Structural aspects of calcium-release activated calcium channel function

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Introduction

Spatio-temporal changes in cytosolic calcium (Ca^{2+}) levels drive a plethora of signaling events in eukaryotic cells that mediate diverse physiological and pathophysiological processes including apoptosis, fertilization, the immune response and transcription, to name a few.1,3 Large differences in intracellular compartmentalized Ca^{2+} levels can induce local transient changes in cytosolic Ca^{2+} concentrations. For example, the ER lumen maintains basal Ca^{2+} levels in the ~sub-mM concentration range, compared with the cytosol which generally keeps basal Ca^{2+} at ~sub-μM levels.1 This steep concentration gradient allows for passive movement of Ca^{2+} from the ER lumen to the cytosol at a minimal energetic cost through, for instance, inositol 1,4,5-trisphosphate receptor (Ins[1,4,5]P_3 R) Ca^{2+} release channels on the ER. Ins[1,4,5]P_3 R channels open after binding the small second messenger inositol 1,4,5-trisphosphate (Ins[1,4,5]P_3), produced by the catalytic conversion of phosphatidylinositol 4,5-bisphosphate (PtdIns[4,5]P_2) into Ins(1,4,5)P_3 and diacylglycerol by phospholipase C after agonist-induced G-protein coupled receptor activation (reviewed in ref. 5). It is imperative that depletion of Ca^{2+} from the ER does not detrimentally effect vital luminal processes such as protein chaperone function;1 hence, Ins(1,4,5)P_3 Rs have evolved a bell-shaped cytosolic Ca^{2+} dependent channel activity, where open probability is highly suppressed at low (i.e., < 10 nM) as well high cytosolic Ca^{2+} concentrations (i.e., > 3 μM) and is maximal at ~100 nM–1 μM.7,8

The Ca^{2+} released from the ER lumen only transiently increases cytosolic Ca^{2+} levels as pump, exchanger and buffer proteins quickly sequester and move the Ca^{2+} into different compartments and organelles such as the mitochondria, golgi and extracellular space. However, some physiological signals require longer term cytosolic Ca^{2+} increases such as T-cell activation that is dependent on transcriptional regulation.4,9,10 Eukaryotes have evolved the process of store operated Ca^{2+} entry (SOCE), which is intimately linked to agonist-induced ER stored Ca^{2+} depletion, to deliver sustained augmentation of cytosolic Ca^{2+} levels. SOCE occurs after ER Ca^{2+} store depletion; further, the reduction in luminal Ca^{2+} (i.e., > 1 mM) to the cytosol (i.e., < 1 μM) is maximal at ~100 nM–1 μM.7,8 SOCE-mediated increases in cytosolic Ca^{2+} not only signal various physiological processes, but also provide Ca^{2+} for the depletion of luminal Ca^{2+} by active pumping through sarco/endoplasmic reticulum Ca^{2+}-ATPases (SERCA) located on the ER.

The molecular players that choreograph SOCE include the ER/SR-resident stromal interaction molecules (STIMs)13,16 and the Orai plasma membrane Ca^{2+} channels.17,22 STIMs are single pass, type I transmembrane (TM) proteins that sense changes in ER luminal Ca^{2+} levels via communicative interactions between EF-hand and sterile α motif (SAM) domains, conserved from lower to higher eukaryotes.16,22-25 This sensory luminal domain machinery is separated from 3 conserved cytosolic coiled-coil (CC) domains (i.e., CC1, CC2, and CC3) by an ~20 amino acid TM domain (Fig. 1A). The cytosolic STIM domains undergo a structural reorganization after luminal depletion,26-29 dependent on the allostery and intermolecular interactions of the Ca^{2+} sensing EF-hand and SAM domains.26-33 This structural change permits STIMs...

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physically couple to Orai subunits at ER-PM junctions, mediating stoichiometric channel assembly as well as gating of these PM Ca\textsuperscript{2+} release activated Ca\textsuperscript{2+} (CRAC) channels. The Orai family of proteins which constitute CRAC channels are 4 TM spanning proteins with cytosol-oriented N- and C-terminal domains (Fig. 1B) (reviewed in ref. 34). STIM coupling to both N- and C-terminal domains is implicated in recruitment and gating of the channel. 35-40 Orai CRAC channels assemble with the TM1 segments constituting the central pore 17,20,41-45 of a tetrameric 46-50 or hexameric architecture. 41,51 Recent structural information on the CRAC channel components has provided a wealth of information on the molecular basis of channel regulation. Structures of the EF-hand together with the SAM domain (EF-SAM) have improved our understanding of how the luminal STIM region senses changes in Ca\textsuperscript{2+} levels; further, high resolution data on the conserved cytosolic domains has increased our comprehension of the supercoiling events that play a role in mediating the STIM quiescent and active conformations. The elucidation of an Orai channel structure has revealed atomic features, vital for assembly of the channel as well as permeation and selectivity of Ca\textsuperscript{2+} through the TM1-constituted pore. Further, a structure of the ubiquitous and cytosolic Ca\textsuperscript{2+} sensor/regulator calmodulin (CaM) in complex with the Orai1 N-terminal domain has exposed a mechanism for negative channel regulation. This review focuses on the structural details of the aforementioned high resolution data and discusses the implications for CRAC channel regulation and function.

**STIM EF-SAM Structure and Function**

Vertebrates encode 2 STIM isoforms, and although STIM1 and STIM2 are ubiquitously expressed in mammalian tissues, STIM2 is more enriched in neuronal and brain cells compared with STIM1. 24,25 Nonetheless, STIM1 appears to play a dominant role in regulating the ON/OFF capacity of CRAC channels, 14,15 whereas STIM2 plays a prevailing role in basal intracellular Ca\textsuperscript{2+} homeostasis. 53,54 At basal ER Ca\textsuperscript{2+} levels (i.e., ~400 \(\mu\)M), the majority of STIM1 is in a quiescent state; on the other hand, a considerable fraction of STIM2 is in an activated state, coupling to and opening PM CRAC channels that contribute to the maintenance of resting intracellular Ca\textsuperscript{2+} levels. 53 Upon ER Ca\textsuperscript{2+} depletion, a large fraction of STIM1 molecules become activated compared with the more modest level of remaining STIM2 activation. This difference in Ca\textsuperscript{2+} sensitivity is partly attributable to distinctions in Ca\textsuperscript{2+} binding properties of STIM1 compared with STIM2. In vitro Ca\textsuperscript{2+} binding experiments have demonstrated that isolated and purified STIM1 EF-SAM domains bind Ca\textsuperscript{2+} with an equilibrium dissociation constant (\(K_d\)) of ~400 \(\mu\)M, 30,31 compared with STIM2 EF-SAM which shows weaker affinity (i.e., \(K_d\) ~700 \(\mu\)M). 32,55 Binding of Ca\textsuperscript{2+} keeps EF-SAM in a monomeric conformation and prevents dimerization/oligomerization and the accompanying conformational change prerequisite to the translocation of STIM to ER-PM junctions where coupling with Orai occurs.

High resolution solution nuclear magnetic resonance (NMR) structures of STIM1 and STIM2 EF-SAM have revealed that
functional \( \text{Ca}^{2+} \) sensing distinctions between the isoforms involve factors in addition to \( \text{Ca}^{2+} \) affinity. The \( \text{Ca}^{2+} \)-loaded STIM1 EF-SAM domain is primarily \( \alpha \)-helical and folds into a compact and stable monomer (Fig. 2A). While primary sequence prediction reveals only a single canonical \( \text{Ca}^{2+} \) binding EF-hand motif (i.e., \( \alpha 1\text{-loop1-} \alpha 2 \)), a second non-canonical EF-hand is, indeed present (i.e., \( \alpha 3\text{-loop2-} \alpha 4 \)), exhibiting mutual \( \text{loop1:loop2} \) stabilization via backbone hydrogen bonding between the carbonyl oxygen and amide hydrogen atoms of Val83 (loop1) and Ile115 (loop2). This STIM1 helix-loop-helix EF-hand pair forms a concave hydrophobic pocket comprised of residues from both the canonical and non-canonical EF-hands. The Val68, Ile71, His72, Leu74, Met75, and Leu92 from the canonical EF-hand and Leu96, Lys104, Phe108, Ile115, and Leu120 from the non-canonical EF-hand create this hydrophobic cleft (Fig. 2B). The STIM1 SAM domain folds into a 5 \( \alpha \)-helix bundle (i.e., \( \alpha 6\text{-} \alpha 10 \)), and a short linker helix (i.e., \( \alpha 5 \)) connects the 2 luminal domains in sequence space (Fig. 2A). However, the EF-hand and SAM domains intimately interact through the hydrophobic EF-hand domain pocket and critical hydrophobic residues near the C-terminus of the \( \alpha 10 \) helix on the SAM domain (i.e., Leu195 and Leu199) (Fig. 2C). These \( \alpha 10 \) anchor residues pack into the EF-hand cleft and stabilize EF-SAM as a single entity.

The intimate EF-hand:SAM domain interface revealed in the STIM1 EF-SAM structure was the first interaction involving well-characterized EF-hand and SAM domains identified in nature and has important implications on understanding the \( \text{Ca}^{2+} \) sensing function of STIM proteins. Upon \( \text{Ca}^{2+} \) depletion, STIM1 EF-SAM becomes markedly destabilized; further, this destabilization promotes homomeric EF-SAM:EF-SAM dimerization and oligomerization, a vital initiation step that...
leads to a conformational change in the cytosolic STIM1 region required for binding to and activating Orai channels at ER-PM junctions. The separate introduction of mutations in the EF-hand cleft and SAM domain anchors aimed at disruption of the structurally elucidated EF-hand:SAM domain interaction induces a similar destabilization in the EF-SAM context in vitro and activates full-length STIM1 in live cells without any perturbations in Ca\(^{2+}\) binding. These data suggest that a compact and well-folded EF-SAM domain in the presence of Ca\(^{2+}\) maintains STIM in a quiescent conformation, while Ca\(^{2+}\)-depletion-induced destabilization of EF-SAM and the accompanying partial unfolding drives the oligomerization of the N-terminal region which initiates the downstream molecular mechanisms of SOCE activation.

Examination of isoform-specific EF-SAM stability revealed that STIM2 EF-SAM is more stable than STIM1 in both the presence and absence of Ca\(^{2+}\). This enhanced stability, despite the lower binding affinity for STIM2 EF-SAM compared with STIM1 implies that other structural factors must compensate for attenuated stability conduced by the higher \(K_d\). Indeed, the high resolution solution NMR structure of Ca\(^{2+}\)-loaded STIM2 EF-SAM exposed 4 critical structural features which enhance the stability of STIM2 EF-SAM compared with STIM1. Overall, STIM2 EF-SAM is comprised of 10 \(\alpha\)-helices forming a globular and compact shape similar to STIM1 (Fig. 2D). The EF-hand domain contains a canonical Ca\(^{2+}\) binding EF-hand motif (i.e., \(\alpha\)-loop1-\(\alpha\)-2) as well as a non-canonical EF-hand motif (i.e., \(\alpha\)-3-loop2-\(\alpha\)-4) that does not bind Ca\(^{2+}\), but plays a stabilizing role through backbone hydrogen bonding of the 2 adjacent loops (i.e., N(H) of Ile87 in loop1: C(O) of Ile119 in loop2). This EF-hand pair creates a more extensive hydrophobic pocket than observed for STIM1, as 13 residues with hydrophobic character contribute to the cleft, compared with only 11 for STIM1. Specifically, Leu72, Ile75, His76, Met79, Ile87, Phe95, and Met100 from the canonical EF-hand and Lys103, Lys108, Leu112, Ile119, Leu124, and Trp128 from the non-canonical EF-hand make up the STIM2 EF-hand pocket (Fig. 2E). The increased hydrophobicity of the STIM2 EF-hand cleft is due to Lys103 and Trp128, which are directed out of the pocket in the case of STIM1 (i.e., His99 and Trp124 STIM1 numbering) (Fig. 2B and E). Lys103 is involved in the second structural factor which enhances the stability of STIM2 EF-SAM compared with STIM1, as this residue plays a dual role in contributing to the hydrophobicity of the EF-hand cleft via the aliphatic CH\(_2\) groups and forming a close ionic interaction with the acidic Asp200 located on the SAM domain through the positively charged ε-amino group.

The third structural factor enhancing the stability STIM2 EF-SAM is the hydrophobicity of the SAM domain core. The STIM2 SAM domain folds into a 5 \(\alpha\)-helix bundle (i.e., \(\alpha\)-6-\(\alpha\)-10) with 12 hydrophobic residues at least 95% solvent inaccessible compared with only 8 in the STIM1 SAM core. This enhanced hydrophobic packing is influenced by Ile180 which is not conserved in STIM1, and also involves the inclusion of Phe158 and Lys193 side chains which project out of the core in STIM1 SAM (i.e., Phe154 and Lys189 STIM1 numbering) (Fig. 2C and F). The final factor enhancing STIM2 SAM stability is the number of hydrophobic anchors at the \(\alpha\)10 helix that pack into the EF-hand cleft. The STIM2 \(\alpha\)10 helix conserves the 2 anchor residues identified in STIM1 EF-SAM (i.e., Leu199 and Leu203 STIM2 numbering); however, STIM2 contains an additional hydrophobic Val201 that is not conserved in STIM1, increasing the stability of the EF-hand:SAM domain interaction (Fig. 2C and F). Functionally, a STIM1/STIM2 EF-SAM chimeric approach was used to corroborate the significance of stability to the Ca\(^{2+}\) sensing function of STIMs. Specifically, by delineating the canonical EF-hand motif, non-canonical EF-hand motif and the SAM domain as the building blocks of EF-SAM and engineering every STIM1/STIM2 chimeric combination, both ‘super-stable’ and ‘super-unstable’ EF-SAM domains were created. Importantly, when swapped with the endogenous EF-SAM in the full-length STIM1 context and expressed in live cells the ‘super-stable’ chimera which consists of the STIM1 canonical EF-hand, the STIM2 non-canonical EF-hand and the STIM2 SAM domain (i.e., ES212) demonstrates suppressed maximum attainable inward current densities indicative of decreased CRAC channel activity and increased times to maximal activation compared with wild-type STIM1. The ‘super-unstable’ chimera comprised of the STIM2 canonical EF-hand, the STIM1 non-canonical EF-hand and the STIM1 SAM domain (i.e., ES211) activates CRAC channels as efficiently as wild-type STIM1; however, the activation is spontaneous and independent of ER stored Ca\(^{2+}\). Therefore, the regulatory distinctions between STIM1 and STIM2 are facilitated, at least in part, by a balance between the EF-hand Ca\(^{2+}\) binding affinity and SAM domain stability; further, STIM1 has evolved as the principal SOCE regulator with a higher binding affinity and lower SAM stability, rendering it robustly responsive to agonist-induced ER Ca\(^{2+}\) store depletion, whereas STIM2 has evolved as a regulator of basal Ca\(^{2+}\) homeostasis with a lower binding affinity and higher SAM stability rendering a fraction of molecules persistently active at resting ER Ca\(^{2+}\) and the remaining fraction less responsive to small fluctuations in ER Ca\(^{2+}\) store depletion.

**CRAC-Activating Domain Structure and Function**

The cytosolic domains of STIM1 contain the machinery required to couple to and gate Orai1 channels on the PM. Three independent groups identified similar boundaries within the conserved CC region as the minimal CRAC channel activating fragments with the CRAC activating domain (CAD, residues 342–448), STIM-Orai activating region (SOAR, residues 344–442) and coiled-coil boundary fragment 9 (ccb9, residues 346–450). Co-overexpression of any of these fragments with Orai1 in live cells constitutively assembles and gates Orai1 channels. At resting ER Ca\(^{2+}\) levels, these regions are inaccessible by Orai1 and/or in a quiescent conformation in the full-length context of STIM1. Ca\(^{2+}\)-depletion-induced oligomerization of full-length STIM1, initiated by the N-terminal ER luminal domains and apposition of the CC1 region induces a constricted-to-extended structural change in the cytosolic domains which releases CAD/SOAR/ccb9 in a conformation which is conducive to recruiting and gating Orai1 CRAC channels at ER-PM junctions.
Figure 3. Human and C. elegans structures of the cytosolic CAD/SOAR/ccb9 domains. (A) Crystal structure of a Leu374Met/Val419ala/Cys436Thr triple mutant human CAD/SOAR/ccb9 dimer. The 4 α-helices making each monomer are labeled. The residues which were mutated to stabilize the dimer and facilitate crystallization are shown as sticks (blue). The region of intramolecular supercoiling between CC2 and CC3 helices are shaded teal. The intermolecular angle between the CC2 helices at the Tyr361 pivot point is indicated. (B) Crystal structure of the C. elegans CC1-CAD/SOAR/ccb9 dimer. The 2 α-helices making up each monomer are labeled. The CC1 helix demonstrated to modulate STIM1 activation is shaded magenta. Unresolved regions of low electron density are shown as broken black lines. In (A and B), the amino and carboxy termini are denoted by N and C, respectively. The human and C. elegans structure images were created with 3TEQ.pdb and 3TER.pdb coordinates, respectively.

Figure 4. D. melanogaster Orai channel structure. (A) Cytosolic view of the Orai hexamer structure in the presumably closed state. An individual dimer unit building block is bounded by a broken black box. The TM1, TM2, TM3 and TM4 segments from a single monomer are labeled. (B) Residue composition of the TM1-constituted pore region. Only 2 TM1 segments exhibiting the greatest separation are shown for clarity. The acidic (red), hydrophobic (green), and basic (blue) pore-lining residues are shown relative to the extracellular space and the cytosol. The residue position mutated in a heritable form of severe combined immunodeficiency disease (i.e., r 91W in human numbering) is labeled as ‘SCID’. The direction of the Ca\(^{2+}\) gradient (i.e., high to low concentration) is indicated with an arrow. (C) The TM4 C-terminal extension within the dimer unit. The antiparallel interaction between the C-terminal extensions is shown with hydrophobic residues involved in stabilizing this dimer interface depicted as sticks (green). The hinge regions responsible for creating the antiparallel orientation of the C-terminal extensions are indicated. In (A–C), color is consistent with Figure 2, and the amino and carboxy termini are labeled N and C, respectively. The D. melanogaster structure images were created with the 4HKR.pdb coordinate file.
The crystal structure of a protein construct similar to human STIM1 CAD/SOAR/ccb9 (i.e., residues 345–444) bearing a triple Leu374Met/Val419Ala/Cys436Thr mutation that stabilizes the dimeric state was solved at -1.9 Å. Although 3 distinct dimer interfaces were identified in the asymmetric unit, the biologically relevant dimer was defined from the interface with the largest buried surface area (i.e., -1800 Å²). The dimerizer identified in this manner shows the 2 monomers arranged in a 2-fold symmetrical V-shape (Fig. 3A). Consistent with this conformation, an -2.6 Å crystal structure of Caenorhabditis elegans STIM cytosolic residues 212-410 (i.e., aligned with human residues -233-465) identified from a single dimer interface in the asymmetric unit displays a similar symmetrical V-shape (Fig. 3B). Each monomer in the human protein contains a single extended α-helix in the putative CC2 region followed by 2 short and 1 extended α-helix (i.e., α2-α4) in succession and antiparallel relative to α1 within predicted CC3; however, the C. elegans structure exhibits only a single extended α-helix and unstructured polypeptide chain in the region corresponding to the α2 and α3 human helices, raising questions about the relevance of the short α-helices in the human CC3 region (Fig. 3A and B). Analysis of the human V-shaped dimer using the SOCKET program which unambiguously and objectively identifies CC regions based on atomic resolution side chain packing reveals a single region of intramolecular left-handed coiling between 2 helices (i.e., supercoiling) involving residues Lys366-Ala376 of CC2 and Ile409-Ala419 of CC3 within each monomer (Fig. 3A). It is noteworthy that no intermolecular CC interactions were identified in the human STIM1 CAD/SOAR/ccb9 dimer. Triple mutants aimed at perturbing the dimer interface in the CC2 (i.e., Leu347Ala/Trp350Ala/Leu351Ala) and CC3 regions (i.e., Trp430Ala/Ile433Ala/Leu436Ala) abolish co-localization of this STIM1 fragment with Orai1; however, the effects on dimerization in the fragmental and full-length STIM1 context have not been determined.

Interestingly, the larger C. elegans structure reveals an α-helix in putative CC1 (i.e., C. elegans residues 257-279), flanked by unstructured regions exhibiting low electron density; further, this helix packs against both extended helices corresponding to CC2 and CC3 in an intramolecular and intermolecular manner, respectively (Fig. 3B). Since deletion of the equivalent CC1 helix in human STIM1 results in constitutive activation of CRAC channels when co-overexpressed with Orai1 in live cells, this helix was initially labeled as ‘inhibitory helix’. However, more recent functional analyses have shown that mutations in this CC1 helix, rather than full deletion, can both constitutively activate as well as inhibit CRAC entry in live cells. Therefore, this region of CC1 likely plays an important modulatory role in the quiescent-to-active conformational change that occurs in the cytosolic domains, requisite for the recruitment and activation of Orai CRAC channels. While only intramolecular supercoiling was revealed in the cytosolic domain crystal structures, recent solution NMR studies in the Ikura laboratory (Ontario Cancer Institute) has shown that the cytosolic domains of STIM1 are also capable of forming intermolecular supercoils between CC1 as well as CC2 regions (Stathopoulos PB and Ikura M, unpublished data). Taken together, the different interactions elucidated in the apo crystal structures and via solution NMR suggest that dynamic CC interplay is involved in the transition of the cytosolic STIM domains from a quiescent to Orai activation-competent conformation.

**Drosophila melanogaster Orai Structure and Channel Formation**

At present, there are no human Orai channel high resolution structures; nevertheless, a crystal structure of *D. melanogaster* Orai has been determined in which the native protein, consisting of 351 residues, was N- and C-terminally truncated in a construct encompassing residues 132–341 and carrying a Cys224Ser/Cys283Thr/Pro276Arg/Pro277Arg quadruple mutation to produce well-diffracting crystals (i.e., ~3.4 Å). Importantly, this *D. melanogaster* protein shares >70% sequence identity with human Orai1 through the TM regions and, after reconstitution in liposomes, has the ability to conduct Ca²⁺ when constitutively activated with the Val174Ala (i.e., Val102Ala in human Orai1 numbering) mutation. Remarkably, this Orai crystal structure is formed by a trimer of dimer unit building blocks, exhibiting a 6-fold central axis of symmetry along the pore and overall 3-fold symmetry (Fig. 4A). The hexameric quaternary state is in contrast to several studies suggesting that Orai1 assembles as a functional tetramer. The TM1 helices line the pore in the hexameric structure, and a ring composed of Glu178 (i.e., Glu106 in human numbering) vital for Ca²⁺ binding and ion permeability is located near the extracellular surface of the assembled channel (Fig. 4B). The pore widens near the intracellular side and is lined by basic residues (i.e., Arg155, Lys159 and Lys163, corresponding to Arg83, Lys87, and Arg91 in human Orai1 numbering) (Fig. 4B). Sandwiched between the anionic and cationic pore regions is a hydrophobic section made up of Leu167, Phe171, and Val174 (i.e., corresponding to human Orai1 Leu95, Phe99, Val102) (Fig. 4B). This acidic-hydrophobic-basic hierarchy implies a mechanism of permeation where extracellular Ca²⁺ is attracted to and binds the oppositely charged anionic ring; moreover, the widening of the pore would permit Ca²⁺ to move relatively freely into the central hydrophobic segment. It has been suggested that the basic region at the pore outlet binds anions, and these anions would have to be displaced for Ca²⁺ to permeate through to the cytosol. A widening of the pore on the intracellular side could dilute the electropositive potential facilitating the loss of the anions and permitting Ca²⁺ to diffuse into the cytosol. Interestingly, the Lys163Trp mutant *D. melanogaster* structure, crystallized using the same construct as the pseudo-wild-type and corresponding to human Arg91Trp that causes severe combined immunodeficiency, has a remarkably similar structure (i.e., Cα atom root mean square deviation of ~0.1 Å) as the pseudo-wild-type. The Lys163Trp residues extend the hydrophobicity of the central channel region and point into the center axis of the pore. It is tempting to speculate that the loss in function caused by the Lys163Trp (i.e., Arg91Trp in human numbering) mutation is due to an inability to undergo pore dilation required for Ca²⁺ permeability, restricted by inter-subunit Lys163Trp hydrophobic...
interactions. Although the precise role, if any, of anion binding to Ca\(^{2+}\)-permeation through Orai channels has not been established, it should be noted that the Lys163Trp mutation abolishes at least one anion binding site.\(^{51}\)

STIM1 CAD/SOAR/ccb9 interactions with both the N- and C-terminal Orai cytosolic regions are required for the recruitment and gating of the channel at ER-PM junctions.\(^{35-40}\) In the \textit{D. melanogaster} crystal structure, the TM1 helix of each subunit linearly extends beyond the predicted depth of the inner PM leaflet into the cytosol without any breaks in the continuity of the helix (Fig. 4B). Further, studies show that human Orai remains active after deletion of residues 1–73; however, deletion of residues 73–84, corresponding to 145–156 in the \textit{D. melanogaster} Orai structure that comprise most of the N-terminal extension (i.e., residues 144–157), abolishes interactions with CAD and eliminates CRAC entry,\(^{39}\) suggesting that the TM1 helical extension plays an important role in binding to STIM and the mechanism of Orai activation.

The C-terminal helices are located on the periphery of the hexameric channel and also extend into the predicted cytosol; however, the cytosolic TM4 extensions which maintain an \(\alpha\)-helical conformation run mostly parallel to the inner plane of the PM due to a hinge (i.e., residues 305–308) which bends the extensions (Fig. 4C). These cytosolic TM4 extensions adopt 1 of 2 conformations in the dimer unit building blocks such that the \(\alpha\)-helices pair in an antiparallel manner via hydrophobic interactions (Fig. 4C). Critical residues which stabilize the inter-subunit antiparallel C-terminal domain interaction include Ile316 which packs against Leu319 of the opposite subunit (i.e., Phe270 and Leu273 in human numbering). Considering that deletion of the C-terminal cytosolic extension abrogates interactions with STIM1,\(^{55,67,68}\) the antiparallel helix pair may represent a recruitment handle for Ca\(^{2+}\)-depleted and oligomerized STIM1 at ER-PM junctions.

\section*{Ca\(^{2+}\)-CaM:Orai1-N Structure and CRAC Entry Regulation}

Ca\(^{2+}\)-dependent inactivation (CDI) of CRAC channels is an important regulatory mechanism that confers limits on local Ca\(^{2+}\) concentrations influenced by SOCE; moreover, both STIM as well as calmodulin (CaM) have been linked to CDI of CRAC channels.\(^{14,69,70}\) STIM1 residues 470–491 are required for fast CDI of CRAC channels, although it is unclear whether this region serves as a bona fide Ca\(^{2+}\) sensor involved in the inactivation mechanism.\(^{70}\) Nevertheless, it has been demonstrated that CaM binds to the Orai1 N-terminal domain residues 68–91 in a Ca\(^{2+}\) dependent manner, and this CaM interaction is necessary for CDI of CRAC channels.\(^{70}\) A crystal structure of Ca\(^{2+}\)-CaM in complex with an Orai1 construct encompassing N-terminal residues 69–91 solved at 1.9 Å shows that CaM is in the well-characterized dumbbell conformation, with an extended \(\alpha\)-helix linking the 2 EF-hand lobes that coordinate 2 Ca\(^{2+}\) ions each (Fig. 5A).\(^{71}\) The Orai1-N peptide binds to the hydrophobic pocket created by the CaM-C lobe and retains the \(\alpha\)-helical conformation observed in the \textit{D. melanogaster} Orai crystal structure (see above); further, the CaM-N lobe remains free of Orai1-N in the crystal structure. The CaM-C:Orai1-N interaction is stabilized primarily by hydrophobic side chain packing. In particular, the Trp76, Leu79 and Tyr80 residues of Orai1-N bind in a cleft formed by the CaM-C lobe and retains the \(\alpha\)-helical conformation observed in the \textit{D. melanogaster} crystal structure (see above); further, the CaM-N lobe remains free of Orai1-N in the crystal structure. The CaM-C:Orai1-N interaction is stabilized primarily by hydrophobic side chain packing.

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\caption{Figure 5. Rat Ca\(^{2+}\)-CaM structure in complex with a human Orai1 N-terminal fragment. (A) Dumbbell structure of Ca\(^{2+}\)-loaded CaM forming a complex with a fragment of the Orai1 N-terminal domain (i.e., residues 69–91) through interactions at the C-lobe. The locations of the CaM lobes, central linker helix between lobes and the Orai1 N-terminal helix (beige) are labeled. (B) The Ca\(^{2+}\)-CaM C-lobe hydrophobic cleft. The residues constituting the hydrophobic cleft are depicted as purple sticks. The anchor residues from the Orai1 N-terminal fragment which pack into the CaM C-lobe hydrophobic cavity are shown as red sticks. In (A and B), the amino and carboxy termini are labeled N and C, respectively, and the Ca\(^{2+}\) atoms are shown as orange spheres. The complex structure images were created with The PyMOL Molecular Graphics System, Version 1.5.04 Schrödinger, LLC.}
\end{figure}
(i.e., residues 70–91; $K_d$ ~250 μM). These stark differences in affinity for the Orai N-terminal domain advocate a CaM-dependent mechanism of CDI that is, at least in part, mediated by the free energy differences of the STIM1 and CaM complexes with Orai1-N. Increased Ca$^{2+}$ levels local to the CRAC channel complex will favor the more stable Ca$^{2+}$-CaM:Orai1-N interaction over STIM1:Orai1-N; however, it is tempting to speculate that, in addition to the displacement of the STIM1 cytosolic domains from Orai1-N by Ca$^{2+}$-CaM, CDI may also involve the bridging of two Orai1-N domains by a single Ca$^{2+}$-CaM molecule, thereby constricting the channel pore. It should also be noted that Ca$^{2+}$-CaM can bind to the polybasic tails of both STIM1 (i.e., residues 667–685) and STIM2 (i.e., residues 730–746) with $K_d$ -μM; moreover, this affinity is reduced by 2 orders of magnitude in the absence of Ca$^{2+}$. Therefore, CaM may also downregulate SOCE via binding to STIMs in a Ca$^{2+}$-dependent manner after localized cytosolic Ca$^{2+}$ level increases, thereby inhibiting and/or disrupting ER-PM targeting of STIM1 molecules which is dependent on the polybasic stretches of STIMs.

**Concluding Remarks**

The current available high resolution data of CRAC channel components and regulators have revealed a wealth of knowledge on the mechanisms controlling SOCE. In particular, the Ca$^{2+}$-loaded STIM1 and STIM2 EF-SAM solution NMR structures have exposed the bases by which the luminal domains distinctly sense changes in ER stored Ca$^{2+}$, signaling the cytosolic domains to induce CRAC entry and maintain basal intracellular Ca$^{2+}$ homeostasis, respectively, via communication with Orai proteins; moreover, the balance between the EF-hand affinity for Ca$^{2+}$ and SAM domain stability has evolutionarily defined these specific physiological roles for each isoform in mammals. Further, differences in the CAD/SOAR/ccb9 (i.e., CC2-CC3), CC1-CAD/SOAR/ccb9 crystal structures and solution NMR data on CC1-CC2 fragments (Stathopulos P.B. and Ikura M., Ontario Cancer Institute, unpublished data), both solved in the absence of Orai, are in line with the dynamic closed-to-open transition that the STIM1 cytosolic domains must undergo to adopt an Orai recruitment- and activation-competent conformation. The closed hexameric *D. melanogaster* Orai channel crystal structure reveals an extracellular-to-intracellular, anionic-hydrophobic basic pore hierarchy that mediates Ca$^{2+}$ permeation; moreover, the TM1 helices that make up the pore contiguously extend into the cytosol exposing vital binding sites for, at least, STIM$^{35,38-40}$ and CaM$^{70}$ regulatory partners. Each lobe of CaM is capable of binding one TM1 extension with higher apparent affinity than the STIM1 cytosolic domain during CDI. The TM4 helices of *D. melanogaster* Orai also extend into the cytosol, stabilizing Orai dimer units through antiparallel intermolecular interactions. The antiparallel C-terminal domain extensions of TM4 make up a recruitment-like handle which may present Orai dimer C-terminal binding sites to STIM molecules at ER-PM junctions during the activation process.

Despite the tremendous progress in our comprehension on the structural biology of the CRAC entry molecular players, numerous targets of prime importance remain. For example, minimal high resolution structural information exists on the apo (i.e., Ca$^{2+}$-free) luminal STIM domains which initiate the process of SOCE. Further, the structural basis for STIM CC3:CC3’ interactions involved in oligomerization is lacking, and it is currently unknown how or even whether CC3 directly interacts with the Orai proteins during the activation process. The human Orai channel structure in either the closed or open state has not been determined; moreover, the basis for Orai1 N-terminal or C-terminal interactions with STIM1 which have been implicated in channel recruitment and gating has not been disseminated. Additionally, the exact stoichiometry in the functional STIM:Orai complex has not been elucidated. The non-conserved far C-terminal domains of STIM are involved in targeting to ER-PM junctions via phosphoinositide binding, and high resolution structural data on the basis for this targeting is also missing.

In addition to these questions on static states, vital information on dynamic processes involved in the activation of CRAC channels remains outstanding. How is the Ca$^{2+}$ binding event in the luminal domains transmitted to the cytosolic side of STIM in order to activate Orai? What conformational change in Orai is required for channel opening and how does STIM association induce this transformation? How do the numbers of active STIM:Orai complexes change at ER-PM junctions during the course of SOCE activation and how do these clusters of multiple active channels effect the functional properties of CRAC entry. Future studies on these aforementioned structural targets and questions are needed to paint a more complete picture of the molecular mechanisms governing SOCE which is vital to countless physiological processes in all eukaryotes.

**Disclosure of Potential Conflicts of Interest**

No potential conflicts of interest were disclosed.

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**References**

1. Bootman MD, Lipp P. (2001). Calcium signalling and regulation of cell function. *Encyclopedia of Life Sciences, 1-7.*
2. Berridge MJ. (2009). Cell Signalling Biology, Vol. 2009. Portland Press, Ltd., London.
3. Berridge MJ, Bootman MD, Roderick HL. Calcium signalling: dynamics, homeostasis and remodeling. *Nat Rev Mol Cell Biol 2003; 4:517-29; PMID:12838355; http://dx.doi.org/10.1038/ nrm1395*
4. Frede S. Calcium signalling in lymphocyte activation and disease. *Nat Rev Immunol 2007; 7:690-702; PMID:17703229; http://dx.doi.org/10.1038/nri2152*
5. Stathopulos PB, Seo MD, Enomoto M, Amador FJ, Iishiyma N, Ikura M. Themes and variations in ER/SR calcium release channels: structure and function. *Physiology (Bethesda) 2012; 27:351-42; PMID:23233627; http://dx.doi.org/10.1152/ physiol.00013.2012*
6. Berridge MJ. *The endoplasmic reticulum: a multifunctional signalling organelle.* Cell Calcium 2002; 32:235-49; PMID:12543086; http://dx.doi.org/10.1006/lPe.2006.001823
A nature04147
STIM1 is a Ca2+ sensor that activates CRAC channels by a STIM1 coiled-coil mutant. Therefore, transmembrane segment 1 cause STIM1-independent release-activated Ca(2+) (CRAC) current, structure, and function. CRAC channel activity. J Biol Chem 2006; 281:21088-96; PMID:16588458.

20. Varnai P, Emr SM, Tkatch AT, Ikura M, Gething MJ, Schindl P, et al. Identification and functional characterization of the Ca(2+)-sensing domain of STIM1. J Biol Chem 2003; 278:47766-75; PMID:12963858.

21. Gouaux E, Kang E, Weis WI, Kobilka BK, Held M, Schertler GF, et al. Structure, function, and evolution of the voltage sensor of voltage-gated ion channels. Cell 2005; 121:935-54; PMID:15733836.

22. Zhu Z, Liu C, Zhou W, Liu H, Zhang J, Liu P, et al. Identification and functional characterization of the Ca(2+)-sensing domain of STIM1. J Biol Chem 2003; 278:47766-75; PMID:12963858.

23. Gouaux E, Kang E, Weis WI, Kobilka BK, Held M, Schertler GF, et al. Structure, function, and evolution of the voltage sensor of voltage-gated ion channels. Cell 2005; 121:935-54; PMID:15733836.

24. Gouaux E, Kang E, Weis WI, Kobilka BK, Held M, Schertler GF, et al. Structure, function, and evolution of the voltage sensor of voltage-gated ion channels. Cell 2005; 121:935-54; PMID:15733836.
58. Covington ED, Wu MM, Lewis RS. Essential role for the CRAC activation domain in store-dependent oligomerization of STIM1. Mol Biol Cell 2010; 21:1897-907; PMID:20357143; http://dx.doi.org/10.1091/mbc.E10-02-0145

59. Korzeniowski MK, Popovic MA, Szepetnyzer Z, Varnai P, Stojilkovic SS, Balla T. Dependence of STIM1/ORAI-mediated calcium entry on plasma membrane phosphoinositides. J Biol Chem 2009; 284:21027-35; PMID:19483082; http://dx.doi.org/10.1074/jbc.M109.012252

60. Stathopoulos PB, Ikura M. Partial unfolding and oligomerization of stromal interaction molecules as an initiation mechanism of store operated calcium entry. Biochem Cell Biol 2010; 88:175-83; PMID:20453920; http://dx.doi.org/10.1139/O09-125

61. Stathopoulos PB, Ikura M. Structurally delineating stromal interaction molecules as the endoplasmic reticulum calcium sensors and regulators of calcium release-activated calcium entry. Immunol Rev 2009; 231:113-31; PMID:19754893; http://dx.doi.org/10.1111/j.1600-065X.2009.00814.x

62. Huang GN, Zeng W, Kim JY, Yuan JP, Han L, Mualem S, Worley PF. STIM1 carboxyl-terminus activates native SOCE, (Icra) and TRPC1 channels. Nat Cell Biol 2006; 8:1003-10; PMID:16906149; http://dx.doi.org/10.1038/nclb454

63. Kawasaki T, Lange I, Feske S. A minimal regulatory domain in the C terminus of STIM1 binds to and activates ORAI1 CRAC channels. Biochem Biophys Res Commun 2009; 385:49-54; PMID:19433061; http://dx.doi.org/10.1016/j.bbrc.2009.05.020

64. Yang X, Jin H, Cai X, Li S, Shen Y. Structural and mechanistic insights into the activation of Sromol interac in STIM1. Proc Natl Acad Sci U S A 2012; 109:5657-62; PMID:22491994; http://dx.doi.org/10.1073/pnas.1118947109

65. Walshaw J, Woolfson DN. Sockey: a program for identifying and analysing coiled-coil motifs within protein structures. J Mol Biol 2001; 307:1427-50; PMID:11292393; http://dx.doi.org/10.1006/jmbi.2000.4545

66. Feske S, Gwack Y, Prakriya M, Srikanth S, Puppel SH, Tanasa B, Hogan PG, Lewis RS, Daly M, Rao A. A mutation in Orail causes immune deficiency by abrogating CRAC channel function. Nature 2006; 441:179-85; PMID:16582901; http://dx.doi.org/10.1038/nature04702

67. Zheng H, Zhou MH, Hu C, Kuo E, Peng X, Hu J, Kuo L, Zhang SL. Differential roles of the C and N termini of Orail1 protein in interacting with stromal interaction molecule 1 (STIM1) for Ca2+ release-activated Ca2+ (CRAC) channel activation. J Biol Chem 2013; 288:11263-72; PMID:23447534; http://dx.doi.org/10.1074/jbc.M113.450254

68. McNally BA, Somasundaram A, Jairaman A, Yamashita M, Prakriya M. The C- and N-terminal STIM1 binding sites on Orail1 are required for both trapping and gating CRAC channels. J Physiol 2013; 591:2833-50; PMID:2361525

69. Derler I, Fahrner M, Muik M, Lackner B, Schindl R, Groschner K, Romainin C. A Ca2+-release-activated Ca2+ (CRAC) modulatory domain (CMD) within STIM1 mediates fast Ca2+-dependent inactivation of ORAI1 channels. J Biol Chem 2009; 284:24933-8; PMID:19622747; http://dx.doi.org/10.1074/jbc.C109.024083

70. Mullins FM, Park CY, Dolmetsch RE, Lewis RS. STIM1 and calmodulin interact with Orail1 to induce Ca2+-dependent inactivation of CRAC channels. Proc Natl Acad Sci U S A 2009; 106:15495-500; PMID:19766428; http://dx.doi.org/10.1073/pnas.0906781106

71. Liu Y, Zheng X, Mueller GA, Sobhany M, DeRose EF, Zhang Y, London RE, Birnbaumer L. Crystal structure of calmodulin binding domain of orail in complex with Ca2+ calmodulin displays a unique binding mode. J Biol Chem 2012; 287:43030-41; PMID:23109337; http://dx.doi.org/10.1074/jbc.M112.380964

72. Derler I, Plenk P, Fahrner M, Muik M, Jardin I, Schindl R, Gruber HJ, Groschner K, Romainin C. The extended transmembrane Orail1 N-terminal (ETON) region combines binding interface and gate for Orail1 activation by STIM1. J Biol Chem 2013; 288:29025-34; PMID:23943609; http://dx.doi.org/10.1074/jbc.M112.390530

73. Bauer MC, O’Connell D, Cahill DJ, Linse S. Calmodulin binding to the polybasic C-termini of STIM proteins involved in store-operated calcium entry. Biochemistry 2008; 47:6089-91; PMID:18484746; http://dx.doi.org/10.1021/bi800496a

74. Calloway N, Owens T, Corwitt K, Rodrigers W, Holowka D, Baird B. Stimulated association of STIM1 and Orail is regulated by the balance of PtdIns(4,5)P2 between distinct membrane pools. J Cell Sci 2011; 124:2602-10; PMID:21750194; http://dx.doi.org/10.1242/jcs.084178

75. Walsh CM, Chovanov M, Haynes LP, Petersen OH, Tepikin AV, Burgoyne RD. Role of phosphoinositides in STIM1 dynamics and store-operated calcium entry. Biochem J 2010; 425:159-68; PMID:19843011; http://dx.doi.org/10.1042/Bj20090884