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Calmodulin binds to Drosophila TRP with an unexpected mode

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Running title: An unexpected binding mode between CaM and TRP

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Abstract

Drosophila TRP is a calcium-permeable cation channel essential for fly visual signal transduction. During phototransduction, Ca^{2+} mediates both positive and negative feedback regulation on TRP channel activity, possibly via binding to calmodulin (CaM). However, the molecular mechanism underlying Ca^{2+} modulated CaM/TRP interaction is poorly understood. Here, we discover an unexpected, Ca^{2+} -dependent binding mode between CaM and TRP. The TRP tail contains two CaM binding sites (CBS1 and CBS2) separated by a ~70-residue linker. CBS1 binds to the CaM N-lobe and CBS2 recognizes the CaM C-lobe, so each TRP only binds to one CaM. Mutations introduced in both CBS1 and CBS2 completely eliminated CaM binding in full length TRP, but surprisingly had no effect on the response to light under physiological conditions, suggesting alternative mechanisms governing Ca^{2+} -mediated feedback on the channel activity. Finally, we discover that TRPC4, the closest mammalian paralog of *Drosophila* TRP, adopts a similar CaM binding mode.

INTRODUCTION

Transient receptor potential channels (TRP channels) are extensively expressed in different species of animals and involved in diverse physiological processes such as responding to light, pressure, pain, taste, temperature, and other stimuli (Clapham, 2003; Montell, 2005). Based on their sequence similarities and functional properties, TRP channels in mammals can be classified into 7 subfamilies: the canonical TRP channels (TRPCs), the melastatin TRP channels (TRPMs), the vanilloid TRP channels (TRPVs), the polycystin channels (TRPPs), the ankyrin transmembrane protein 1 channels (TRPA), the mucolipin channels (TRPML) and mechanosensing TRPN channels (*Drosophila* NOMPC) (Li, 2017; Montell, 2005).

The archetype TRP channel was discovered ~30 years ago following investigations of a spontaneous *Drosophila* mutant which had abnormal electroretinogram during prolonged intense light stimulation (Cosens and Manning, 1969). The *Drosophila* TRP (referred to as TRP from hereon) protein was later shown to be a Ca²⁺ permeable cation channel essential for excitation and light adaptation in *Drosophila* phototransduction (Hardie and Minke, 1992; Minke and Selinger, 1996; Montell and Rubin, 1989; Suss-Toby et al., 1991). Underneath the plasma membranes of rhabdomere in *Drosophila* photoreceptor cells, TRP together with eye-specific protein kinase C (ePKC) and phospholipase C β (NORPA) are assembled by a master scaffold protein called inactivation no after potential D (INAD) into a large and stoichiometric supra-molecular assembly termed the signalplex or transducisome (Chevesich et al., 1997; Hardie and Raghu, 2001; Montell, 2012; Tsunoda et al., 1997; Ye et al., 2018; Ye et al., 2016).

As well as being a Ca^{2+} -permeable channel, the *Drosophila* TRP is also known to be regulated by Ca^{2+} ions (Hardie, 1991, 1995; Scott et al., 1997) and Ca^{2+} influx via the channels can exert both positive and negative feedback regulation on TRP activities. In the presence of physiological Ca^{2+} concentration, TRP can be activated and terminated with extremely fast kinetics (Hardie, 1991, 1995; Reuss et al., 1997).

 Ca^{2+} influx also mediates light adaptation (Gu et al. 2005) and is critical for dark noise threshold maintenance in Drosophila photoreceptors (Chu et al., 2013; Katz and Minke, 2012). However, the molecular mechanisms by which Ca^{2+} regulates TRP activities are largely unknown. At least two potential non-exclusive mechanisms exist. First, Ca²⁺ may directly bind to TRP to regulate channel activities, as in the case recently demonstrated for mammalian TRPMs (Wang et al., 2018; Yin et al., 2019). Second, Ca²⁺ might indirectly regulate channel activity by modulating the interaction between TRP and calmodulin (CaM) (Chevesich et al., 1997; Sun et al., 2018; Tang et al., 2001). CaM, as a principal Ca^{2+} signal decoder in all eukaryotic cells, is known to bind to and regulate activities of various ion channels including voltage-gated K⁺ channels (Chang et al., 2018), Na⁺ channels (Herzog et al., 2003), Ca²⁺ channels (Qin et al., 1999), etc. For the TRP superfamily ion channels, CaM also serves as a channel regulator with versatile functions (Zhu, 2005). For example, CaM is critical for Ca²⁺-mediated termination of TRPV5 and TRPV6 activity (Hughes et al., 2018; Singh et al., 2018). On the contrary, CaM seems to act as an activator of TRPC4 and TRPC5 (Ordaz et al., 2005; Otsuguro et al., 2008). In Drosophila compound eyes, reduction of CaM can markedly impair the termination process of phototransduction (Liu et al., 2008; Porter et al., 1995; Scott et al., 1997). CaM was demonstrated to directly bind to a large fragment of the TRP tail (aa L683-A976) using a CaM-overlay assay (Chevesich et al., 1997). However, the mechanisms for CaM to regulate TRP is poorly understood. Currently, there is no structural study on the binding of CaM to TRP.

In this study, we characterized the interaction between CaM and the C-terminal tail of TRP in detail. We demonstrated that the TRP tail contains two CaM binding sites (CBSs) separated by a flexible linker with more than 70 amino acid residues. Unexpectedly, we found that the TRP tail containing both CBSs binds to one molecule of Ca²⁺-CaM with its CBS1 engaging the CaM N-lobe and CBS2 binding to the CaM C-lobe. Lys75 in the N-lobe of CaM plays a critical role in determining the specificity of the two CBS sites for binding to N- and C-lobes of CaM. Guided by the

biochemical binding mechanism and the structure of the TRP/CaM complex uncovered in this study, we searched for potential binding between CaM and mammalian TRPCs, and found that the mouse TRPC4 C-terminal tail binds to Ca^{2+} -CaM in a mode similar to that between CaM and *Drosophila* TRP. Together, our discoveries serve as a structural framework for future investigations of TRP channel activity regulation by Ca^{2+} and CaM.

RESULTS

Characterization of the interaction between *Drosophila* TRP C-terminal tail and CaM

We began our study by performing a detailed biochemical characterization of the interaction between CaM and the TRP tail. The tail of TRP begins with residue S717 and ends at residue L1275 (Figure 1A). Searching the Calmodulin Target Database (http://calcium.uhnres.utoronto.ca/ctdb/no_flash.html) (Yap et al., 2000) showed that the N-terminal part of the TRP tail (aa 717-940) contains several potential CaM binding sites, and the rest of the tail (aa 940-1275) does not contain signs of CaM binding sequence. Consistently, the C-terminal part of the tail (aa 940-1275, purified as a thioredoxin (Trx)-tagged fusion protein, had no detectable binding to CaM either in the presence or absence of Ca²⁺ (Figure 1B). In contrast, the N-terminal part of the TRP tail (aa 717-940) specifically bound to Ca²⁺-CaM based on analytical gel filtration chromatography coupled with static light scattering (SLS). Using both Trx-tagged and tag-removed proteins, we calculated that TRP (aa 717-940) bound to Ca²⁺-CaM with a 1:2 stoichiometry (Figure 1C), indicating that there are at least two CaM binding sites within the fragment.

We further divided the 717-940 fragment into two. Interestingly, the fragment containing as 783-940 (we chose to start the fragment from N783 as this residue marks the beginning of non-conserved regions among the TRP channels following the

coiled-coil domain; Figure 1F) was found to bind to CaM in a Ca^{2+} -dependent manner, and this fragment (referred to as CBS-B in Figure 1A) binds to Ca²⁺-CaM with a 1:1 stoichiometry (Figure 1D). The TRP fragment containing 717-783 could not be expressed in soluble forms alone or in complex with CaM in bacterial cells. We therefore used a synthetic peptide to test whether a certain segment in this region may bind to CaM. As was shown in an earlier study, a synthetic peptide corresponding aa 728-754 of TRP bound to Ca²⁺-CaM with a Kd of $3.75\pm0.62 \mu$ M (Tang et al., 2001) (Figure 1E), and we refer this CaM binding site of TRP as CBS-A (Figure 1A). The residues corresponding to CBS-A largely overlap with the entire connecting helix in the closed state structures of TRPC3, TRPC4, TRPC5 and TRPC6 (Duan et al., 2018; Duan et al., 2019; Tang et al., 2018) (Figure 1F). The residues corresponding to aa 728-754 of TRP are predicted to form a single connecting helix that is completely buried in the structures of TRPCs (Figure 1G). Therefore, it is unlikely that the fragment corresponding to the CBS-A site of TRP is accessible to CaM in the closed state full-length channel. Thus, the interaction between CBS-A peptide and CaM may originate from an artifact of the TRP tail fragmentation from the intact channel. Indeed, we demonstrated that the CBS-A site in the full-length endogenous TRP is not involved in binding to CaM (see data in Figure 4 below). However, we cannot exclude the possibility that CBS-A may function as a CaM binding site of TRP under certain conditions when this part of TRP gets exposed. The above biochemical study, together with structure-based sequence analysis of TRP and TRPC channels, establishes that the TRP tail fragment encompassing residues 783-940 contains the functional CaM binding element.

The TRP tail contains two discrete CaM binding sites acting together to bind one CaM in a Ca²⁺-dependent manner

Detailed sequence analysis of the 783-940 fragment of the TRP tail revealed two potential CaM binding sites (CBSs) with sequences conserved in insects, though not

in mammalian TRPCs (Figure 2A). These two potential CBS sequences are defined as CBS1 (aa T802-K862) and CBS2 (aa M899-D940), which we used for the following experiments, unless specified otherwise. We used purified Trx-fused CBS1 and CBS2 to test whether the predicted CBS fragments may indeed bind to CaM. Isothermal titration calorimetry (ITC)-based assays showed that CBS1 binds to Ca²⁺-CaM with a K_d of 0.35±0.09 μM, and with a 1:1 stoichiometry (Figure 2B1). CBS2 also binds to Ca²⁺-CaM with a 1:1 stoichiometry and a K_d of 0.25±0.03 µM (Figure 2B2). Neither CBS1 nor CBS2 showed detectable binding to apo-CaM (titration curves in red in Figure 2B). The data in Figure 2B reveal that both CBS1 and CBS2 can bind to CaM in a Ca²⁺-dependent manner and with quite strong binding affinities. Unexpectedly, the analytical gel filtration chromatography coupled with static light scattering (SLS) analysis (Figure 1D), and analytical ultracentrifugation (AUC) sedimentation velocity analysis (Figure 2C) showed that a large fragment of TRP tail encompassing both CBS1 and CBS2 (aa N783-D940) formed a stable complex with Ca²⁺-CaM with a 1:1 stoichiometry. In fact, the TRP(783-940)/Ca²⁺-CaM complex could only be obtained by co-expressing the two proteins together in bacteria cells. Removal of Ca^{2+} from the complex by addition of an excess amount of EDTA led to dissociation of TRP(783-940) from CaM. Moreover, the dissociated TRP(783-940) precipitated due to its extremely low solubility (data not shown).

There are two possible explanations for the above observations. CBS1 and CBS2 may compete with each other for binding to CaM, so only one of the two CBS sites engages CaM in the TRP(783-940)/Ca²⁺-CaM complex. Alternatively, CBS1 and CBS2 may simultaneously interact with one molecule of CaM, thus forming the 1:1 TRP(783-940)/Ca²⁺-CaM complex. To differentiate between these two possibilities, we mixed CBS1, CBS2 and CaM at a 1:1:1 molar ratio and subjected the mixture to analytical gel-filtration coupled with SLS analysis. The CBS1/CBS2/CaM mixture eluted as a single peak with a volume smaller than the elution volume of the CBS1/CaM complex or the CBS2/CaM complex (Figure 2D, top). SDS-PAGE

analysis of the fractions of the elution peak of the CBS1/CBS2/CaM mixture also showed that CaM formed a complex simultaneously with Trx-CBS1 and Trx-CBS2 (Figure 2D, bottom). Taken together, the above analysis suggested that a stable triple complex was formed when CBS1, CBS2 and CaM were mixed at a 1:1:1 molar ratio (i.e. both CBS1 and CBS2 bind to the same Ca²⁺-CaM moiety).

To further understand the molecular mechanism governing the interaction between CBS12 and CaM, we resorted to NMR spectroscopic studies. We compared the ¹H-¹⁵N HSQC spectrum of Ca²⁺-CaM co-expressed with CBS12 complex with the spectrum of Ca²⁺-CaM and found that residues from both the N- and C-lobes of Ca²⁺-CaM underwent significant CBS12 binding-induced chemical shift changes (Figure 2E, and Figure S1A&B for the whole spectra). These data indicate that both lobes of CaM are involved in the interaction with CBS12. To further dissect the interaction, we overlaid the ¹H-¹⁵N HSQC spectrum of the ¹⁵N-Ca²⁺-CaM/CBS12 complex with that of the ¹⁵N-Ca²⁺-CaM/¹⁴N-CBS1 complex or with that of the ¹⁵N-Ca²⁺-CaM/¹⁴N-CBS2 complex. The ¹H-¹⁵N HSQC spectrum of the Ca²⁺-CaM in Ca²⁺-CaM/CBS12 complex roughly overlapped with the summed spectra of ¹⁵N-Ca²⁺-CaM/¹⁴N-CBS1 and ¹⁵N-Ca²⁺-CaM/¹⁴N-CBS2 (Figure 2E and Figure S1B), suggesting that CBS1 and CBS2 bind to distinct lobes of Ca²⁺-CaM. We also compared the ¹H-¹⁵N HSQC spectrum of ¹⁵N-labelled Ca²⁺-CaM with that of ¹⁵N-labelled Ca²⁺-CaM in complex with unlabeled CBS1 or CBS2 (Figure S1A). The binding of CBS1 induced large chemical shift changes to the N-lobe of Ca²⁺-CaM, but only caused relatively small shift changes to the C-lobe of CaM. Conversely, binding of CBS2 induced large chemical shift changes to the C-lobe of Ca²⁺-CaM, but chemical shift changes to the N-lobe of CaM was small (see the signature Gly residues from each EF-hand of Ca²⁺-CaM in the zoomed in region in Figure S1A). The above NMR analysis suggested that CBS1 and CBS2 specifically bind to the N-lobe and C-lobe of Ca²⁺-CaM, respectively, such that Ca²⁺-CaM and CBS12 form a stable 1:1 complex. To confirm the conclusion derived from the NMR-based study,

we generated CaM derivatives in which we mutated Ca^{2+} -binding sites in the N-lobe or the C-lobe by substituting the last Glu in EF-hands 1&2 with Gln (i.e. E31/67Q) or EF-hands 3&4 with Gln (i.e. E104/140Q). The E31/67Q-CaM lost its binding to CBS1 but retained its binding to CBS2. Conversely, the E104/140Q-CaM lost its CBS2 binding but retained CBS1 binding (Figure S2). Taken together, we formulate an interaction model between TRP(783-940) and Ca²⁺-CaM (Figure 2F). In this model, CBS1 and CBS2 selectively bind to the N-lobe and C-lobe of Ca²⁺-CaM, respectively, forming a stable 1:1 complex. It is noteworthy that the complete CaM binding region of the TRP tail spans a total of ~140 amino acid residues (Figure 2A).

Finally, we asked whether CBS12 might be able to bind to Ca^{2+} -CaM with a higher affinity due to potential synergistic actions of CBS1 and CBS2. Since we could not obtain isolated CBS12 for ITC-based assay, we used analytical ultracentrifugation (AUC) sedimentation equilibrium to measure the dissociation constant between CBS12 and CaM (Figure 2G). The K_d value (0.10±0.01 µM) derived from AUC for the Ca²⁺-CaM/CBS12 complex is comparable to that of the Ca²⁺-CaM/CBS1complex or the Ca²⁺-CaM/CBS2 complex (Figure 2B), indicating that there is very little conformational coupling (or synergism) between CBS1 and CBS2 in binding to CaM.

Structural Characterization of the CaM/TRP CBS12 Complex

We attempted to uncover the detailed molecular mechanism governing the unexpected binding between TRP tail and CaM by determining the crystal structure of the CBS12/Ca²⁺-CaM complex. Despite extensive trials, we could not crystallize the complex, likely due to the conformational flexibility between the two lobes of CBS12-bound Ca²⁺-CaM (Figure 2F and the NMR data in Figure 2E and Figure S1). Since CBS1 and CBS2 independently bind to the N- and C-lobes of Ca²⁺-CaM, we decided to crystallize the CBS1/N-lobe_CaM and CBS2/C-lobe_CaM complexes separately. We were able to determine the structures of CBS1/N-lobe_CaM complex

and CBS2/C-lobe_CaM complex at resolutions of 1.78Å and 2.15Å, respectively (Figure 3A and Figure S3 and Table 1). For easy viewing, we connected the two lobes of CaM by a dotted line representing the flexible central linker connecting α D and α E of Ca²⁺-CaM.

Both lobes of CaM adopt an open conformation with a Ca^{2+} ion occupying each EF-hand (Zhang et al., 1995). Interestingly, CBS1 forms a "helix-turn-helix" structure interacting with the hydrophobic surface of N_lobe_CaM. The first helix (a1) spans Q816-M827 and the second helix (a2) encompasses I836-G845. The residues connecting the $\alpha 1$ and $\alpha 2$ are not defined in the structure. Both helices are amphipathic in nature (Figure 2A). The hydrophobic residues from α 2 make extensive contacts with CaM and the side chain of F837 from a2 functions as the anchoring residue inserting into the deep pocket of N-lobe_CaM (Figure 3B). The hydrophobic residues from $\alpha 1$ of CBS1 also interact with CaM and with $\alpha 2$. The formation of a helix-turn-helix conformation of CBS1 while binding to one lobe of Ca²⁺-CaM is rather unusual among the numerous CaM/target complexes determined to date (Hoeflich and Ikura, 2002; Tidow and Nissen, 2013). This also explains why an elongated fragment (>30 amino acid residues) is required for CBS1 to bind to the N-lobe_CaM. It is worth noting that the binding mode between the C-lobe of Ca²⁺-CaM to a variant of the SK2 K⁺-channel tail resembles the CBS1/N-lobe CaM structure in this study, because the SK2 peptide also forms a helix-turn-helix conformation (Zhang et al., 2012). However, the first helix (corresponding to al in CBS1) in the SK2 channel does not make direct contact with CaM, whereas al in CBS1 interacts with CaM directly (Figure 3B).

CBS2 in the complex with the C-lobe_CaM forms a single α -helix (α 3) spanning A917-K929 (Figure 2A and Figure 3C). Like most of other Ca²⁺-CaM binding helical peptides (Tidow and Nissen, 2013), CBS2 α 3 is also a positively charged amphipathic α -helix (Figure 2A). The hydrophobic residues of α 3 contact the hydrophobic surface of the C-lobe_CaM and the aromatic ring of F925 from α 3 inserts into the deep pocket

of C-lobe_CaM (Figure 3C).

The N/C-lobes of CaM bind to TRP CBS1 and CBS2 respectively with high specificity, although the two lobes share very high structural similarities. An analysis of the two structures determined here offers the molecular basis for such binding specificity. Lys75 in the N-lobe of CaM interacts with S838 in CBS1 (Figure 3D1). The residue corresponding to Lys75 in the C-lobe of CaM is the last residue of the proteins and does not form α -helix, which may partly explain why CBS1 prefers to bind to CaM N-lobe. We next modeled the CBS2 peptide onto CaM N-lobe by superimposing the N-lobe and C-lobe structures of CaM shown in Figure 3A. In this modeled structure, Lys924 from CBS2 physically crashes into Lys75 from CaM (Figure 3D2). Thus, CBS2 is unfit to bind to the CaM N-lobe. In line with this analysis, when Lys924 of CBS2 is substituted with Ala, the mutant CBS2 became to be a CaM N-lobe binder (Kd: 7.94±0.88 μ M, Figure S2C). Thus, the selectivity of CBS2 for the C-lobe concomitantly determines the binding of CBS1 to the N-lobe of CaM.

Collectively, the structures of the CBS1/N-lobe_CaM and CBS2/C-lobe_CaM complexes, together with our detailed biochemical studies, demonstrate that two discrete fragments in the tail of TRP separated by a flexible linker of >70 residues simultaneously and non-synergistically bind to the two lobes of Ca²⁺-CaM forming a stable complex with a 1:1 stoichiometry. Amino acid sequence analysis of TRP CBS12 from different insect species revealed that residues involved in the CaM binding are highly conserved (Figure 2A).

We generated a series of CBS1 and CBS2 mutants to validate the structures of their respective complexes with N- and C-lobes of CaM (Figure 3E and Figure S4). We used ITC assays to measure the binding of each of these mutants to CaM. Replacing hydrophobic residues in the α 1 helix of CBS1 (F819 and V820 together or L823 alone) with Ala significantly weakened the CBS1/CaM binding (Figure 3E), supporting our structure-based analysis showing the role of α 1 in CBS1's binding to

CaM. As expected, replacing hydrophobic residues in the α2 helix of CBS1 with Ala also invariably weakened its binding to CaM (Figure 3E). In particular, when the anchoring F837 and the neighboring I836 were both replaced by Ala, the mutated CBS1 showed no detectable binding to CaM (Figure S4A4). Similarly, replacing the anchoring F925 together with its neighboring M926 with Ala totally abolished CBS2's binding to CaM (Figure S4B4). Finally, the IF836,837AA and FM925,926AA quadruple mutant of CBS12 exhibited no detectable binding to CaM (Figure 3F).

Ca²⁺-dependent interaction between TRP and CaM

A previous study showed that CaM can interact in the presence or absence of Ca²⁺ with the C-terminal tail of TRP, as well as TRP extracted from *Drosophila* head lysate (Chevesich et al., 1997), a finding that is different from our current in vitro study using the TRP tail fragment. It is possible that the full-length TRP may contain additional CaM binding site(s) or that the tail of the channel may adopt a different conformation in the context of the full-length protein thereby having different CaM binding properties from the fragmented tail purified in our in vitro study. To resolve this issue, we attempted to purify TRP from *Drosophila* head lysates for CaM binding assay. We took advantage of our previous finding that the NORPA C-terminal domain (NORPA_CT, aa E863-A1095), which contains coiled-coil and PDZ binding motif of the NORPA protein, binds to INAD with a Kd ~10 nM (Ye et al., 2018). Therefore, we used NORPA_CT as an affinity tag to purify the INAD organized INAD/TRP/ePKC complex from Drosophila head lysates. We then eluted the INAD associated TRP with a GB1-tagged 15-residue peptide corresponding to the last 15 residues of TRP (TRP_CT, aa R1261-L1275), which was shown to be responsible for the specific interaction between TRP and INAD PDZ3 (Ye et al., 2016). Using this method, we purified sufficient amount of highly homogeneous TRP from Drosophila heads for direct CaM binding assays. GST-CaM pull-down assays using the purified TRP from wild-type flies showed that, in the presence of Ca^{2+} , TRP robustly interacted with CaM (Figure 4A, *left*). Chelating of Ca^{2+} by EDTA totally eliminated the interaction between TRP and CaM (Figure 4A, *left*).

To validate whether the CaM binding sites within CBS12 delineated from our *in vitro* biochemical study are indeed responsible for the TRP/CaM interaction, we compared the binding of CaM to the wild-type full-length TRP (TRP-WT) or to TRP containing the IF836,837AA and FM925,926AA quadruple mutations in CBS12 (Figure 3E). No detectable binding between CaM and TRP with the quadruple mutation purified from transgenic fly heads was observed, either in the presence or absence of Ca²⁺ (Figure 4A, *right*). The complete loss of CaM binding of the quadruple mutant of TRP indicated that CBS12 is solely responsible for TRP to bind to CaM. This result also supports our earlier analysis showing that the CBS-A site corresponding to the connecting helix of the channel (Figure 1) is not likely a functional CaM binding site in the full-length TRP.

Amino acid sequence analysis indicated that the CaM binding tail and the tetramer forming coiled coil domain of TRP are separated by a relatively short stretch of connecting residues (Figures 1F and 2A). We next tested whether CaM binding may promote polymerization of TRP. We created a homotetramer of TRP CBS12 by fusing a tetrameric leucine zipper GCN4 coiled-coil domain (Harbury et al., 1993) to aa 783-940 of TRP, so that the GCN4-CBS12 fusion protein resembles the actual tail assembly mode of the full-length TRP (see Figure 1F&G) (Duan et al., 2018; Tang et al., 2018). The GCN4-CBS12 and Trx-CaM complex could be purified to high homogeneity. AUC sedimentation velocity assay showed that the GCN4-CBS12/Trx-CaM complex formed a single peak with a molecular mass of ~225 kD (Figure 4B), corresponding to a 4:4 molar ratio complex. Negative staining electron microscopy further showed that addition of CaM, either in the apo- or in the Ca2+-saturated forms, to the full-length TRP purified from Drosophila heads did not change the oligomerization state of the channels (Figure S5). Thus, CaM binding is not likely to cause multimerization of TRP tetramers via its C-terminal tail.

Combining the above biochemical and structural results, we built an interaction model between TRP C-terminal tail and CaM by modeling the TRP CBS1/N-lobe complex and CBS2/C-lobe complex crystal structures onto the cryo-EM structure of mouse TRPC4 (mTRPC4, PDB: 5Z96) (Duan et al., 2018) (Figure 4C&D). We believe that the most likely mode of the interaction is that each Ca²⁺-CaM binds to CBS1 and CBS2 with the same subunit of the TRP tetrameric tail (Figure 4C), as we could only detect a stable 1:1 Ca²⁺-CaM/CBS12 complex in our study (Figure 1D&2C). Since CBS1 and CBS2 are connected by a flexible linker of >70 residues, it is nonetheless possible that the two lobes of CaM may bind to CBS1 and CBS2 from two neighboring TRP tails forming an inter-subunit cross-connected TRP tail assembly (Figure 4D). Further work will be required to differentiate which mode might be adopted by the full-length TRP.

CaM binding to the tail does not directly modulate TRP channel activity under physiological conditions

We next asked whether Ca^{2+} -dependent binding of CaM to the C-terminal tail of TRP plays a role in the channel activity of TRP during phototransduction by recording whole-cell voltage clamped responses to light in photoreceptors from dissociated ommatidia. In photoreceptors expressing wild-type TRP channels brief light flashes delivered in normal bath (containing 1.5 mM Ca²⁺) elicit rapid responses that peak within ~40 ms and return to baseline after ~100 ms. As previously reported (e.g. Hardie, 1991; Reuss et al, 1997; Henderson et al, 2000) the rapid kinetics are dependent upon sequential positive and negative feedback mediated by Ca²⁺ influx via the TRP channels, and in the absence of extracellular Ca²⁺ (0 Ca²⁺, 1 mM EGTA) kinetics of both excitation and inactivation are slowed ~10-fold (Figure 5A).

We generated transgenic flies expressing TRP with double Ala substitutions in both CBS1 (IF836,837AA) and CBS2 (FM925,926AA) respectively, as well as a quadruple mutant of TRP with both CBS1 and CBS2 disrupted. Although our biochemical assays showed that these mutations render CaM binding to TRP undetectable, amplification, rapid kinetics and their dependence upon Ca²⁺ appeared essentially unaffected in any of these mutants (Figure 5A-D, quantified in Figure 5E). These surprising results suggest that the Ca²⁺ dependence of TRP channel activity in phototransduction must be mediated by another mechanism(s). Nevertheless, upon closer inspection, a subtle but consistent phenotype was detected in all three mutants. Thus, whilst light response amplitudes in physiological calcium were similar to those of TRP-WT, in Ca²⁺ free bath they were ~3 times larger than responses observed in TRP-WT under the same conditions (Figure 5A-D, quantified in Figure 5F). It appears that, without Ca²⁺-CaM restriction, the open probability of TRP somehow can increase. This extracellular Ca²⁺ independent effect of the CBS mutations is not readily understood but may imply some indirect (allosteric) effect of this region on channel open probability. Future studies will be required test this possibility.

TRPC4 binds to Ca²⁺-CaM with a similar mode as TRP does

Among the TRP subfamilies, TRPCs share the highest sequence similarity to TRP (Montell, 2005; Ramsey et al., 2006). Within the TRPC subfamily, TRPC4 and TRPC5 are most closely related to TRP, and both are also activated by the phospholipase C-mediated signaling cascade. Recently, TRPC3, TRPC4, TRPC5 and TRPC6 cryo-EM structures have been determined (Duan et al., 2018; Duan et al., 2019; Tang et al., 2018), and all of them share similar conformations in their structured regions observable by cryo-EM. Each TRPC channel contains N-terminal ankyrin repeats followed by six transmembrane helices. TRPC C-terminal tails contain a common TRP box, a connecting helix, and a coiled-coil domain, followed by a stretch of disordered sequences with very different lengths (Figure 6A, also see Figure 1F&G). Regrettably, the very C-terminal tails after coiled-coil domain of the TRPC channels are invisible in the cryo-EM structures. Interestingly, previous studies

demonstrated that the C-terminal tail of TRPC4 also contains 2 CBSs (CBS1 aa D785-N812 and CBS2 aa E827-E854 corresponding to the mouse TRPC4 sequence) (Tang et al., 2001; Trost et al., 2001), while TRPC5 only contains 1 CBS (aa K829-S858) (Ordaz et al., 2005). To test whether the Ca²⁺-CaM interaction mode found in the *Drosophila* TRP tail might have been adopted by mammalian TRPC members, we investigated the binding of CaM to each of the TRPC tails.

First, we assayed the binding of CaM to each member of the TRPC subfamily (Figure 6A) by GST pull down assay. Each of the TRPC tails used for the binding assays contains the coiled-coil domain to the C-terminal end. GST-CaM was used to pull down GFP-TRPC tail expressed in HEK293T cells. This pull-down assay showed that, among the seven TRPC tails, only the TRPC4 and TRPC5 showed positive Ca²⁺-CaM binding (Figure 6B&C). Analytical gel-filtration coupled with SLS analysis demonstrated that both Trx-TRPC4 tail and Trx-TRPC5 tail formed 1:1 complexes with CaM in a Ca^{2+} -dependent manner (Figure 6D and Figure S8). Detailed mapping using ITC-based quantitative binding analysis revealed that CaM interacts with the first half (CBS1, aa K758-S817) and the second half (CBS2, aa A818-L974) of TRPC4 tail with a K_d values of 0.70 ± 0.06 µM and 1.17 ± 0.22 µM, respectively (Figure 6E and Figure S6). The K_d value of CaM to the entire tail of TRPC4 (aa K758-L974) is 0.34±0.06 µM (Figure 6E), likely due to the mild synergism resulting from the relatively short spacing between CBS1 and CBS2 of TRPC4 (Figure S6A). Interestingly, the mutation eliminating Ca^{2+} binding to the N-lobe (E31/67Q) of CaM abolished its binding to CBS1, and the mutation eliminating Ca²⁺ binding to the C-lobe (E104/140Q) of CaM had very little impact on its binding to CBS1, showing that similar to the Drosophila TRP CBS1/N-lobe interaction, CBS1 of TRPC4 also specifically binds to the N-lobe of CaM. However, different from the TRP CBS2/C-lobe_CaM interaction, the direct binding between TRPC4 CBS2 and C-lobe_CaM was too weak to detect. In the absence of CBS1, CBS2 can bind to Ca²⁺-CaM with a Kd of 1.17±0.22 µM. This binding is also

contributed by the N-lobe of CaM, as elimination of Ca^{2+} binding to the N-lobe of CaM also abolished CBS2/CaM binding (Figure 6E). Based on the above data, we propose that the N-lobe of CaM plays a dominant role in the binding to TRPC4 via CBS1 and the C-lobe plays a minor role in binding to CBS2.

To characterize the molecular mechanism underlying the interaction between TRPC4 and CaM, we tried to determine the structure of TRPC4 CBS12/CaM complex. Although the crystallization of TRPC4 CBS12/CaM complex failed after numerous trials, the crystal structure of the TRPC4 CBS1/N-lobe_CaM complex was determined at a resolution of 1.9 Å (Figure 6F and Table 1). Interestingly, TRPC4 CBS1 in the complex also adopts a "helix-turn-helix" conformation (Figure 6F&G). In the complex, The N-lobe of CaM adopts an open conformation. Both α helices of TRPC4 CBS1 participate in binding to N-lobe_CaM. Different from the TRP CBS1, the α 1 of TRPC4 CBS1 is the main helix contacting N-lobe_CaM with L792 serving as the anchoring residue inserting into the deep pocket of N-lobe_CaM (Figure 6F2). Consistent with this structural analysis, substitutions of L792 together with its neighboring F793 with Ala totally abolished TRPC4 CBS1's binding to CaM (Figure S7B).

Using alanine scanning, we were able to show that the hydrophobic residues from TRPC4 CBS2 also contribute to TRPC4's binding to CaM. Replacing the hydrophobic residues in the CBS2 weakened TRPC4 CBS12's binding to CaM by ~3-fold. The combination of the LF792,793AA substitutions in CBS1 and the hydrophobic residue substitutions in CBS2 totally eliminated TRPC4 tail/CaM binding (Figure S7C5&6).

Although the amino acid sequence of *Drosophila* TRP CBS12 is not conserved in TRPC4 and TRPC5 (Figure 6H), *Drosophila* TRP and mammalian TRPC4 still share a similar bidentate CaM interaction mode. In particular, both *Drosophila* TRP CBS1 and mammalian TRPC4 CBS1 form a helix-turn-helix conformation upon binding to CaM N-lobe. It is possible that this bidentate CaM binding mode of *Drosophila* TRP

and mammalian TRPC4 may be a result of evolution. However, since the CaM binding sequences of *Drosophila* TRP and mammalian TRPC4 share very low similarities (Figure 6H), the bidentate CaM binding mode observed for the two channels could be a simple coincidence.

As for the TRPC5 tail, the residues corresponding to the α 1 helix in TRPC4 CBS1 do not exist (Figure S8B), and thus the TRPC5 tail may not contain a CaM-binding CBS1 found in TRPC4. However, the residues corresponding to E831-K852 (based on mouse TRPC5 sequence) can be aligned with the sequence of TRPC4 CBS2 (Figure S8B). The TRPC5 tail fragment containing E831-K852 was shown to bind to CaM with a Kd of $1.25\pm0.32 \mu$ M (Figure S8C). Substitution of the hydrophobic residues within E831-K852 (i.e. residues highlighted with blue dots in Figure S8B) totally abolished the binding between TRPC5 tail and CaM. Therefore, TRPC5 should only contains one continuous CaM binding segment spanning E831-K852 and the binding between TRPC5 tail to CaM is about 4 times weaker than the binding between TRPC4 tail and CaM.

DISCUSSION

Previous studies have reported that Ca^{2+} plays both positive feedback and negative feedback regulations on the TRP channel activation and termination processes. However, how Ca^{2+} acts on the TRP channel to regulate channel activity remains to be elucidated. On the other hand, although CaM has been reported to bind to the TRP channel, the molecular mechanism underlying CaM/TRP interaction is not known.

Here, we have elucidated the detailed molecular mechanism underlying the interaction between TRP and CaM. We show that CaM utilizes its two lobes to grasp two CBSs of TRP separated by more than 70 residues in a Ca^{2+} -dependent manner. Upon Ca^{2+} -CaM binding, TRP CBS1 forms a unique "helix-turn-helix" structure

while CBS2 forms a canonical single α helix (Figure 3A). The separation of N- and C-lobe_CaM binding sites by an elongated flexible linker has recently been observed in the cryo-EM structures of TRPV5 and TRPV6 (Dang et al., 2019; Hughes et al., 2018; Singh et al., 2018). However, both CBSs on TRPV5 and TRPV6 form a single amphipathic α helix structure upon CaM binding. In TRPV5 and TRPV6, the distal CBS2-bound C-lobe of Ca²⁺-CaM functions to block the channel Ca²⁺ conductance by inserting the side chain of trimethylated K115 into the channel pore by forming a highly unusual trimethyllysine/(Trp)₄ cation- π cage (Dang et al., 2019; Hughes et al., 2018; Singh et al., 2018; Zhang et al., 1994). This trimethyl lysine/(Trp)₄ cation- π cage presumably explains why only Lys115 out of a total of 8 Lys residues in CaM is trimethylated (Roberts et al., 1986; Zhang et al., 1994).

TRP shares greater similarity with the TRPC subfamily channels. TRPC channels include a four-helix bundle formed by the coiled-coil domain after the connecting helix, which functions as a fence separating the channel pore from the rest of the C-terminal tail (Figure 1G). Therefore, the CBS12 bound CaM is unlikely to directly block the channel pore (Figure 4C&D). Additionally, there exists a stretch of flexible sequences of 35-residues connecting the end of the four-helix bundle to the N-terminus of CBS1 in TRPC4 (aa R755-N789) (Duan et al., 2018) (Figure 6H). Similarly, there exists a 33-residue flexible linker between the predicted four-helix bundle and CBS1 in TRP (aa N783-F815) (Figure 4C&D). Thus, we believe that the Ca²⁺-dependent binding of CaM to the TRP tail (or to TRPC4 tail) is not likely to directly alter the channel pore properties. Indeed, in Drosophila eyes, replacing WT TRP with TRP variants containing the CBS1 CaM binding deficient mutation (IF836,837AA), or the CBS2 CaM binding mutation (FM925,926AA), or both CBS1 and CBS2 mutations (the IF836,837AA and FM925,926AA quadruple mutant; characterized in Figure 3D and Figure 4A) did not affect light responses mediated by the channels under physiological conditions. In particular and surprisingly, both the positive and negative feedback mediated by Ca²⁺ appeared fully intact even in the quadruple mutant, which is incapable of binding Ca^{2+} -CaM (Figure 5). This suggests that Ca^{2+} must mediate its pronounced feedback effects on the channel by a as yet unidentified molecular mechanism. One possibility is that Ca^{2+} may directly, instead of via CaM, bind to and regulate TRP activity. It is also possible that CaM binding to the tail of TRP may be required for the downstream signaling or the INAD-mediated signalplex organization. For example, CaM may shuttle between TRP and other CaM binding proteins in response to light-dependent Ca^{2+} concentration fluctuations in rhabdomere (Porter et al., 1995; Porter et al., 1993). Concentrating CaM within the rhabdomere by several proteins including TRP and NINAC may provide a mechanism for fast Ca^{2+} -dependent signaling in *Drosophila* photoreceptors. Future studies will be required to tease out the physiological role of Ca^{2+} -dependent CaM binding to TRP tail.

In summary, we discovered an unexpected Ca^{2+} -dependent interaction mode between CaM and TRP C-terminal tail. Our biochemical and structural studies of the interaction between CaM and TRP tail reveal the remarkable target binding capacity of CaM. We further showed that CaM can bind to TRPC4 tail with a mode like that between CaM and TRP. The remarkable versatility of CaM in binding to its targets reminds us that CaM, the master Ca²⁺ signal modulator, may bind to and regulate activities of many other ion channels or transmembrane receptors with previously unknown interaction modes.

MATERIALS AND METHODS

Key resources table

Reagent type	Designation	Source or	Identifiers
(species) or		reference	
resource			
Constructs	·		
Strain, strain	BL21(DE3)	Novagen	Cat #69450
background (
E.coli)			
Strain, strain	Rosseta(DE3)	Novagen	Cat #70954
background (
E.coli)			
Cell line	HEK293T	ATCC	Cat #CRL-3216;
(human)			RRID: CVCL_0063
Transfected	Flag-Drosophila TRP	This paper	NCBI: NM_057420.4
construct	CBS12: aa N783-D940		
(plasmid)			
Transfected	GFP-TRPC1 Mouse: aa	This paper	NCBI: NM_011643.4
construct	T757-N793		
(plasmid)			
Transfected	GFP-TRPC2 Mouse: aa	This paper	NCBI: AF111108.1
construct	D1042-S1172		
(plasmid)			
Transfected	GFP-TRPC3 Mouse: aa	This paper	NCBI: FJ207475.1
construct	N788-E836		
(plasmid)			
Transfected	GFP-TRPC4 Mouse: aa	This paper	NCBI: NM_016984.3
construct	T732-L974		
(plasmid)			
Transfected	GFP-TRPC5 Mouse: aa	This paper	NCBI: NM_009428.3
construct	T739-L975		
(plasmid)			
Transfected	GFP-TRPC6 Mouse: aa	This paper	NCBI: NM_013838.2
construct	N879-R930		
(plasmid)			
Transfected	GFP-TRPC7 Mouse: aa	This paper	NCBI: NM_012035.3
construct	N805-I862		
(plasmid)			
Antibodies			
Antibody	Anti-Flag (M-2) (mouse	Sigma	Cat #F3165;
	mAb)		RRID:AB_259529

Antibody	Anti-Mouse IgG (Goat	Sigma	Cat #A4416;
	polyAb)		RRID: AB_258167
Recombinate	Proteins and Synthesized	peptides	
Recombinant	Drosophila Calmodulin:	This paper	NCBI: NM_078986.3
protein	aa A1-K148		
Recombinant	Drosophila Calmodulin	This paper	NCBI: NM_078986.3
protein	N_lobe: aa A1-D78		
Recombinant	Drosophila Calmodulin	This paper	NCBI: NM_078986.3
protein	C_lobe: T79-K148		
Recombinant	Mouse Calmodulin: aa	This paper	NCBI: M19381.1
protein	A1-K148		
Recombinant	Drosophila TRP	This paper	NCBI: NM_057420.4
protein	CBS-AB: aa S717-D940		
Recombinant	Drosophila TRP	This paper	NCBI: NM_057420.4
protein	CBS-B: aa N783-D940		
Recombinant	Drosophila TRP	This paper	NCBI: NM_057420.4
protein	CBS12: aa N783-D940		
Recombinant	Drosophila TRP CBS1:	This paper	NCBI: NM_057420.4
protein	aa T802-K862		
Recombinant	Drosophila TRP CBS2:	This paper	NCBI: NM_057420.4
protein	aa M899-D940		
Recombinant	Mouse TRPC4	This paper	NCBI: NM_016984.3
protein	C-terminal tail: aa		
	K758-L974		
Recombinant	Mouse TRPC4 first half	This paper	NCBI: NM_016984.3
protein	of C-terminal tail: aa		
	K758-S817		
Recombinant	Mouse TRPC4 second	This paper	NCBI: NM_016984.3
protein	half of C-terminal tail:		
	aa A818-L974		
Recombinant	Mouse TRPC5	This paper	NCBI: NM_009428.3
protein	C-terminal tail: aa		
	K765-L975		
Recombinant	Mouse TRPC5 first half	This paper	NCBI: NM_009428.3
protein	of C-terminal tail: aa		
	K765-S837		
Recombinant	Mouse TRPC5 second	This paper	NCBI: NM_009428.3
protein	half of C-terminal tail:		
	aa G823-L975		
Synthesized	Mouse TRPC4 CBS1:	Shenzhen	NCBI: NM_016984.3
peptide	aa D758-N812	PepBiotic	
		Co., Ltd.	
Synthesized	Mouse TRPC4 CBS2:	Shenzhen	NCBI: NM_016984.3

peptide	aa E827-E854	PepBiotic		
		Co., Ltd.		
Commercial a	ssays and Softwares	1		
Commercial	Clone Express II,	Vazyme Biot	Cat #C112	
assay or kit	One-Step Cloning Kit	ech Co., Ltd		
Commercial	ViaFect transfection	Promega	Cat #E4981	
assay or kit	reagent	Corporation		
Software,	ASTRA6.1	Wyatt	http://www.wyatt.	
algorithm		Technology	com/products/software/astra.html	
		Corporation		
Software,	Origin7.0	OriginLab	http://www.originlab. com/;	
algorithm			RRID: SCR_002815	
Software,	GraphPad Prism	GraphPad	http://www.graphpad.com/scientif	
algorithm		Software Inc.	ic-software/prism;	
			RRID: SCR_002798	
Software,	HKL2000	HKL	http://www.hkl-xray.com/	
algorithm		Research Inc.		
Software,	CCP4	PMID:	http://www.ccp4.ac.uk/;	
algorithm		21460441	RRID: SCR_007255	
Software,	PHENIX	PMID:	http://www.phenix-online.org/;	
algorithm		20124702	RRID: SCR_014224	
Software,	Coot	Emsley et al.,	https://www2.mrc-lmb.cam.ac.uk/	
algorithm		2010	Personal/pemsley/coot/	
Software,	PyMOL	DeLano	http://www.pymol.org/;	
algorithm		Scientific	RRID: SCR_000305	
		LLC		
Software,	NMRPipe	NIH	https://spin.niddk.nih.gov/NMRPi	
algorithm			pe/ref/index.html	
Software,	Sparky	UCSF	https://www.cgl.ucsf.edu/home/sp	
algorithm		Sparky	arky/	
Deposited Data	a			
Crystal	X-ray diffraction	This paper	PDB code: 7CQV	
structure of				
TRPCBS1/N				
-lobe_CaM				
complex				
Crystal	X-ray diffraction	This paper	PDB code: 7CQH	
structure of				
TRPCBS2/C				
-lobe_CaM				
complex				
Crystal	X-ray diffraction	This paper	PDB code: 7CQP	
structure of				

TRPC4CBS1			
/N-lobe_Ca			
M complex			
Crystal	X-ray diffraction	Rupp et al.,	PDB code: 1UP5
structure of		1996	
chicken			
Calmodulin			
cryo-EM	cryo-EM	Duan et al.,	PDB code: 5Z96
structure of		2018	
mouse			
TRPC4			
cryo-EM	cryo-EM	Tang et al.,	PDB code: 5YX9
structure of		2018	
TRPC6			

Plasmids, protein expression and purification

Plasmids carrying different fragments and mutants of TRP, TRPCs and CaM were generated by the standard PCR-based method. All mutations were confirmed by DNA sequencing. Recombinant proteins with N-terminal 6x His-tag were expressed in *Escherichia coli* BL21 (DE3) or Rosseta (DE3) cells in LB medium at 16°C and purified using Ni sepharose 6 fast flow affinity column, followed by size-exclusion and anion exchange chromatography (GE Healthcare). Purified proteins were prepared in 50 mM Tris-HCl (pH 7.5), 100 mM NaCl, 1 mM CaCl₂ and 2 mM β -Mercaptoethanol for most of the experiments. Uniformly ¹⁵N-labeled samples were prepared by growing bacteria in M9 minimal medium using ¹⁵NH₄Cl (Cambridge Isotope Laboratories Inc.) as the sole nitrogen source.

Analytical gel filtration chromatography coupled with static light scattering

Molar mass measurements were performed on an AKTA FPLC system (GE Healthcare) coupled with a static light scattering detector (mini-DAWN, Wyatt) and a differential refractive index detector (Optilab, Wyatt). 50 μ M protein samples were loaded to a Superdex 200 Increase 10/300 GL column (GE Healthcare) pre-equilibrated by a buffer composed of 50 mM Tris-HCl (pH 7.5), 100 mM NaCl, 1

mM CaCl₂ and 2 mM β -Mercaptoethanol.

Analytical ultracentrifugation analysis

Sedimentation equilibrium and sedimentation velocity experiments were performed using a Beckman Proteomelab XL-I ultracentrifuge equipped with Beckman 50Ti rotor. To preform sedimentation equilibrium experiment, TRP CBS12/CaM complex was centrifuged at 16,000 rpm and equilibrated for 72 h at 16,000 rpm at six sector cells with 3 different concentrations (absorption at 280 nm of 0.3, 0.6, and 0.9, respectively), and scans were taken every 8 h for 2 times, 6 h for 4 times and 4 h for 8 times. To preform sedimentation velocity experiment, TRP CBS12/CaM complex was centrifuged at 40,000 rpm and scans were taken for 400 times with intervals of 1 minute at 2 sector cells at absorption at 280 nm of 0.5. GCN4-CBS12/CaM complex was centrifuged at 24,000 rpm and scans were taken for 400 times with intervals of 1 minute at 2 sector cells at absorption at 280 nm of 0.5. All data were calibrated with the buffer as the background. Sedimentation equilibrium and sedimentation velocity data analyzed using Sedfit and Sedphat programmes were (http://www.analyticalultracentrifugation.com/default). Data fitting was performed using a heterogeneous interaction model and with simulated annealing algorithms (Lebowitz et al., 2002).

NMR experiments

NMR samples contained 0.2 mM of Ca²⁺-CaM or TRP CBS12/Ca²⁺-CaM complex in 50 mM Tris-HCl (pH 8.0, with 2 mM DTT) in 90% H₂O/10% D₂O. HSQC spectra were acquired at 30°C on a Varian Inova 800-MHz spectrometer equipped with an actively z-gradient shielded triple resonance probe. Backbone resonance assignment of CaM were obtained from previous paper (Mal and Ikura, 2006). Spectra were analyzed using NMRPipe (Delaglio et al., 1995) and Sparky (T.D. Goddard and D.G. Kneller, SPARKY 3, University of California, San Francisco).

Protein crystallography

Crystals of TRP CBS1/N-lobe_CaM complex (2.5 M Ammonium sulfate, 0.1 M BIS-TRIS propane pH 7.0), TRP CBS2/C-lobe_CaM complex (11.4% w/v Polyethyleneglycol 20,000, 150 mM Sodium acetate, 7% v/v Ethyleneglycol, pH 4.5) and TRPC4 CBS1/N-lobe_CaM complex (55.7 % w/v Polyethylene glycol 1,000, 150 mM MES, 2 % v/v 2-Methyl-2,4-pentanediol, pH 6.0) were obtained by sitting-drop vapor-diffusion method at 16°C. Diffraction data were collected at the Shanghai Synchrotron Radiation Facility (BL17U or BL19U) at 100K. Data were processed and scaled using HKL2000 (Otwinowski and Minor, 1997).

All three complex crystal structures were determined by molecular replacement with the model of N-lobe_CaM (PDB: 1UP5, Rupp, 1996) using PHASER (McCoy et al., 2007). Further manual model building and refinement of the structures were completed iteratively using Coot (Emsley et al., 2010) and PHENIX (Adams et al., 2010). The final models were validated by MolProbity and the statistics are summarized in Table 1. All structure figures were prepared with PyMOL (http://www.pymol.org).

Isothermal titration calorimetry assay

ITC experiments were performed on a MicroCal ITC200 calorimeter (Malvern, UK) at 25°C. Proteins with high concentrations (300 μ M in the TRP cases or 400 μ M in the TRPC cases) were loaded into the syringe and titrated into the cells containing corresponding interactors with low concentrations (30 μ M in the TRP cases or 40 μ M in the TRPC cases). The sample in the syringe was subsequently injected into the cell with a time interval of 120 s (0.5 μ l for the first injection and 2 μ l each for the following 18 injections). Titration data were analyzed using the Origin7.0 software and fitted with the one-site binding model. The ITC experiments were performed by 3 independent repeats.

GST pull down assay

GST-tagged CaM or GST alone was incubated with purified endogenous TRP or

HEK293T cell lysates expressing Flag-tagged target proteins for 1 h in 4°C. The mixture was mixed with 20 μ l Glutathione Sepharose 4B beads (GE Healthcare) slurry in TBS with 1 mM CaCl₂ or 1 mM EDTA for 30 min in 4°C. After washing twice, the proteins captured by the beads were eluted by boiling with SDS-PAGE loading buffer, resolved by SDS-PAGE and detected using Coomassie brilliant blue R250 staining or Western blot.

Transgenic Drosophila expressing mutated TRP

Mutations of "IF836,837AA", "FM925,926AA" and "IF836,837AA & FM925,926AA" were individually introduced into a 6.5-kb *trp* genomic DNA (Montell et al, 1985), and further cloned into the pattB vector. Three corresponding transgenic lines were generated by PhiC31 integrase-mediated transgenesis to insert at the attp40 site (BestGene), and these transgenes were individually recombined into $trpl^{MB10553}$ and $trp^{MB03672}$ double null mutant background. Homozygote with two transgene copies of each TRP variant were analyzed for whole-cell patch clamp recordings and calmodulin binding.

Drosophila head preparation and endogenous TRP purification

Wild type or mutant *Drosophila* were first quick-frozen in liquid nitrogen and shaken hard to break heads from bodies. Then the heads were filtrated by sieves and mechanically crashed and extracted in extraction buffer (v/v, 1:10) containing 50 mM Tris-HCl pH 8.0, 150 mM NaCl, 6 mM (0.3%) n-dodecyl-β-D-maltoside (DDM), 1x cocktail protease inhibitor (Roche). After extraction, the mixture was centrifuged at 14,000 g for 10 min followed by 100,000 g for 45 min at 4°C. Total protein concentrations of supernatant (about 4 mg/ml) were determined by BCA protein assay. The supernatant was then used in the 3xStrep-NORPA_CT (aa E863-A1095) pull-down assays (similar procedure with the GST pull down assay). After the enrichment of INAD/ePKC/TRP complex by 3xStrep-NORPA to Streptavidin beads (GE Healthcare), TRP were competed from INAD complex by adding excessive

amount of GB1-TRP_CT (aa R1261-L1275) protein. The TRP elution was used for further GST-CaM pull down assay and Transmission Electron Microscopy (TEM) experiments.

Transmission Electron Microscopy

Determination of the polymeric state of endogenous TRP/CaM complex was achieved by negative-staining TEM. Purified endogenous TRP was eluted as a tetramer in Superose 6 Increase 10/300 GL column (GE Healthcare) in the buffer containing 50 mM Tris-HCl pH 8.0, 150 mM NaCl, 0.03% DDM, 2 mM DTT. 4 μ L diluted (about 0.01 mg/ml) TRP or TRP/CaM complex (1:1 molar ratio) sample was spotted to glow-discharged carbon-coated copper grids (Beijing Zhongkejingyi Technology). After 60s of absorption, the sample was blotted, and then stained with 2% uranyl acetate. Images were recorded using a transmission electron microscope TEM HT7700 (HITACHI) equipped with a field emission gun and operated at an acceleration voltage of 100 kV.

Cell culture and transfection

HEK293T cells were cultured in Dulbecco's Modified Eagle Medium (DMEM) supplemented with 10% fetal bovine serum (FBS), and 1% of penicillin-streptomycin at 37°C with 5% CO₂. HEK293T cells were transfected with Flag-tagged target proteins using ViaFect Transfection Reagent (Promega, Madison, WI) following the manufacturer's instruction. After 24-36 h, cells were harvested and stored at -80°C preparing for the pull-down assay.

Whole-cell patch clamp recordings

Whole-cell patch clamp recordings of photoreceptors from dissociated ommatidia from newly eclosed adult flies of either sex were performed as previously described (e.g. Reuss et al 1997). Standard bath contained (in mM): 120 NaCl, 5 KCl, 10 *N*-Tris-(hydroxymethyl)-methyl-2-amino-ethanesulphonic acid (TES), 4 MgCl₂, 1.5 CaCl₂, 25 proline and 5 alanine, pH 7.15. For Ca²⁺ free bath CaCl₂ was omitted and 1 mM Na₂EGTA added; the solution being applied locally from a nearby puffer pipette. The intracellular pipette solution was (in mM): 140 K gluconate, 10 TES, 4 Mg-ATP, 2 MgCl₂, 1 NAD and 0.4 Na-GTP, pH 7.15. Chemicals were obtained from Sigma-Merck (Germany) and VWR (USA). Recordings were made at room temperature ($22 \pm 1^{\circ}$ C) at -70 mV using electrodes of resistance 10-15 M Ω . Data were collected and analyzed using Axon amplifiers and pCLAMP v.9 or 10 software (Molecular Devices, Union City, CA). Photoreceptors were stimulated via a green (522 nm) ultrabright light-emitting-diode (LED) controlled by a custom-made LED driver; intensities were calibrated in terms of effectively absorbed photons by counting quantum bumps at low intensities. Differences between experimental groups were tested with one-way Anova followed by the Tukey's test.

Quantification and Statistical Analysis

Statistical parameters including the definitions and exact values of n (e.g., number of experiments), distributions and deviations are reported in the Figures and corresponding Figure Legends. For TRP channel amplitudes recordings, the results were expressed as mean \pm SD using one-way Anova with the Turky's test.

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Author contributions

WC and Z Sun performed biochemical experiments; Z Shen and FY were responsible for structural studies; SA performed electrophysiology experiments, ZC generated transgenic flies; CM, RH, WL, MZ designed and supervised the research, all authors analyzed the data; WC and MZ wrote the manuscript with input from all other authors. MZ coordinated the research.

Competing interests

The authors declare no competing interests.

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Data Collection				
Dataset	TRP CBS1/N-lobe	TRP CBS2/C-lobe	TRPC4 CBS1/N-lobe	
Space group	H3 ₂	I41	P6122	
Wavelength	0.97890	0.97890	0.97890	
Unit cell (a,b,c,Å)	98.449, 98.449,	58.344, 58.344, 92.21	64.46, 46.46, 120.75	
	134.26			
Unit cell $(\alpha, \beta, \gamma, \circ)$	90, 90, 120	90, 90, 90	90, 90, 120	
Resolution range (Å)	49.27-1.78(1.81-1.78)	49.31-2.15(2.19-2.15)	50.00-1.90(1.97-1.90)	
No. of	24105(1192)	8342(369)	12754(1169)	
unique reflections				
Redundancy	19.6(19.5)	12.6(9.1)	7.4(7.5)	
I/sigma	55.8(7.27)	50.18(2.71)	19.11(2.29)	
Completeness (%)	100.0(100.0)	99.1(88.5)	99.7(99.5)	
Rmerge ^a (%)	6.1(35.6)	5.0(36.8)	10.2(100)	
Structure Refinement				
Resolution (Å)	49.27-1.78(1.85-1.78)	49.31-2.15(2.19-2.15)	50.00-1.90(2.09-1.90)	
$R_{cryst}^{b}/R_{free}^{c}(\%)$	20.48/23.39	21.40/25.32	21.94/24.98	
Rmsd bonds (Å)	0.007/0.963	0.0107/1.45	0.006/0.909	
/angles (°)				
Average B factors	30.88	68.58	27.54	
(Å ²)				
No. of atoms				
Protein atoms	1444	664	725	
Water	73	0	30	
Other molecules	4	2	2	
No. of reflections				
Working set	22889(2530)	7910(513)	11637(2800)	
Test set	1202(127)	415(34)	607(140)	
Ramachandran plot				
regions				
Favored(%)	98.40	98.85	98.90	
Allowed(%)	0.53	0.00	1.10	
Outliers(%)	0.53	1.15	0.00	

Table 1: Statistics of X-ray Crystallographic Data Collection and Model Refinement

Numbers in parentheses represent the value for the highest resolution shell.

a. $R_{merge} = \Sigma |I_i - \langle I \rangle| / \Sigma |I_i|$, where I_i is the intensity of measured reflection and $\langle I \rangle$ is the mean intensity of all symmetry-related reflections.

b. $R_{cryst} = \Sigma ||F_{calc}| - |F_{obs}|| / \Sigma F_{obs}$, where F_{obs} and F_{calc} are observed and calculated structure factors.

c. $R_{free} = \Sigma T ||F_{calc}| - |F_{obs}|| / \Sigma F_{obs}$, where T is a test data set of about 5% of the total unique reflections randomly chosen and set aside prior to refinement.



Figure 1: Identification of CaM binding sites at the *Drosophila* TRP C-terminal Tail

- (A) Schematic diagrams showing the domain and topographic organizations of *Drosophila* TRP channel. The two CaM binding sites are designated as CBS-A and CBS-B. ANK stands for ankyrin repeats and PBM represents PDZ domain binding motif.
- (B) Analytical gel filtration chromatography analysis showing that there is no detectable binding between TRP (940-1275) and CaM both in the presence and absence of Ca^{2+} .
- (C) Analytical gel filtration chromatography coupled with static light scattering analysis showing that TRP CBS-AB (717-940), with or without the Trx-tag, binds to Ca²⁺-CaM forming a 1:2 molar ratio complex.
- (D) Analytical gel filtration chromatography coupled with static light scattering analysis showing that TRP CBS-B (783-940), with or without the Trx-tag, binds to Ca²⁺-CaM forming a 1:1 molar ratio complex.
- (E) Isothermal titration calorimetry (ITC)-based measurement of the binding of Ca²⁺-CaM to the TRP CBS-A peptide (728-754).
- (F) Sequence alignment of the connecting helix and the coiled-coil domain of TRP, TRPL and mouse TRPC1-7. Based on the cryo-EM structure of TRPC6, the connecting helix is shown as a bar in salmon and the coiled-coil domain is indicated as an orange bar above the alignment. The CBS-A peptide of TRP used in panel E is indicated by a pink box.
- (G) Ribbon diagram representation of the close state cryo-EM structure of TRPC6 (PDB: 5YX9). The region highlighted in the dashed box contains the connecting helix (colored in salmon) and the coiled coil domain (colored in orange) of TRPC6. For simplicity, only one monomer subunit of the connecting helix and the coiled coil domain are colored in both the ribbon and ribbon combined with the surface model shown on the right (the rest part colored in gray). The other three monomer subunits are colored in green. The structure shows that the connecting helix is completely embedded in the tetramer structure of TRPC6 and not accessible to CaM.



Figure 2: CaM binds to TRP C-terminal tail with an unexpected binding mode

- (A) Sequence alignment of TRP CBS-B from different insect species. The two CaM binding sites, CBS1(802-862) and CBS2(899-940), are highlighted and their secondary structures when in complex with Ca²⁺-CaM are indicated. In this alignment, the completely conserved residues are shaded in red, highly conserved residues shown in red, and non-conserved residues in black.
- (B) ITC-based measurements of the bindings of CaM to TRP CBS1 (B1) and CBS2 (B2) in the presence (black curve) or absence (red curve) of Ca²⁺.
- (C) Analytical ultracentrifugation sedimentation velocity analysis showing that Ca^{2+} -CaM binds to TRP CBS12 forming a homogenous 1:1 molar ratio complex.

- (D) Analytical gel filtration coupled with static light scattering showing that CBS1, CBS2 and CaM can form a stable ternary complex with a 1:1:1 stoichiometry. The SDS-PAGE with Coomassie blue staining at the bottom panel shows the protein composition of the elution peak of the CBS1, CBS2 and CaM mixture shown in red at the top panel.
- (E) Overlay of the ¹H,¹⁵N-HSQC spectra of the ¹⁵N-Ca²⁺-CaM/¹⁵N-CBS12 complex (red) with those of ¹⁵N-Ca²⁺-CaM in complex with ¹⁴N-CBS1 (blue) and ¹⁴N-CBS2 (purple), respectively.
- (F) Cartoon showing the binding mode between TRP CBS12 and Ca²⁺-CaM.
- (G) Dissociation constants of the TRP CBS12/Ca²⁺-CaM complex derived from the analytical ultracentrifugation sedimentation equilibrium analysis.



Figure 3: Crystal structures of Ca²⁺-CaM in complex with TRP CBS1 and CBS2 and validation of the structure

(A) Ribbon diagrams showing the structures of the TRP CBS1/N-lobe_CaM complex and the CBS2/C-lobe_CaM complex. The N/C-lobe_CaM are colored in salmon. TRP CBS1 is colored in pale blue, and the CBS2 is colored in deep blue. The two lobes of CaM are arbitrarily connected with a dotted line. CBS1 and CBS2 in the complex are separated by a 73-residues linker indicated by a dashed line.

- (B,C) The combined surface (N/C-lobe_CaM) and ribbon (TRP CBS1/CBS2) models showing the interaction between TRP CBS1/CBS2 and Ca²⁺-CaM. In the surface model, the positively charged amino acids of N/C-lobe_CaM are highlighted in blue, the negatively charged residues in red, the hydrophobic residues in yellow, and the others in white. The sidechains of residues from TRP CBS1/CBS2 that are involved in binding to CaM are shown with the stick model.
- (D) The ribbon models of the original structure of CBS1/N-lobe complex (D1) and the aligned structure of CBS2/N-lobe complex (D2). The critical K75 from the CaM N-lobe are highlighted by red dashed box.
- (E) Table summarizing the binding affinities of various mutants of CBS1 and CBS2 to Ca²⁺-CaM derived from ITC-based assays.
- (F) GST pull-down assay showing that when both IF836,837AA and FM925,926AA are introduced into TRP-CBS12, the interaction between CBS12 and CaM is totally abolished. Note that we intentionally used excess amount of the IF836,837AA/FM925,926AA quadruple mutant in the pull-down assay to show that even weak interaction could not be detected.



Figure 4: Ca²⁺-dependent interaction between full-length TRP and CaM

- (A) GST pull-down assay showing that endogenous TRP purified from fly heads interacted with CaM in a Ca²⁺-dependent manner. GST-CaM had no detectable binding to TRP when Ca²⁺ in the assay buffer was chelated by EDTA. When IF836,837AA and FM925,926AA quadruple mutations were introduced to TRP (the quadruple mutations were introduced in a *trp* genomic rescue in a *trpl^{MB10553};trp^{MB03672}* double null mutant background), the interaction between TRP and Ca²⁺-CaM was totally abolished.
- (B) Analytical ultracentrifugation sedimentation velocity assay showing the formation of a 4:4 complex between GCN4-CBS12 (TRP CBS12 fused to the C-terminal tail of a GCN4 coiled-coil tetramer) and Ca²⁺-CaM.
- (C&D) Schematic and ribbon diagrams showing that the two lobes of Ca²⁺-CaM bind to CBS1 and CBS2 from the same molecule of the TRP tail (i.e. the intra-molecular binding model; C); or the two lobes of Ca²⁺-CaM bind to CBS1

and CBS2 from two different subunits the TRP tails (i.e. the inter-molecular binding model; D). The models were generated by docking the TRP CBS1/N-lobe_CaM complex and TRP CBS2/C-lobe_CaM complex crystal structures on to the cryo-EM structure of TRPC4, PDB:5Z96.



Figure 5: Disruptions of CaM binding to CBS1, CBS2 or CBS12 do not directly affect Ca²⁺-mediated feedback on TRP channel activity

- (A) Example responses to 1 ms flashes (arrows) containing ~150 effective photons in whole-cell patch-clamped TRP wild-type photoreceptors (in the *trpl* background) in the presence and absence of bath Ca^{2+} .
- (B-D) Example responses to 1 ms flashes (arrows) (~150 effective photons) in photoreceptors from TRP CBS1 (B), CBS2 (C) and CBS1+2 mutants (in the $trpl^{MB10553}$; $trp^{MB03672}$ background), in the presence and absence of bath Ca²⁺.
- (E) Quantification of peak response amplitude of photoreceptors expressing either TRP-WT (*trpl*) or Ca²⁺-CaM binding mutants of TRP, in the presence of physiological Ca²⁺ (1.5 mM). Individual values and means ± SD are shown for WT (n=23 cells) and mutants (n=16-19). The four groups were not significantly different (p=0.26; one-way ANOVA)
- (F) Percentage ratios of peak response amplitudes in 0 Ca²⁺ over physiological Ca²⁺ in the TRP-WT and mutants. Individual values and means \pm SD are shown for WT

(n=14 cells) and mutants (n=11-15). The four groups were significantly different (p<0.0001; one-way ANOVA). In particular, all three mutants had larger ratios (i.e. larger relative 0 Ca²⁺ responses) compared to WT (p<0.0001 for each pair; Tukey's post test). Moreover, CBS2 and CBS1+2 both had significantly larger ratios compared to CBS1 (p<0.001 and p<0.01; Tukey's test).



Figure 6: TRPC4 binds to Ca²⁺-CaM following a similar mode as TRP does

- (A) Schematic diagram showing the domain organizations of the C-terminal tails of *Drosophila* TRP, TRPL and the mouse TRPC subfamily members 1-7.
- (B) GST pull down assay showing that Ca²⁺-CaM only binds to the tails of TRPC4 and TRPC5.
- (C) Ponceau S staining of GST and GST-CaM used in the pull-down assay in panel B.
- (D) Analytical gel filtration coupled with static light scattering showing that TRPC4 forms a 1:1 molar ratio complex with CaM in a Ca²⁺-dependent manner.
- (E) Table summarizing the binding affinities of CaM to TRPC4 CBS12 (758-974), CBS1 (758-817), and CBS2 (818-974). The table also shows that the impact of Ca²⁺ binding deficient mutants of CaM in binding to CBS1, CBS2, and CBS12.
- (F) Crystal structure of the TRPC4 CBS1 in complex with the N-lobe_CaM shown in the ribbon (F1) and surface combined with ribbon (F2) models with the same coloring scheme as in Figure 3B.
- (G) Alignment of the TRP CBS1/N-lobe_CaM and the TRPC4 CBS1/N-lobe_CaM complexes showing the formation of the "helix-turn-helix" conformation of CBS1 from TRP (yellow) and from TRPC4 (cyan) upon binding to N-lobe of Ca²⁺-CaM. The N-lobe_CaM structures were superimposed to each other in this alignment.
- (H) Sequence alignment of CBS12 regions of TRP, TRPC4 and TRPC5 with the same coloring scheme as that in Figure 2A. The residues from TRPC4 CBS1 and CBS2 that are critical for CaM binding are indicated by red dots. The residues critical for CaM binding in TRP CBS1/2 are indicated by blue dots.

Supplemental Figures



Figure S1: Overlay of the ¹H,¹⁵N-HSQC Spectra of ¹⁵N-Ca²⁺-CaM with different CaM binding sites of TRP

(A) Overlay of the ¹H,¹⁵N-HSQC spectra of ¹⁵N-Ca²⁺-CaM alone (red) and with those of ¹⁵N-Ca²⁺-CaM in complex with ¹⁴N-TRP CBS1 (black) and CBS2 (purple), respectively. The region shown in the dashed box are the residues corresponding to the conserved Gly residue in each EF-hand of Ca²⁺-bound CaM and is enlarged at the right for detailed comparison.

(B) Overlay of the ¹H,¹⁵N-HSQC spectrum of ¹⁵N-Ca²⁺-CaM in complex with

¹⁵N-CBS12 (blue) with those of ¹⁵N-Ca²⁺-CaM in complex with ¹⁴N-TRP CBS1 (black, left panel) and CBS2 (purple, right panel), respectively.



Figure S2: ITC-based measurements of the binding affinities of CaM mutants to TRP CBS1 and CBS2

(A) ITC-based measurements of the bindings of TRP CBS1 to CaM and its mutants: CaM WT (A1), CaM_E31/67Q (A2), CaM_E104/140Q (A3).

(B) ITC-based measurements of the binding of TRP CBS2 to CaM and its mutants: CaM WT (B1), CaM_E31/67Q (B2), CaM_E104/140Q (B3).

(C) ITC-based measurement of the binding of TRP CBS2 K924A to CaM E104/140Q.



Figure S3: Omit map of the TRP CBS1/N-lobe_CaM complex and the TRP CBS2/C-lobe_CaM complex

(A) Omit map of the $\alpha 1$ and $\alpha 2$ helices of CBS1 in complex with CaM N-lobe. The Fo-Fc density map was generated by deleting $\alpha 1$ and $\alpha 2$ and contoured at 2.5 σ .

(B) Omit map of CBS2 α 3 bound to CaM C-lobe. The Fo-Fc density map was generated by deleting α 3 and contoured at 2.5 σ .



Figure S4: ITC-based measurements of the binding affinities of Ca²⁺-CaM to TRP CBS1 and CBS2 mutants

(A) ITC-based measurements of the bindings of Ca²⁺-CaM to different TRP CBS1 mutants: CBS1 WT (A1), CBS1 FV819,820AA (A2), CBS1 L823A (A3), CBS1 IF836,837AA (A4), CBS1 L840A (A5), and CBS1 VI843,844AA (A6).

(B) ITC-based measurements of the bindings of Ca^{2+} -CaM to TRP CBS2 mutants: CBS2 WT (B1), CBS2 V920A (B2), CBS2 Y922A (B3), and CBS2 FM925,926AA (B4).



Figure S5. Negative staining TEM images of purified endogenous TRP with or without addition of CaM.

(A) TRP alone in 1mM CaCl₂. (B) TRP:CaM = 1:1 in 1mM CaCl₂. (C) TRP:CaM(E104/140Q) = 1:1 in 1mM CaCl₂. (D) TRP:CaM = 1:1 in 1mM EDTA.



Figure S6: ITC-based measurements of the binding affinities of CaM mutants to TRPC4 fragments

(A) ITC-based measurements of the binding affinities of CaM (A1) and its Ca^{2+} binding deficient mutants E31/67Q (A2) and E104/140Q (A3) to TRPC4 CBS12.

(B) ITC-based measurements of the binding affinities of CaM (B1) and its Ca^{2+} binding deficient mutants E31/67Q (B2) and E104/140Q (B3) to TRPC4 CBS1 (758-817).

(C) ITC-based measurements of the binding affinities of TRPC4 CBS2(817-974) to CaM (C1) and its Ca^{2+} binding deficient mutants: E31/67Q (C2) and E104/140Q (C3).

(D) ITC-based measurement of the binding affinity of CaM to the TRPC4 C-terminal fragment (854-974).



Figure S7: ITC-based measurements of the bindings of CaM to TRPC4 mutants

(A) Sequence alignment of CBS12 of TRPC4 from different species with the same coloring scheme as that in Figure 2A. The residues from TPRC4 CBS1 and CBS2 that are critical for CaM binding are indicated by blue dots.

(B) ITC-based measurement of the binding affinity of CaM to TRPC4 CBS1(758-817) LF792,793AA.

(C) ITC-based measurements of the binding affinities of CaM to TRPC4 (758-974) mutants: TRPC4 WT (C1), TRPC4 LF792,793AA (C2), TRPC4

FV834,835AA&FGLF841,842,843,844AGAA (C3), TRPC4 LFtoAA&FVtoAA (C4),TRPC4LFtoAA&FGLFtoAGAA (C5),andTRPC4LFtoAA&FVtoAA&FGLFtoAGAA (C6).



Figure S8: TRPC5 C-terminal tail interacts with CaM in a Ca²⁺-dependent manner

(A) Analytical gel filtration coupled with static light scattering showing that TRPC5 tail forms a 1:1 complex with CaM in a Ca^{2+} -dependent manner.

(B) Sequence alignment between TRPC4 and TRPC5 corresponding to the CBS12 region of TRPC4. Note that the TRPC5 tail does not contain critical residues corresponding to CBS1 in TRPC4. The residues for critical for CaM interaction in TRPC4 and TRPC5 CBSs are indicated by blue dots.

(C) ITC-based measurements of the binding affinities of CaM and its mutants to different TRPC5 C-terminal fragments: CaM WT to TRPC5 (765-837, C1), CaM WT to TRPC5 CBS (822-975, C2), CaM WT to TRPC5 very C-terminal fragment(859-975, C3), CaM E31/67Q to TRPC5 CBS (822-975, C4), CaM E104/140Q to TRPC5 CBS (822-975, C4), and CaM WT to TRPC5 FM838,839AA&F843A&LGLF846,847,848,849AGAA (765-975, C6).