1 was titrated with injections of 1.5 μ L of each peptide (300–400 μ M), with a time gap of 120 s between each injection. For AtCaM1 - full-length AtPCaP1 interaction, 200 μ L of 18 μ M full-length AtPCaP1 was titrated with injections of 1.5 μ L of AtCaM1 (160 μ M), with a time gap of 120 s between each injection. Following the manufacturer's instructions, a first injection of $0.4 \ \mu L$ was made, and the first data point was removed in the fitting procedure. Data analysis was performed using the MicroCal ITC Origin 7 Analysis Software (MicroCal, Malvern Ltd., Malvern, UK) to obtain the apparent dissociation constant (K_d), the enthalpy changes (ΔH), and the apparent entropy change (ΔS) of the interaction. Results represent mean values \pm SEM of more than three independent experiments using at least two protein preparations.

2.3. Nuclear magnetic resonance spectroscopy

Nuclear Magnetic Resonance (NMR) experiments were performed on a 600 MHz Bruker Avance III spectrometer equipped with a triple resonance Prodigy cryo-probe (Bruker, Karlsruhe, Germany). All the experiments were performed at 298 K in 50 mM Tris, 50 mM KCl, 0.5 mM DTT (pH 7.5), supplemented with 5% D₂O. Titration experiments were performed by supplementing the Ca²⁺- bound ¹⁵N-AtCaM1 (0.55 mM) sample with increasing amounts of AtPCaP1p or myr-AtPCaP1p till reaching a final molar ratio of 1:2 (¹⁵N-AtCaM1:peptide) as described in (Bombardi et al., 2022; Conter et al., 2021). ¹H–¹⁵N-HSQC (Heteronuclear Single Quantum Correlation spectroscopy) experiments were acquired after each peptide addition with 2048 complex points in the direct dimension and 256 complex points in the indirect dimension. In total, 8 transients were acquired with a recycle delay of 0.2 s.

The combined ${}^{1}H^{-15}N$ Chemical shift perturbation (CSP) was calculated in agreement with the following equation:

$$CSP = \sqrt{\frac{\left(\Delta H\right)^2 + \left(\frac{\Delta N}{5}\right)^2}{2}} \tag{1}$$

Sequence-specific backbone resonance assignment of $^{15}N_{-}^{13}C_{-}$ AtCaM1 in its Ca²⁺-bound form in the presence or absence of AtP-CaP1peptides was achieved by recording sequential 3D HNCA, HNCACO, HNCACB, HNCO experiments, selected from the Bruker library, using 20% of randomly spread data points (Sattler et al., 1999; Schmieder et al., 1994). The sequential assignment was confirmed with a 3D¹⁵N-edited NOESY-HSQC experiment selected from the Bruker library, acquired with a matrix of 2048(F3) x 64(F2) x 128(F1) complex points and 16 transients, with a recycle delay of 1.2 s.

Backbone assignment was performed by recording a standard 3D $^{1}H^{-15}N$ NOESY-HSQC selected from the Bruker library. The experiment was acquired with a matrix of 2048(F3) x 64(F2) x 128(F1) complex points and 16 transients, with a recycle delay of 1.2 s.

All the spectra were processed using the software Topspin (Bruker) and NMRpipe (Delaglio et al., 1995) and successively analyzed with the program ccpnmr Analysis 2.2.1 (Vranken et al., 2005). Non uniform sampling (NUS) experiments were processed using the Sparse multidimensional iterative lineshape-enhanced (SMILE) reconstruction algorithm integrated in NMRpipe (Ying et al., 2017). Backbone assignment of Ca²⁺ bound AtCaM1 has been deposited into the BioMagResBank (BMRB) (BMRB entry 52071).

2.4. Circular dichroism spectroscopy

Far-UV circular dichroism (CD) spectra at 25 °C were recorded between 200 (or 190) and 250 nm using 0.1 cm quartz cuvettes on a Jasco J-1500 spectropolarimeter equipped with a temperature control device. Protein and peptide samples were dissolved in a buffer containing 50 mM Tris–HCl, 150 mM KCl, 0.5 mM DTT (pH 7.5) and supplemented with 5 mM CaCl₂. Spectra were collected as an average of three scans as previously described (Bombardi et al., 2022; Spyrakis et al., 2011).

2.5. Ca^{2+} binding assay with chromophoric chelator

The binding of Ca²⁺ to AtCaM1 in the absence or presence of AtP-CaP1 peptides was studied by 5,5'-Br₂-BAPTA [5,5'-dibromobis-(o-aminophenoxy) ethane N,N,N',N'-tetra-acetic acid] competition essay (Andre et al., 2002; Astegno et al., 2014; Linse et al., 1991). Briefly, the decrease in the absorbance ($\lambda = 263$ nm) of 25 μ M 5,5'Br₂-BAPTA (Ca²⁺ affinity = 2.3 μ M in 0.15 M KCl) in the presence of 25 μ M AtCaM1 was followed upon addition of 3 μ M Ca²⁺ for each titration point at 25 °C. Decalcified proteins were solubilized in decalcified 50 mM Tris–HCl pH 7.5, 150 mM KCl buffer (initial Ca²⁺ concentration in the buffer ranged between 0.3 and 0.5 μ M). The stoichiometric ratio AtCaM1: AtPCaP1 peptide was set to 1 : 1.5. Data were normalized as follows:

normalized
$$A = \frac{A_{263} - A_{min}}{A_{max} - A_{min}}$$
 (2)

where A_{263} is the absorbance at each titration point, A_{\min} and A_{\max} are the absorbance values at the lowest and highest Ca²⁺ concentration, respectively.

The normalized signal decrease upon Ca^{2+} titration was fitted to a four-site binding model for AtCaM1 (Astegno et al., 2016) using CaLigator (Andre et al., 2002) to estimate individual macroscopic binding constants.

2.6. Kinetic rate measurements

 Ca^{2+} dissociation rate constants from AtCaM1 complexed with AtPCaP1 peptides were measured using the SX-18 stopped-flow apparatus (Applied Photophysics, Leatherhead, UK) taking advantage of the Ca^{2+} chelator Quin-2 whose fluorescence changes upon binding of Ca^{2+} .

Stopped flow experiments were performed at 11 °C in 50 mM Tris-HCl, 150 mM KCl (pH 7.5) as previously described (Troilo et al., 2022). The change of Quin-2 fluorescence signal as a function of time was monitored using an excitation wavelength $\lambda_{ex} = 332$ nm, and a 435 nm emission cut-off filter. Solutions of 6 μ M AtCaM1 or AtCaM1 variants (B12Q and B34Q) and 18 μ M AtPCaP1 or myr-AtPCaP1 peptides, in 50 μ M Ca²⁺, were rapidly mixed with 150 μ M Quin-2.

For each trace, data from 4 to 8 injections were averaged and fitted with either a single (Eq (3)) or double (Eq (4)) exponential function.

$$F = a \times e^{(-k_{obs} \times t)} + F_{max} \tag{3}$$

Where *F* is the observed fluorescence intensity at time *t*; *a* is the amplitude of the fluorescence change; k_{obs} is the rate constant at which the change in fluorescence is occurring, representing the Ca²⁺ dissociation rate (k_{off}) from the proteins; and F_{max} is the fluorescence signal at infinite time.

$$F = a_1 \times e^{(-k_{obs1} \times t)} + a_2 \times e^{(-k_{obs2} \times t)} + F_{max}$$
(4)

Where: a_1 and a_2 are the individual amplitudes of each component of the fluorescence change; k_{obs1} and k_{obs2} are the corresponding rate constants.

Data were normalized to maximum (1) and minimum (0) fluorescence. Graphs were obtained using Prism-GraphPad 6. All the traces were fitted excluding data points in the dead-time of the stopped-flow (< 2 ms).

3. Results

3.1. Identification of CaM-binding site in AtPCaP1

A putative CaM-binding region was identified in the first 26 N-terminal amino acids (²GYWNSKVVPKFKKLFEKNSAKKAAA²⁶) of the AtPCaP1 sequence using the Calmodulin Target Database (Yap et al., 2000). This region possesses the typical 1–8–14 motif, and an α -helical wheel representation of the predicted sequence shows that the positively charged residues are located on one side of the wheel, while the hydrophobic amino acids are on the other side when the peptide forms an α -helical structure (Fig. 1A) (Tidow et al., 2013). Peptides mimicking this putative CaM-binding domain in AtPCaP1 (AtPCaP1p) were synthesized in both myristoylated (at Gly2) and non-myristoylated forms. The far-UV CD spectra of AtPCaP1p (Fig. 1B) and myr-AtPCaP1p (Fig. S1A) recorded in the presence of varying amounts of 2,2,2-Trifluoroethanol (TFE) confirmed that they have a strong propensity to form α -helical structures. TFE strengthens the hydrogen bonding inducing and stabilizing α -helices in sequences with helical propensity, therefore mimicking the hydrophobic environment of these regions in the intact protein. In aqueous solution, the AtPCaP1 peptides were largely unstructured with a CD spectrum characterized by a negative peak centered around 200 nm (Fig. 1B and Fig. S1A). However, when the TFE concentration was increased to 20% v/v, the peptides adopted an α -helical structure, with two peaks at 208 and 222 nm. The helical content further increased at higher [TFE], thus confirming the ability of peptides to acquire *a*-helical conformation in hydrophobic environments.

Since in most CaM-peptide interactions, the peptide undergoes a major secondary structure rearrangement from a random coil, flexible structure when isolated to an α -helix structure upon complex formation, we next explored this aspect in the AtCaM1-AtPCaP1p interaction using CD spectroscopy. The far-UV CD spectrum of AtCaM1 is typical of an alpha-helical protein (Astegno et al., 2014, 2016), while the free peptides appear to be unstructured (Fig. 1C and Fig. S1B). However, the addition of AtPCaP1 peptides is accompanied by an increase in the dichroic signal, which can be attributed to a coil-to-helix transition of the peptide as it binds to AtCaM1 (Fig. 1C and Fig. S1B), as reported for many other CaM-target peptides (Astegno et al., 2016, 2017; La Verde et al., 2018a). Overall, these analyses clearly indicate that the first 26



residues of AtPCaP1 possess the features of the typical CaM-binding motifs.

3.2. Thermodynamics of AtCaM1-AtPCaP1 interaction

The energetics of AtPCaP1 peptides binding to AtCaM1 were studied using isothermal titration calorimetry (ITC). The thermodynamic parameters of the interaction are reported in Table 1. Representative thermograms and the binding curves for the titration of AtPCaP1p or myr-AtPCaP1p into AtCaM1 in the presence of Ca²⁺ or EGTA are shown in Fig. 2.

Our data showed that in the presence of Ca^{2+} , all the binding reactions take place with an exothermic heat exchange (negative enthalpy) and a 1:1 stoichiometry. For the Ca^{2+} -AtCaM1-AtPCaP1p couple, the experimental data could be best fitted to a one-site binding model, giving a dissociation constant (K_d) of 911 ± 63 nM. Notably, the presence of the myristoyl-group at Gly2 significantly enhances the affinity of AtCaM1 for the peptide ($K_d = 66 \pm 12$ nM). In contrast, no significant binding of AtPCaP1 peptides to AtCaM1 was detected in the absence of Ca^{2+} (i.e., in the presence of the metal chelator EGTA), thus indicating the strong dependence on Ca^{2+} of the AtCaM1-AtPCaP1 interaction. The negative T Δ S value (Table 1), which is largely compensated by the negative enthalpic contribution of the reaction, confirmed a conformational rearrangement of AtPCaP1 peptides to an ordered helix upon binding to AtCaM1 as shown by CD spectroscopic analysis (Fig. 1).

Table 1

Thermodynamic parameters of the interaction between AtCaM1 and AtP-CaP1 peptides in the presence of 5 mM CaCl₂ at 25°C. The reported parameters are the mean \pm standard error of the mean (SEM) of at least three independent titrations using two different protein preparations.

	n	$K_{\rm d}({\rm nM})$	ΔH (kcal mol ⁻¹)	-T Δ S (kcal mol ⁻¹)	
Ca ²⁺ /AtCaM1 + myr- AtPCaP1p	0.9 ± 0.1	66 ± 12	-12.5 ± 0.8	2.7 ± 0.8	
Ca ²⁺ /AtCaM1 + AtPCaP1p	$\begin{array}{c} 1.1 \ \pm \\ 0.0 \end{array}$	$\begin{array}{c} 911 \pm \\ 63 \end{array}$	-17.0 ± 0.5	8.7 ± 0.5	

Fig. 1. Analysis of CaM binding region of AtP-CaP1. (A) The primary sequence of AtPCaP1 is reported together with the CaM binding score obtained by the Calmodulin Target Database search (Yap et al., 2000) and the wheel model of the first 26 N-terminal residues. (B) CD spectra of AtPCaP1p (in aqueous solution or at different [TFE] mixtures). (C) CD spectra of AtCaM1 (black line) and AtPCaP1p (red line), and the protein-peptide complex (blue line) in the presence of 5 mM CaCl₂. The green line results from subtracting the blue line (complex) and the black line (AtCaM1). All spectra were recorded using 0.2 mg mL⁻¹ of AtCaM1. For the complex, a 1:2 M ratio of the peptide was added.

We also produced the recombinant full-length AtPCaP1 in *E. coli* and purified it to homogeneity, as evidenced by SDS-PAGE (Fig. S2A). The structural integrity of the protein was further validated through far-UV CD analysis, which revealed a characteristic spectrum indicative of a protein with alpha helix content, displaying double minima at 208 and 222 nm (Fig. S2B).

The thermodynamic properties of the interaction between the fulllength AtPCaP1 (in the non-myristoylated form) and AtCaM1 was studied by ITC in the presence of Ca²⁺ or EGTA. ITC data (Fig. S3 and Table S1) showed that the AtPCaP1-AtCaM1 interaction is a Ca²⁺dependent exothermic binding reaction with a favorable enthalpy (–9.4 \pm 0.5 kcal mol⁻¹) but an unfavorable entropy (1.1 \pm 0.4 kcal mol⁻¹). The dissociation constant (K_d) for the complex is 859 \pm 62 nM and the stoichiometry is 0.8 \pm 0.1, indicating high affinity 1:1 complex formation (Table S1). In the presence of EGTA, no binding was observed.

ITC results obtained with the full-length AtPCaP1 are consistent with those obtained using the non-myr peptide approach; this provides strong evidence that the N-terminal region of AtPCaP1 fully recapitulates the interaction of AtPCaP1 with AtCaM1, making it a suitable representative model for further studies.

3.3. The myristoyl moiety has a significant effect on the conformation of the AtCaM1-AtPCaP1 complex

An NMR-based approach was used to structurally characterize Ca²⁺-AtCaM1 when bound to AtPCaP1 peptides. We collected twodimensional ¹H, ¹⁵N-HSQC NMR spectra of ¹⁵N-labeled Ca²⁺-AtCaM1 in the absence and presence of unlabeled AtPCaP1 peptides. As shown in Fig. 3A, several resonances of AtCaM1 underwent significant chemical shift perturbation (CSP) and intensity variation following the addition of both myr and non-myr peptides, indicating the physical interaction between the protein and the peptides and the formation of a well-organized Ca²⁺-protein-peptide complex.

To further investigate the structural basis of AtCaM1 interaction with AtPCaP1 peptides, we assigned the backbone chemical shifts of free Ca^{2+} - AtCaM1 and Ca^{2+} -AtCaM1 when bound to myr and non-myr AtPCaP1 peptides (Fig. S4). The chemical shift assignments were obtained using standard double- and triple-resonance NMR experiments



conducted on uniformly ¹⁵N- and ¹³C/¹⁵N -labeled protein samples. The goodness of the assignment was also confirmed using a 3D ¹⁵N-edited NOESY, a technique that provides information on nuclei closed in space (usually ~5–6 Å) and by comparing the NMR resonances of Ca²⁺-AtCaM1 from different species that have been previously assigned by several groups (Ikura et al., 1990, 1992; Chou et al., 2001). The backbone chemical shifts of AtCaM1 were assigned with 97% completion for free Ca²⁺-AtCaM1 and Ca²⁺-AtCaM1 in complex with the non-myr AtPCaP1 peptide, excluding the two Pro residues and the first three N-terminal residues (Met, Ala, and Asp), due to their intrinsic flexibility. Instead, upon addition of myr-AtPCaP1p several resonances underwent significant CSP and signal broadening, thus we could assign approximately the 87% of the signals present in the ¹⁵N-HSQC spectrum.

Structural changes in Ca²⁺-AtCaM1 induced upon peptide binding were probed by comparing the backbone 1 H and 15 N chemical shifts of

free Ca²⁺-AtCaM1 and Ca²⁺-AtCaM1 bound to each peptide. Differences in these chemical shifts were quantified and the CSPs plotted as a function of residue number (Fig. 3B). Notably, the degree and the extent of backbone CSPs were markedly different in our two protein-peptide complexes (Fig. 3B–D). Upon addition of myr-AtPCaP1p to AtCaM1, most of the ¹⁵N-AtCaM1 resonances that underwent extensive chemical shift changes (>0.2 ppm) were dispersed throughout the protein backbone and were present within both the N- and C- terminus of the ¹⁵N-AtCaM1 sample, suggesting that a global conformational transition of AtCaM1 occurs upon binding of the myr-peptide (Fig. 3B and C). In particular, the region spanning from residue I64–K74 corresponding to the linker between the N-terminal and C-terminal lobes shifted significantly with an average CSP of ~0.35 ppm. We also noticed that many hydrophobic residues namely F20, V56, F93, V109, M110, N111, L112 and M146 disappeared from the spectrum after peptide binding and

Fig. 2. AtPCaP1 peptides binding to AtCaM1 studied by ITC in the presence and absence of Ca²⁺. Representative thermograms (top panels) and the derived binding isotherms (bottom panels) of titration of AtPCaP1p (A, B) and myr-AtPCaP1p (C, D) into AtCaM1 in the presence of 5 mM CaCl₂ (A, C) or 5 mM EGTA (B, D) at 25 °C. The ligand dilution blank experiment (peptide titrated into buffer) was subtracted from the binding isotherm obtained in the presence of protein. A first injection of 0.4 μ L was made, and the first data point was removed from the data fitting.



Fig. 3. Interaction between AtCaM1 and AtPCaP1 peptides revealed by NMR. (A) Overlaid HSQC spectra of Ca²⁺-AtCaM1 (black) with Ca²⁺-AtCaM1-AtPCaP1p (light blue) or Ca²⁺-AtCaM1-myr-AtP-CaP1p (orange). A subset of assigned peaks displaying significant movement upon peptide binding are indicated. (B) Backbone amide 1H and 15N CSPs induced by the binding of either non-myr (light blue) or myr-AtPCaP1p (black) to Ca²⁺ saturated AtCaM1 as a function of amino acid residue number. Secondary structural elements, derived from the structure of human CaM in complex with Ca²⁺ (PDB:1CLL) are displayed on the top of the figure as α -helices (grey rectangle) and β -sheets (black rectangle). Regions undergoing significant chemical shift changes are color coded in panels C and D onto the 3D structure of Ca²⁺ AtCaM1 from white (CSP <0.1 ppm) to red (CSP >0.3 ppm). (C, D) Three-dimensional representation of perturbed residues in Ca²⁺-AtCaM1 (model calculated using AlphaFold2) upon binding of the synthetic myr-AtPCaP1p (C) or synthetic non-myr AtPCaP1p (D). Residues displaying ¹H-¹⁵N chemical shift perturbation are mapped as colored spheres on the protein structure from vellow (CSP > 0.1 ppm) to red (CSP >0.3 ppm). Residues with a CSP <0.1 ppm are displayed in white. Ca²⁺ atoms are displayed as blue spheres.

could not be assigned. These residues were primarily found in the hydrophobic clefts of the N and C-domain of AtCaM1 and seem to be directly involved in the binding of the myristoyl moiety providing hydrophobic contacts in the human CaM (PDB: 1L7Z) (Matsubara et al., 2004).

On the other hand, although a few backbone resonances in the N-lobe and the central linker of AtCaM1 were affected by the binding of non-myr AtPCaP1p, most changes map to AtCaM1's C-lobe (residues 86–149), suggesting a crucial role for the C-terminal domain in the target recognition (Fig. 3B and D).

Notably, inspection of the downfield resonances corresponding to the glycine residues at position six of the four Ca^{2+} -binding EF-loops (G26, G62, G99 and G135) revealed that the peaks belonging to G26 (EF-1) and G62 (EF-2) of AtCaM1 remain unperturbed upon non myrpeptide binding, supporting the conclusion that AtCaM1 N-terminal domain is not critically involved in the binding of non-myr AtPCaP1p. On the contrary, all the four peaks corresponding to G26, G62, G99 and G135 were characterized by a clear chemical shift variation upon AtCaM1-myr AtPCaP1p complex formation.

Overall, our NMR data suggest that the binding of the two AtPCaP1 peptides to AtCaM1 results in complexes of different final conformations with the myristoyl group that has a large impact on AtCaM1 conformation.

Moreover, an accurate analysis of NMR titration data for binding of unlabeled AtPCaP1p or myr-AtPCaP1p to Ca²⁺-AtCaM1 indicates that, upon increasing concentrations of peptides, most of the cross-peaks in the ¹H–¹⁵N HSQC spectrum of ¹⁵N-AtCaM1 display a strong intensity decrease without additional line broadening. This is a classical behavior of resonances in slow exchange on the NMR time scale, which typically corresponds to an intermolecular dissociation constant on the order of 10^{-7} M or less (Latham et al., 2009; Favretto et al., 2020); furthermore, only a single set of AtCaM1 resonances was observed when bound to

peptides, pointing to a 1:1 stoichiometry, in agreement with the above-described ITC results.

0.3

We also analyzed the solution properties of the AtCaM1 alone and in complex with both peptides by size exclusion chromatography (SEC) in the presence of Ca^{2+} (Fig. S5A). Specifically, the myr-AtPCaP1p-AtCaM1 complex elutes later compared to AtCaM1 alone. This suggests that the binding of the myristoylated peptide induced a conformational change in AtCaM1 that leads to a smaller hydrodynamic radius, likely due to a more compact entity. On the other hand, the AtPCaP1p-AtCaM1 complex elutes earlier than AtCaM1 alone, suggesting a larger hydrodynamic radius for this complex. These observations are consistent with NMR data and support the notion that myr-AtPCaP1p and AtPCaP1p interact with AtCaM1 in different ways, leading to complexes with distinct final conformations.

SEC analysis performed with the full-length AtPCaP1 again recapitulates results obtained with the AtPCaP1 peptide (Fig. S5B). Indeed, the AtPCaP1-AtCaM1 complex elutes earlier than AtCaM1 alone, highlighting the specific and relatively high-affinity binding between these two proteins in the presence of Ca^{2+} .

3.4. Effect of AtPCaP1 peptides on AtCaM1 Ca^{2+} affinity and Ca^{2+} dissociation kinetics

The effect of AtPCaP1 peptides on the Ca²⁺ binding properties of AtCaM1 was quantitatively evaluated using a competition assay based on the Ca²⁺ chelator 5,5'-Br₂-BAPTA, whose absorption decreases upon ion binding (Andre et al., 2002; Linse et al., 1991). Representative Ca²⁺ titration curves of AtCaM1 alone and in the presence of AtPCaP1 peptides are shown in Fig. 4. Since AtCaM1 has been demonstrated to possess four Ca²⁺ binding sites (Astegno et al., 2014, 2016), data were fitted with a four-site binding model. Titration of AtCaM1 with 5, 5'-Br₂-BAPTA allowed to calculate an overall K_d of ~16 µM for Ca²⁺



Fig. 4. Ca^{2+} titration curves obtained by absorption spectroscopy in the presence of the chromophoric Ca^{2+} chelator 5,5'Br₂-BAPTA. Experimental points for AtCaM1 alone (black circles) and for AtCaM1 in the presence of myr-AtPCaP1p (red triangle) or AtPCaP1p (blue square) are shown together with the optimal curves calculated by computer fitting to a four-states model. Calculated Ca^{2+} -binding constants are reported in Table 2; model fitting and data normalization are detailed in the Methods section.

binding to AtCaM1, which is in perfect agreement with the value ($K_d \sim 13 \ \mu$ M) previously obtained (Table 2) (Astegno et al., 2016). Interestingly, all the four individual macroscopic binding constants significantly changed in the presence of the myr-peptide, revealing an overall K_d value of ~180 nM. Thus, the apparent affinity for Ca²⁺ of AtCaM1 was increased of approximately 90-fold in the presence of this peptide. On the other hand, in the presence of the non-myr peptide an apparent K_d value of ~0.9 μ M was measured, such that the affinity for Ca²⁺ was increased of only ~18-fold (Table 2). In this case, the most notable changes in macroscopic binding constants were observed only for two out of the four EF-hands (K_{d1} and K_{d2}) (Table 2).

Next, we analyzed the effects of the peptides binding on the kinetics of Ca²⁺ dissociation from AtCaM1 by stopped-flow spectroscopy, taking advantage of the fluorescence increase of the Ca²⁺chelator Quin-2 upon Ca²⁺ binding. As reported in our previous work, in the absence of targets, Ca²⁺ dissociation from the N-lobe of AtCaM1 occurs significantly faster than from the C-lobe ($k_{obs} > 500 \text{ s}^{-1}$ and $k_{obs} = 15.6 \text{ s}^{-1}$, respectively, Table 3) (Troilo et al., 2022). Notably, in the presence of both AtPCaP1 peptides, the Ca^{2+} dissociation kinetics is a biphasic process (Fig. 5) characterized by two rate constants: $k_{obs1}=3.6\pm0.02$ s^{-1} and k_{obs2} = 0.12 \pm 0.005 s^{-1} with myr-AtPCaP1p (Fig. 5A) and k_{obs1} = 34.6 \pm 0.23 s^{-1} and k_{obs2} = 0.20 \pm 0.001 s^{-1} with the non-myr peptide (Fig. 5D, Table 3). To assign the Ca^{2+} dissociation rates to each lobe of the protein, we used AtCaM1 mutants deficient in Ca²⁺ binding at sites 1 and 2 of the N-lobe (B12Q-AtCaM1) or sites 3 and 4 of the C-lobe (B34Q-AtCaM1) in which only the C-lobe or the N-lobe is competent for Ca²⁺ binding, respectively (Troilo et al., 2022). Ca²⁺ dissociation from the N-lobe (B34Q-AtCaM1) gives rise to a dissociation rate of 3.8 \pm 0.01 s⁻¹ and 28 \pm 0.12 s⁻¹ in the presence of

Table 3

 Ca^{2+} dissociation kinetics. The reported parameters are the mean \pm standard error of the mean (SEM) of at least three independent titrations using two different protein preparations.

	N-lobe	C-lobe
	k_{obs} (s ⁻¹)	$k_{obs} (s^{-1})$
AtCaM1 ^a	>500	15.6 ± 0.1
B34Q-AtCaM1 ^a	>500	
B12Q-AtCaM1 ^a		14.4 ± 0.12
AtCaM1 + myr-AtPCaP1p	3.6 ± 0.02	0.12 ± 0.005
B34Q-AtCaM1 + myr-AtPCaP1p	$\textbf{3.8} \pm \textbf{0.01}$	
B12Q-AtCaM1 + myr-AtPCaP1p		0.24 ± 0.001
AtCaM1 + AtPCaP1p	34.6 ± 0.23	$\textbf{0.2} \pm \textbf{0.001}$
B34Q-AtCaM1 + AtPCaP1p	28 ± 0.12	
B12Q-AtCaM1 + AtPCaP1p		$\textbf{0.28} \pm \textbf{0.004}$

^a Values from (Troilo et al., 2022).

myr-AtPCaP1p and AtPCaP1p, respectively (Fig. 5B and E, Table 3); while the Ca²⁺ dissociation from the C-lobe (B12Q-AtCaM1) results in a $k_{obs} = 0.24 \pm 0.001 \text{ s}^{-1}$ in the presence of myr-AtPCaP1p, and $k_{obs} = 0.28 \pm 0.004 \text{ s}^{-1}$, in the presence of AtPCaP1p (Fig. 5C and F, Table 3). Therefore, the binding of both peptides causes an overall decrease in the Ca²⁺ dissociation rate constants from both AtCaM1 lobes (N-lobe: from >500 to 3.8 for the myr peptide and to 28 s⁻¹ for the non-myr peptide, respectively; C-lobe: from 14.4 to 0.24 and 0.28 s⁻¹ for the myr-AtPCaP1p and AtPCaP1p, respectively) (Table 3) in agreement with the BAPTA experiments which demonstrated an overall increase in the AtCaM1 Ca²⁺ affinity in the presence of the target peptides. Interestingly, the release of Ca²⁺ from the N-lobe occurs \cong 7-fold faster in the case of AtPCaP1p ($k_{obs} = 28 \text{ s}^{-1}$) compared to myr-AtPCaP1p ($k_{obs} = 3.8 \text{ s}^{-1}$) (Table 3).

4. Discussion

PCaP1 belongs to the *Arabidopsis* DREPP family which comprises proteins that interact peripherally with the PM by various mechanisms, including combinations of N-myristoylation and protein-lipid electrostatic interactions due to the presence of a polybasic cluster (Vosolsobě et al., 2017). DREPP proteins have been described as signal proteins and their specific mechanism of PM association has been suggested to be crucial for their localization in signaling platforms in membrane microdomains and thus for their specific function.

According to the AlphaFold model (Fig. S6) more than one-third of the AtPCaP1 protein, particularly the C-terminal region, is predicted to be disordered, while the first 125 residues exhibit α -helical structure which is consistent with the far-UV CD spectrum of the recombinant protein (Fig. S2B). The structured region of AtPCaP1 is characterized by a central core composed of helices from α 3 to α 7 which is stabilized by hydrophobic interactions. The last part of helix α 1 folds over the central core and, together with α 2, contributes to capping on one side of the central core. Additionally, the N-terminal portion of the α 1 extends outwards the core. The electrostatic surface of AtPCaP1 shows that helix α 1 has a strong positive charge, and together with three lysine residues located on helix α 4, it constitutes an extended positively charged surface (Fig. S6).

A closer look at the N-terminal sequence and structural model of AtPCaP1 shows that it contains not only the myristoyl moiety and a polybasic flanking region, which greatly contribute to its interaction

Table 2

Macroscopic Ca²⁺ binding constants for AtCaM1 in the absence and presence of AtPCaP1 peptides.

	log Ka ₁	$K_{\rm d1}~(\mu{\rm M})$	log Ka ₂	$K_{\rm d2}~(\mu{ m M})$	log Ka ₃	$K_{\rm d3}~(\mu{ m M})$	log Ka ₄	$K_{\rm d4}~(\mu{ m M})$	Overall K_d
AtCaM1 ^a AtCaM1 + myr-AtPCaP1p AtCaM1 + AtPCaP1p	$\begin{array}{c} 4.8 \pm 0.1 \\ 8.2 \pm 0.1 \\ 7.1 \pm 0.1 \end{array}$	15.8 0.006 0.08	$\begin{array}{c} 5.8 \pm 0.2 \\ 6.3 \pm 0.1 \\ 7.4 \pm 0.1 \end{array}$	1.6 0.5 0.04	$\begin{array}{c} 3.5 \pm 0.2 \\ 6.7 \pm 0.2 \\ 4.2 \pm 0.1 \end{array}$	316 0.2 63	$\begin{array}{c} 5.5 \pm 0.1 \\ 5.8 \pm 0.1 \\ 5.5 \pm 0.2 \end{array}$	3.2 1.6 3.2	16 μM 180 nM 0.9 μM

^a Values from (Astegno et al., 2016).



Fig. 5. Effect of the AtPCaP1 peptides on Ca²⁺ dissociation kinetics from AtCaM1. Time course of Quin-2-induced Ca²⁺ dissociation from: AtCaM1 in the absence (red) and presence of myr-AtPCaP1p (**A**) and AtPCaP1p (**D**); the N-lobe of AtCaM1 (B34Q variant) in the presence of myr-AtPCaP1p (**B**) or AtPCaP1p (**E**); the C-lobe of AtCaM1 (B12Q variant) in the presence of myr-AtPCaP1p (**C**) or AtPCaP1p (**F**).

with PM, but it also possesses a 1-8-14 CaM binding motif which is a common recognition motif for Ca²⁺-dependent CaM interactions (Fig. 1A). At physiological Ca²⁺ concentrations, the association of PCaP1 with the PM has been shown to be very stable, and not affected by CaM (Nagasaki et al., 2008). Elevated, non-physiological Ca²⁺ levels have been shown to cause the partial dissociation of AtPCaP1 from the membrane to the cytosol (Li et al., 2011; Kato et al., 2010; Nagasaki et al., 2008), while for the *M. truncatula* DREPP Ca²⁺ treatment caused a relocalization of the protein into nanodomains that colocalize with microtubules (Su et al., 2020), indicating that cytoplasmic Ca^{2+} levels are key regulators of the association of DREPP family proteins to the PM. Indeed AtPCaP1, as well as its paralogue AtPCaP2, interacts with Ca²⁺-CaM, and this interaction competitively regulates the binding to PtdInsPs (Li et al., 2011; Kato et al., 2010; Nagasaki et al., 2008), thus adding a further level of sophistication in the fine-tuning of AtPCaP1 associations. However, many questions remain to be answered, such as the definition of the molecular determinants of AtPCaP1-Ca²⁺/CaM interaction, the functional impact of the myristoyl group linked to AtPCaP1 on the complex formation and on the Ca²⁺ sensitivity of CaM.

In this work, we investigated the mutual influence of these aspects on the structural and functional properties of the AtPCaP1-AtCaM1 complex formation. We applied an in vitro approach that, albeit focused on interactions in which the binding partner AtPCaP1 is reduced to a relatively short peptide, offered valuable insights that enhance and complement other strategies described in the literature (Nagata et al., 2016; Giovannoni et al., 2021; Ide et al., 2007; Vallet et al., 2023; Kato et al., 2013; Tanaka-Takada et al., 2019; Nagasaki-Takeuchi et al., 2008) aimed at unraveling the details of the AtPCaP1- Ca²⁺- AtCaM1 interaction network within plant cells. Notably, although the use of a peptide fragment rather than a whole protein as a AtCaM1 target may result in differences in the Ca²⁺-binding affinity of AtCaM1, our ITC studies failed to detect differences in affinity and stoichiometry when AtCaM1 was in complex with either the peptide corresponding to the CaM-binding domain (the first 26 N-terminal residues) of AtPCaP1 ($K_d = 911$ nM) or full-length AtPCaP1 ($K_d = 859$ nM), suggesting that our approach would give representative results and satisfyingly recapitulates the behavior of the entire protein.

We demonstrated that the interaction between AtPCaP1 and AtCaM1 is strictly Ca^{2+} dependent and the myristoylation is directly involved in

the association of the two proteins. These findings provide direct support for the proposal of Matsubara and colleagues (Matsubara et al., 2003; Hayashi et al., 2002) that protein myristoylation plays important roles not only in the interaction with membrane phospholipids but also in the protein-protein interaction with Ca^{2+} -CaM. In recent years, the interest in the role of myristoylation in plant proteins has increased significantly, although the correlation between the presence of the myristoyl groups and specific functions has been demonstrated only for few proteins, including the Arabidopsis Calcineurin B-like protein 4 (Held et al., 2011), the small G-protein Ara 6 (Yin et al., 2017), and the Pto Kinase protein of Tomato (Andriotis et al., 2006).

Our results, by showing that the myr-AtPCaP1 peptide has an affinity to AtCaM1 about 14-fold higher ($K_d \sim 60$ nM) than that of the nonmyristoylated peptide, suggest that the myristoylation might modulate the mechanism of recognition of AtPCaP1 by AtCaM1 and consequently alter the kinetics of complex formation. Interestingly, several proteins that contain myristoylated polybasic N-terminal sequences, including the brain acid soluble protein 1 (CAP23/NAP22) (Takasaki et al., 1999), the viral proteins negative factor (Nef) from HIV1 (Hayashi et al., 2002), and the oncogenic pp60^{v-src} tyrosine-protein kinase from Rous sarcoma virus (Hayashi et al., 2004) have been reported to bind Ca²⁺-CaM.

In the CaM-binding site of these proteins the CaM recognition motif (such as the 1-8-14, 1-5-10, or IQ motif) is absent. As a consequence, binding of these targets to CaM is dependent on the presence of the myristoyl, which, in a linear conformation, occupies the hydrophobic channel formed by the lobes of CaM, while the polybasic amino acid cluster further stabilizes the interaction by contacting the acidic surface of CaM (Grant et al., 2020). In the case of AtPCaP1, the presence of both the myristoyl moiety and a canonical CaM binding motif, in addition to the polybasic cluster (Fig. S6), suggests an additional level of complexity in the mechanism of target recognition by CaM. Our NMR and CD analyses support this conclusion, confirming that the myristoylated AtP-CaP1 peptide can bind Ca²⁺-AtCaM1 with 1:1 stoichiometry and high affinity. The overall structure of Ca²⁺-AtCaM1 changes markedly in the presence of the myristoylated AtPCaP1p, and the peptide adopts a helical conformation in the final complex. On the other hand, the binding of AtCaM1 to the non-myristoylated peptide involves mainly local conformational changes at the C-lobe of AtCaM1.

Notably, we observed that the formation of such complexes

significantly increases the AtCaM1 affinity for Ca²⁺. It is well-known that a fine-tuned equilibrium between $[Ca^{2+}]$, CaM, and its target(s) is crucial for the proper biological function, even if it is still unclear how regulation of CaM's affinity for Ca²⁺ by target proteins is achieved at the molecular level. Interestingly, a 90-fold and 18-fold increase in affinity for Ca²⁺ of AtCaM1 were measured in the presence of the myristoylated and non-myristoylated peptides, respectively, indicating that the myristoylation modulates the Ca^{2+} sensitivity of AtCaM1. Significant changes were observed in all the macroscopic binding constants for the four functional EF-hands of AtCaM1 upon the addition of the myristoylated peptide, while in the case of the non-myristoylated one, the most notable changes were observed for the high-affinity sites, which have been previously assigned to the C-lobe of AtCaM1. These results are nicely supported by the NMR data, suggesting that for the myr-AtPCaP1p, the complex formation is mediated by both AtCaM1 lobes. while for the AtPCaP1p, it is mainly mediated by the C-lobe. Measurements of the Ca²⁺ dissociation rates from both N- and C-lobes of AtCaM1 showed that Ca²⁺ dissociation is remarkably slower in the presence of AtPCaP1 peptides. However, a more significant impact of the myristoyl moiety on the Ca^{2+} dissociation from the N-lobe of AtCaM1 was observed, which is likely the cause of the enhanced Ca^{2+} affinity (\cong 5fold) obtained for the complex in the presence of the myristoylated peptide.

AtPCaP1 regulates many important processes in plant life (Nagata et al., 2016; Qin et al., 2012; Tanaka-Takada et al., 2019), viral cell-to-cell movement (Vijayapalani et al., 2012) and immunity (Giovannoni et al., 2021; Mattei et al., 2016). Interestingly, several studies pointed out the role of N-myristoylation in the immune response against microbial and viral infections (Udenwobele et al., 2017; Wang et al., 2021).

It should also be remembered that AtPCaP1 can bind PtdInsPs which are major components of intracellular signaling (Kato et al., 2010). Thus, the interaction between the Ca²⁺-CaM and myristoylated AtPCaP1 represents an important connection between major signaling pathways in plant cells. In this context, our findings allow us to add novel details to the mechanism whereby PCaP proteins act as molecular switches of the Ca^{2+} signaling mediated by PPIs and CaM (Kato et al., 2010). It is tempting to speculate that at resting conditions, AtPCaP1 is anchored to the PM via N-myristoylation and a polybasic cluster. When the Ca²⁺ concentration increases, the binding to $\mbox{Ca}^{2+}\mbox{-CaM}$ stabilizes a suitable helix conformation of the N-terminal region of PCaP1 that would favor myristoyl exposure and subsequent insertion into the hydrophobic clefts of Ca^{2+} -CaM. The formation of the final complex decreases the Ca^{2+} dissociation rate from AtCaM1, thus ensuring an extended lifetime of the activated protein complex long after the [Ca²⁺] transient has decreased. In parallel, the interaction of AtPCaP1 with Ca²⁺-CaM stimulates the release of PtdInsPs, which may then participate in downstream-specific signaling.

AtPCAP1 is an emerging important player in the cellular Ca^{2+} signaling network with multifaceted physiological roles in plant development and immunity. Our data can help to better understand the molecular mechanisms of regulation of these processes by AtPCaP1 and of signal integration by Ca²⁺-AtCaM1.

CRediT authorship contribution statement

Marco Pedretti: Conceptualization, Formal analysis, Investigation. Filippo Favretto: Formal analysis, Investigation. Francesca Troilo: Formal analysis, Investigation. Moira Giovannoni: Writing – review & editing. Carolina Conter: Investigation. Benedetta Mattei: Writing – review & editing, Funding acquisition. Paola Dominici: Writing – review & editing. Carlo Travaglini-Allocatelli: Investigation, Writing – review & editing. Adele Di Matteo: Conceptualization, Validation, Writing – original draft, Supervision, Project administration, Funding acquisition. Alessandra Astegno: Conceptualization, Validation, Writing – original draft, Supervision, Funding acquisition.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Data availability

Data will be made available on request.

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Appendix A. Supplementary data

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