Coordination of a Single Calcium Ion in the EF-hand Maintains the Off State of the Stromal Interaction Molecule Luminal Domain

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Abstract

Store operated calcium (Ca\(^{2+}\)) entry (SOCE) is the process whereby endoplasmic reticulum (ER) Ca\(^{2+}\) store depletion causes Orai1-composed Ca\(^{2+}\) channels on the plasma membrane (PM) to open, mediating a rise in cytosolic Ca\(^{2+}\) levels. Stromal interaction molecules (STIMs) are the proteins that directly sense ER Ca\(^{2+}\) content and gate Orai1 channels due to store depletion. The trigger for STIM activation is Ca\(^{2+}\) unbinding from the ER lumen-oriented domains, which consist of a nonconserved amino (N) terminal region and EF-hand and sterile a motif (SAM) domains (EF\(_e\)SAM), highly conserved from humans to Caenorhabditis elegans. Solution NMR structures of the human EF\(_e\)SAM domains have been determined at high Ca\(^{2+}\) concentrations; however, no direct structural view of the Ca\(^{2+}\) binding mode has been elucidated. Further, no atomic resolution data currently exists on EF\(_e\)SAM at low Ca\(^{2+}\) levels. Here, we determined the X-ray crystal structure of the C. elegans STIM luminal domain, revealing that EF\(_e\)SAM binds a single Ca\(^{2+}\) ion with pentagonal bipyramidal geometry and an ancillary \(\alpha\)-helix formed by the N-terminal region acts as a brace to stabilize EF\(_e\)SAM. Using solution NMR, we observed EF-hand domain unfolding and a conformational exchange between folded and unfolded states involving the ancillary \(\alpha\)-helix and the canonical EF-hand in low Ca\(^{2+}\). Remarkably, we also detected an \(\alpha\)-helix (−Ca\(^{2+}\)) to \(\beta\)-strand (−Ca\(^{2+}\)) transition at the terminal SAM domain \(\alpha\)-helix. Collectively, our analyses indicate that one canonically bound Ca\(^{2+}\) ion is sufficient to stabilize the quiescent luminal domain structure, precluding unfolding, conformational exchange, and secondary structure transformation.

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In nonexcitable cells, SOCE can be stimulated downstream of any cell surface G protein-coupled receptor (GPCR) activation that results in inositol 1,4,5-trisphosphate (IP$_3$) production and IP$_3$ receptor channel opening on the ER, depleting ER Ca$^{2+}$ levels. However, any mode of ER Ca$^{2+}$ depletion will stimulate SOCE. STIMs contain EF-hand and sterile $\alpha$-motif (SAM) domains oriented in the ER lumen and a series of coiled-coil domains in the cytosol. These luminal and cytosolic domains have a high-sequence similarity among STIM orthologs [15]. A single pass transmembrane (TM) domain separates the luminal from the cytosolic conserved domains.

STIM1 forms dimers on the ER membrane due to interactions by the cytosolic coiled-coil domains [19–24]. These inactive STIM1 dimers pervasively diffuse along the ER [25]. When the ER Ca$^{2+}$ stores are full, the luminal domains remain monomeric due to Ca$^{2+}$-binding dependent stabilization of the EF-hand and SAM domain (EF–SAM) intramolecular interaction [26–29]. When the ER Ca$^{2+}$ stores are depleted, this luminal intramolecular interaction is destabilized, leading to EF–SAM dimerization [26,27,30–32]. This EF–SAM dimerization induces a marked conformational rearrangement of the cytosolic coiled-coils [19,24,33] that leads to trapping of activated STIM1 molecules at ER-PM junctions [25,34]. The trapping of STIM1 occurs due to a release of a polybasic C-terminal region of STIM1 that affords interactions with PM phosphoinositides [35–38] and due to direct interactions by the STIM1 coiled-coils with Orai1 protein subunits [39,40]. These direct STIM1—Orai1 interactions gate the Orai1 channels.

High-resolution structures of the human STIM1 and STIM2 EF–SAM have been determined by solution NMR only in a high molar excess of Ca$^{2+}$ (high Ca$^{2+}$) [27,29]. However, distance restraints coordinating Ca$^{2+}$ were not directly determined in these structures. Furthermore, these human structures lacked the variable N-terminal regions shown to play a modulatory role in the Ca$^{2+}$-sensing mechanism of STIM1 and STIM2 [32,41,42]. Therefore, the precise mechanism of Ca$^{2+}$ coordination, how the variable N-terminal region of STIMs affect core EF–SAM structure, and mechanism of STIM luminal domain dimerization in buffer nominally free of Ca$^{2+}$ (low Ca$^{2+}$) remain major knowledge gaps in the field.

Here, we present the crystal structure of the Caenorhabditis elegans STIM (cSTIM) luminal domain in the presence of high Ca$^{2+}$, revealing a single Ca$^{2+}$ ion coordinated in the canonical EF-hand loop with pentagonal bipyramidal geometry and that the nonconserved N-terminal region forms an ancillary $\alpha$-helix that further stabilizes the core EF–SAM structure. In addition, we used solution NMR to probe the structure and folding of this C. elegans luminal domain in low Ca$^{2+}$, discovering that the EF-hand and ancillary $\alpha$-helices are in conformational exchange between folded and unfolded states. Finally, our NMR data show that the C-terminal helix of the luminal SAM domain, which is intimately involved in maintaining the intramolecular EF-hand:SAM domain interaction in the presence of high Ca$^{2+}$, undergoes a remarkable $\alpha$-helix to $\beta$-strand transition in low Ca$^{2+}$. Collectively, our crystal and solution NMR data provide previously unappreciated atomic level insights into the mechanisms underlying STIM activation.

Results

C. elegans EF–SAM is structurally similar to human STIM2 EF–SAM

In high Ca$^{2+}$, the human STIM1 and STIM2 EF-hand and SAM domains fold into a compact structure made up of ten $\alpha$-helices and two loops, whereas the SAM domain adopts a compact five-helix bundle. In the Ca$^{2+}$-bound state, the terminal SAM helix (i.e., $\alpha$10) packs into a hydrophobic cleft made up of the two EF-hand motifs, thereby promoting the overall compact configuration. This conformation is further buttressed via the small $\beta$-sheet formed by backbone hydrogen (H) bonding between loop I and II of the EF-hand motifs.

Here, our C. elegans crystal structure (1.89 Å resolution) reveals the EF–SAM region of this lower eukaryotic STIM conserves these characteristic features central to regulating the activation state of the molecule and SOCE (Fig. 1A). The crystal structure determined in high Ca$^{2+}$ shows a canonical helix–loop–helix motif ($\alpha$1–$\beta$1–$\alpha$2). Immediately downstream of the canonical EF-hand motif is a second helix–loop–helix motif ($\alpha$3–$\beta$2–$\alpha$4), which pairs with the first EF-hand through backbone H-bonding between $\beta$1 and $\beta$2, forming a short $\beta$-sheet. The second, noncanonical EF-hand loop is not clearly identifiable based on primary structure for neither the human STIMs nor C. elegans (Fig. 1B). A short linker helix ($\alpha$5) links the C. elegans EF-hand domain in sequence space to the five-helix bundle-composed SAM domain ($\alpha$6–$\alpha$10). This SAM domain alone buries 13 residues that are at least 85% solvent inaccessible (Fig. S1).

The EF-hand and SAM domains of the C. elegans STIM form intimate contacts, primarily mediated by nonpolar interactions. In high Ca$^{2+}$, these intimate EF-hand:SAM hydrophobic interactions are a hallmark feature of STIM Ca$^{2+}$ sensing that supports a quiescent conformation [27]. In C. elegans STIM, the EF-hand domain forms a hydrophobic cleft made up...
of at least 13 residues with nonpolar character (Fig. 1C). This extensive hydrophobic pocket serves as a docking site for the hydrophobic L176 and L180 anchor residues found on the α10 helix of the SAM domain (Fig. 1D). In C. elegans, the EF-hand:SAM domain interaction is reinforced through one ionic bond involving R85 within α2 of the noncanonical EF-hand motif and D177 within α10 of the SAM domain (Fig. 1E). These hydrophobic and ionic interactions pull the SAM domain of C. elegans STIM into the EF-hand cleft and promote a compact globular fold.

Despite the continuity in global structure of cSTIM EF−SAM, the electrostatic surface potential is separable based on domain. A highly negative surface potential is apparent over the EF-hand domain, whereas the closely packed SAM domain is slightly electropositive (Fig. 1F). These domain differences in surface potential are exacerbated further when removing the positive charge contribution of the Ca\(^{2+}\) ion from the calculation. The strongly negative surface potential of the EF-hand domain is consistent with a role in the electrostatic guidance of free Ca\(^{2+}\) into the canonical binding loop, which is preserved throughout evolution from C. elegans to humans.

Within the EF-hand domain cleft, the C. elegans SAM domain is rotated away from the canonical EF-hand motif such that the α2 of the canonical EF-hand adopts a nearly parallel orientation relative to the α10 helix of the SAM domain (i.e., −155° α2−α10 interhelical angle; Fig. 2A). This geometry is remarkably similar to that observed for the human STIM2 solution EF−SAM structure in high Ca\(^{2+}\) (i.e., −150° α2−α10 interhelical angle), but distinct from the nearly perpendicular orientation observed for human STIM1 EF−SAM in high Ca\(^{2+}\) (i.e., −94° α2−α10 interhelical angle; Fig. 2B and C) [29]. To accommodate this orientation, both the C. elegans and the human STIM2 EF-hand clefts are expanded (i.e., 13 residues) compared with human STIM1 (i.e., 11 residues); however, the mechanism of the cleft expansion appears to be distinct for C. elegans compared with human STIM2. In human STIM2, W128 and K103 point into the cleft compared with the aligned residues in human STIM1 (i.e., W124 and H99), which are directed away; moreover, the equivalent C. elegans residues (i.e., W105 and R78) are directed away from the cleft similar to STIM1, yet the total number of residues contributing to the cleft is similar to STIM2 due to the unique inclusion of M71 and M77 by C. elegans EF−SAM (Fig. 1C) [29].

**C. elegans EF−SAM coordinates one Ca\(^{2+}\) ion located in the canonical EF-hand loop**

The present C. elegans crystal structure provides the first direct atomic-level view into the mode of Ca\(^{2+}\) binding to the luminal domain of any STIM protein. Examination of the 2F\(_{o}\)−F\(_{c}\) electron density map contoured at 2.0σ shows only a single location of high electron density outside the polypeptide chain that could be attributable to Ca\(^{2+}\); despite the relatively high CaCl\(_2\) concentration (i.e., 10 mM) used for crystallization (Fig. 3A). The coordination of Ca\(^{2+}\) within the canonical EF-hand loop occurs with pentagonal bipyramidal geometry as observed for calmodulin, parvalbumin, and other EF-hand proteins (Fig. 3A) [43,44]. The residues involved in Ca\(^{2+}\) binding within the 12-residue canonical EF-hand loop are D55 (O\(_{g}\), position X), D57 (O\(_{g}\), position Y), S59 (O\(_{γ}\), position Z), S61 (backbone CO, position #), D63 (O\(_{β1}\), position y), and E66 (Oc1 and Oc2, position z). Whereas positions X, Y, and Z provide a single ligand for Ca\(^{2+}\) binding via side chain oxygen atoms, position # contributes to the coordination via a backbone carboxyl oxygen, position y bridges a first coordination shell water molecule through a side chain oxygen atom, and position z provides a bidentate ligand through both side chain oxygen atoms. All seven first coordination shell oxygen ligands are within 2.5 Å of the Ca\(^{2+}\) ion (Fig. 3A).

The Glu at position z (i.e., 12th residue) in the canonical EF-hand loop is critical for the observed pentagonal bipyramidal geometry due to the side chain length and bidentate contribution to the coordination. In other canonical EF-hand containing proteins, when this z position is occupied by Asp, which also has a side chain carboxylate group with two oxygen atoms available for bidentate coordination but has a shorter side chain compared with Glu, Ca\(^{2+}\) is coordinated with octahedral geometry and a preference for Mg\(^{2+}\) is observed [45,46]. In other canonical EF-hand containing proteins, substitution of the Glu in this z position with Gln, which has a side chain of the same length but provides only one oxygen atom for coordination (monodentate), Ca\(^{2+}\) binding affinity is reduced [47,48]. We note that the Glu at position z is highly conserved in the canonical EF-hand of STIMs from higher and lower order organisms [15].

To assess more judiciously the number of possible Ca\(^{2+}\) binding sites within the entire luminal domain, we calculated the F\(_{o}\)−F\(_{c}\) omit map for the dimeric asymmetric unit after removal of the Ca\(^{2+}\) ion from the structure. A series of F\(_{o}\)−F\(_{c}\) omit map views contoured at 2.0, 4.0, 6.0, 8.0, and 16.0σ shows a strong unaccounted-for electron density only in the canonical EF-hand loop, which is persistent through all contour levels shown (Fig. S2A−E). A closer comparison of the omit map in the EF-hand loops reinforces the unaccounted-for electron density in the canonical EF-hand loop contoured at 9.0σ (Fig. 3A), but not the noncanonical EF-hand loop contoured at 9.0σ or 2.0σ (Fig. 3B). Thus, C. elegans STIM EF−SAM stably coordinates only one Ca\(^{2+}\) ion.
Fig. 1. X-ray crystal structure features of Ca\(^{2+}\)-bound Caenorhabditis elegans STIM luminal domain. (A) Cartoon view of the EF–SAM structural fold in the context of the full C. elegans luminal domain (cSTIM\(_{L\text{full}}\)). The SAM domain (brown) docks into a cleft formed by the canonical EF-hand (cyan) and noncanonical EF-hand (light blue) motifs. The EF-hand and SAM domains are linked by the \(\alpha_5\) helix (gray). An ancillary \(\alpha_0\) helix (pink) in the variable N-terminal region is visible. Individual secondary structure components are labeled. (B) Sequence alignment of canonical and noncanonical EF-hand motifs of human and C. elegans STIMs. The secondary structure components of the motifs based on the X-ray crystal structure of cSTIM\(_{L\text{full}}\) are shown above the sequences. The four EF-hand motifs of human calmodulin (hCaM) are shown for comparison. The 12-residue loop of C. elegans STIM are colored red, whereas human STIM1 (hSTIM1), STIM2 (hSTIM2), and hCaM are colored blue. The residues involved in coordination (X, Y, Z, #, x, and z) and the invariant G and I residues are indicated above the loop sequence. The residue ranges are shown at left and right of each sequence based on the accession numbers shown in Table 1 and AAD45181.1 for hCaM. (C) Surface and cartoon view of the EF-hand cleft formed by C. elegans STIM. The 13 cleft forming residues are labeled and shown as yellow sticks. The residues that are
The variable N-terminal region of C. elegans STIM forms an ancillary \( \alpha \)-helix that interacts with both the EF-hand and SAM domains

An isolated peptide derived from the human STIM1 nonconserved N-terminal region (i.e., residues 24–57), upstream of EF–SAM, remains unstructured in solution [49]; nevertheless, paramagnetic relaxation enhancement (PRE) NMR experiments suggest that this peptide interacts with the EF–SAM core near the W121 and K122 residues [42]. Here, our crystal structure of the C. elegans luminal STIM domains reveals the nonconserved N-terminal region forms a previously unseen \( \alpha \)-helix (\( \alpha_0 \)), upstream of the EF–SAM core (Fig. 4A). This \( \alpha_0 \) helix is stabilized by an intrahelix ionic interaction between E36 and R39 (Fig. 4B). Further, this ancillary \( \alpha \)-helix contacts both the EF-hand and SAM domains of C. elegans. Specifically, the A33 carbonyl oxygen of \( \alpha_0 \) is within 3.0 Å of the W102 indole N\( _1 \) atom of the EF-hand domain and the E36 O\( _2 \) atom is within 2.7 Å of the R111 guanidinium N\( _2 \) atom on the SAM domain (Fig. 4B). Remarkably, the W102 of the C. elegans EF–SAM is equivalent to the W121 position on human STIM1, which is the N-terminal region contact site identified on human STIM1 EF–SAM using PRE [42]. Thus, despite the lack of sequence conservation between the N-terminal regions of STIM homologues from higher to lower eukaryotes, interactions of these regions with the respective EF–SAM core may be a conserved structural feature among all STIM proteins.

The nonconserved N-terminal region of C. elegans and human STIMs enhance the stability of the luminal domain

Destabilizing EF–SAM mutations constitutively activate full-length human STIM1 and SOCE [27]. Further, engineering superstable human STIM1/STIM2 EF–SAM chimeras into full-length STIM1 results in inhibited SOCE, whereas unstable EF–SAM chimeras constitutively activate SOCE [29]. In addition, stabilizing luminal domain post-translational modifications inhibit human STIM1 activation [42], whereas destabilizing modifications promote STIM1 activation and SOCE [50,51]. Having elucidated a previously unidentified \( \alpha_0 \) helix in the luminal domain of C. elegans STIM that contacts the EF–SAM core, and given the demonstrated importance of stability to human STIM1 activation, we then probed the effect of extending human STIM1 and human STIM2 EF–SAM constructs to include the respective variable N-terminal regions on stability. We created human STIM1 and STIM2 EF–SAM constructs that were extended N-terminally (\( N_{\text{ext}} \)). Our C. elegans crystal structure construct encompassed all residues of the luminal domain (\( L_{\text{full}} \)), already including both N- and C-terminal extensions relative to EF–SAM; thus, we created a C. elegans construct with a deleted nonconserved N-terminal region (\( L_{\text{AN}} \)).

Far UV circular dichroism (CD) spectra show that all proteins have enhanced \( \alpha \)-helicity in high Ca\( ^{2+} \) compared with low Ca\( ^{2+} \) (Fig. S3A–D). We monitored the change in far-UV CD signal at 225 nm as a function of increasing temperature as an indicator of stability. In high Ca\( ^{2+} \) (i.e., 5 mM CaCl\( _2 \)), the thermal melts showed that the apparent midpoints of temperature denaturation (\( T_m \)) of the cSTIM–\( L_{\text{full}} \) was ~7.5 °C higher compared with the cSTIM–\( L_{\text{AN}} \) construct with the N-terminal deletion (Fig. S3E and F; Table 1). Consistent with a stabilizing role of the variable N-terminal region, the \( T_m \) values of human STIM1–\( N_{\text{ext}} \) and STIM2–\( N_{\text{ext}} \) were enhanced by ~15.5 and 11.5, respectively, compared with the EF–SAM cores alone (Fig. S3G and H; Table 1). A similar increase in thermal stability was observed under low Ca\( ^{2+} \) conditions with the inclusion of the N-terminal regions stabilizing the human STIM1 and human STIM2 EF–SAM domains by ~15.5 and 14.0 °C, respectively, and deletion of the N-terminal region destabilizing the full cSTIM luminal domain by ~12.5 °C (Fig. S3E–H; Table 1). All proteins showed a dramatic decrease in \( T_m \) in low Ca\( ^{2+} \) compared with high Ca\( ^{2+} \) (i.e., \( \Delta T_m \) ranging from ~11.5 to 24 °C; Table 1).

cSTIM–\( L_{\text{full}} \) in high Ca\( ^{2+} \) showed the highest thermal stability of any ortholog examined (Table 1). To examine whether this enhanced stability was related to a change in Ca\( ^{2+} \) binding affinity, we performed a direct radioactive \( ^{45} \)Ca\( ^{2+} \) binding assay. The apparent equilibrium dissociation constant (\( K_d \)) fitted by the Hill equation was 0.40 ± 0.02 mM (Fig. 5). The Ca\( ^{2+} \) binding profile showed high apparent cooperativity with a Hill coefficient of 3.15 ± 0.31. Furthermore, the curve showed
saturation at a $\text{Ca}^{2+}$:protein molar ratio of 1.05 ± 0.02, indicating a single $\text{Ca}^{2+}$ binding site, consistent with our crystal data (Fig. 5). The apparent positive cooperativity may be due to a coupling between $\text{Ca}^{2+}$ binding and protein folding, a highly cooperative process for globular proteins. The fitted $K_d$ for the $C.\text{elegans}$ STIM luminal domain suggests a sub-mM affinity, similar to the ~0.2–0.6 mM and ~0.5–0.8 mM affinities measured for human STIM1 and STIM2 EF–SAM proteins, respectively [26,29]. Thus, the structural features adopted by the $C.\text{elegans}$ luminal domain in the $\text{Ca}^{2+}$ bound state play a dominant role in conferring the highest stability of any of the STIM proteins studied to date.

**The $C.\text{elegans}$ STIM luminal domain shows folded and unfolded conformations in low $\text{Ca}^{2+}$**

Having established that human STIM1, human STIM2, and $C.\text{elegans}$ luminal STIM proteins that are extended to include the respective variable N-terminal domains have enhanced stability compared with the EF–SAM core proteins alone in low $\text{Ca}^{2+}$, we acquired solution $^1\text{H}–^{15}\text{N}$-heteronuclear single quantum coherence (HSQC) spectra of each of the extended proteins to reveal structural details of the conformations. In the presence of excess $\text{Ca}^{2+}$, all proteins showed a well-dispersed $^1\text{H}–^{15}\text{N}$-HSQC spectrum with the expected number of amide H(N) crosspeaks and relatively homogeneous intensities (Fig. S4A–C). In low $\text{Ca}^{2+}$, STIM1–$N_{\text{ext}}$ showed severe peak broadening and a collapse of all visible H(N) peaks into the central ~7.5–8.5 ppm $^1\text{H}$ region of the spectrum, even at relatively low protein concentration (i.e., ~50 µM) and temperature (i.e., 10 °C) (Fig. S4A). On the other hand, the H(N) crosspeaks of STIM2–$N_{\text{ext}}$ and $c\text{STIM–L}_{\text{full}}$ remained better dispersed in the $^1\text{H}–^{15}\text{N}$-HSQC spectra acquired under similar low $\text{Ca}^{2+}$ conditions (Fig. S4B and C). Although the spectra from both the STIM2 and $C.\text{elegans}$ proteins clearly showed an increase in the number of peaks collapsed into the central ~7.5–8.5 ppm $^1\text{H}$ region, the H(N) peak intensities were more homogeneous for $C.\text{elegans}$ (Fig. S4B and C). Thus, we pursued backbone atom chemical shift assignments of $c\text{STIM–L}_{\text{full}}$ as this construct was taken to be the most tractable for atomic resolution structural analyses in low $\text{Ca}^{2+}$.

We completed ~95 and ~75% of the backbone atom chemical shift assignments of $c\text{STIM–L}_{\text{full}}$ in high and low $\text{Ca}^{2+}$, respectively. The chemical shift index (CSI) shows that in solution and high $\text{Ca}^{2+}$, $c\text{STIM–L}_{\text{full}}$ adopts secondary structure elements very similar to the crystal state (Fig. S5A). Remarkably, in low $\text{Ca}^{2+}$, the protein showed two structural forms, as indicated by two distinct tracts of connectivity in the three-dimensional (3D) spectra acquired for the sequential backbone atom assignments and corresponding peak doubling in the
The CSI for each of these conformations show that the $\alpha_0$ and $\alpha_1$ of one state remains structured (form 1), whereas these helices become more unstructured in a second state (form 2) (Fig. S5B). Most unassigned residues in form 1 and form 2 cluster on the canonical EF-hand loop, $\alpha_2$, $\alpha_3$ and the noncanonical EF-hand loop. The difficulty in assigning the chemical shifts for residues in this EF-hand region in low Ca$^{2+}$ is due to poor chemical shift dispersion.

Fig. 3. Ca$^{2+}$ ion coordination geometry in the Caenorhabditis elegans EF-hand loops. (A) The 2Fo$-$Fc electron density map contoured at 2.0$\sigma$ (gray mesh) relative to the polypeptide chain model of the canonical EF-hand loop (sticks), Ca$^{2+}$ ion (yellow sphere), and water molecule (red sphere) is shown at left. At left, the Fc$-$Fc (9.0$\sigma$) omit map (magenta mesh) shows unaccounted-for density in the absence of the Ca$^{2+}$ ion. The pentagonal (green dashed lines) bipyramidal coordination geometry of the Ca$^{2+}$ ion by seven oxygen atoms is shown at right. The distances of each of the liganding oxygen atoms to the Ca$^{2+}$ ion (dashed black lines) are indicated below the atom name. The distance (red dashed line) to the water molecule (red sphere) is indicated below the oxygen atom name in red font. The X, Y, Z, #, x, and z residues corresponding to the loop positions shown in Fig. 1C are labeled adjacent to the side chain. (B) The Fo$-$Fc electron density map contoured at 2.0$\sigma$ (gray mesh) relative to the polypeptide chain model of the noncanonical EF-hand loop (sticks) is shown at left. At right, the Fc$-$Fc (2.0$\sigma$) omit map shows no unaccounted-for density (magenta mesh) in the space encompassed by the loop, indicating no ion coordination in the noncanonical EF-hand loop.
caused by protein unfolding and possibly conformational exchange of residues involved in dimerization/oligomerization. The SAM domain remains largely structured and α-helical for both forms detected in low Ca\(^{2+}\) (Fig. S5B).

Interestingly, the CSI of the C-terminal region of the forms 1 and 2 show a propensity for formation of a short β-strand between ~residues 177–185 (Fig. S5B). In the Ca\(^{2+}\)-bound state, this region of cSTIM–L\(_{\text{full}}\) folds into part of the long α10 helix. Thus, α10 undergoes a structural transition from α-helix to β-strand upon Ca\(^{2+}\) depletion (Fig. S5A and B). Indeed, some of the largest magnitude \(^1\)H(\(^{15}\)N) chemical shift perturbations (CSP) occur at this C-terminal region when comparing the conformations in high and low Ca\(^{2+}\) (Fig. 6B). Additional large CSPs are observed in the α0 and α1 helices, which become more unstructured in low Ca\(^{2+}\) (Fig. 6B).

To assess the structural integrity of the β-strand, which was detected in both low Ca\(^{2+}\) forms of cSTIM–L\(_{\text{full}}\), we used NMR to probe the solution dynamics of the protein. We focused on form 1 because the relative abundance was higher based on peak intensities in the HSQC (Fig. S6). In high Ca\(^{2+}\), the \(^{15}\)N–\(^1\)H-heteronuclear NOEs indicated a stably formed structure with low backbone flexibility on the ~ns timescale. Specifically, the data showed that most residues show saturated/reference peak intensity ratios of >0.8, excluding residues near the N- and C-termini, which show markedly decreased ratios between residues ~23–30 and ~184–192, respectively (Fig. S6). In low Ca\(^{2+}\), these terminal residues similarly show a high degree of flexibility on the ~ns timescale; however, the residues in between the unassigned canonical EF-hand loop, α2, α3, and the noncanonical EF-hand loop show enhanced mobility compared with the state observed in high Ca\(^{2+}\) (Fig. S6). Further, although the mobility of the residues involved in β-strand formation appears unaffected by the loss of Ca\(^{2+}\), the six residues immediately upstream (i.e., residues ~172–178) of the β-strand show an increased mobility on the ~ns timescale (Fig. S6). Thus, the β-strand formation may be facilitated and/or promoted by an unstructuring or increased mobility of residues immediately upstream of the far C-terminal motif.

The far C-terminal region of the C. elegans STIM luminal domain affects folding, stability, and oligomerization kinetics

Having discovered a propensity for the C-terminal residues of α10 to undergo an α-helix (in high Ca\(^{2+}\)) to β-strand (in low Ca\(^{2+}\)) transition, we then assessed the role of the transitioning residues in the folding and stability of cSTIM–L\(_{\text{full}}\) with and without Ca\(^{2+}\). Our crystal structure revealed that L180 is an important anchor residue that packs into the EF-hand cleft in high Ca\(^{2+}\); moreover, our solution NMR data showed the L180 undergoes an α-helix to β-strand transition in low Ca\(^{2+}\). Thus, within the hydrophobic V178–V179–L180–F181 stretch at the C-terminus, we created two separate double mutations to probe how residue variability in this region affects the structure and stability of the entire luminal domain: (i) V178A–V179A–L180–F181 (AALF mutant) and (ii)
Table 1. Thermal stability of STIM luminal domain constructs.

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<tr>
<th>Name</th>
<th>Species</th>
<th>Residue range</th>
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<th>−Ca²⁺</th>
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<td>cSTIM–Lfull(VVAA)</td>
<td>C. elegans</td>
<td>23–192</td>
<td>STIM</td>
<td>44.0</td>
<td>36.0</td>
<td>8.0</td>
<td>8.0</td>
</tr>
<tr>
<td>cSTIM–Lfull(AALF)</td>
<td>C. elegans</td>
<td>23–192</td>
<td>STIM</td>
<td>48.0</td>
<td>12.5</td>
<td>35.5</td>
<td>35.5</td>
</tr>
</tbody>
</table>

a. N denotes N-terminally extended relative to EF–SAM; Lfull denotes full luminal domain; L14 denotes N-terminal deletion relative to the full luminal domain.
b. Tm was taken as the temperature where the fractional change in CD signal was 0.5 in the presence of 5 mM CaCl₂.
c. Tm was taken as the temperature where the fractional change in CD signal was 0.5 in the absence of CaCl₂.
d. ΔTm, CaCl₂ was taken as Tm (+CaCl₂) – Tm (−CaCl₂).
e. ΔTm, mut was taken as Tm [cSTIM–Lfull(wild-type, −CaCl₂)] – Tm [cSTIM–Lfull(mutant, −CaCl₂)].
f. Taken from Refs. [26,27].
g. Taken from Refs. [28,29].

V178−V179−L180A−F181A (VVAA mutant). Far-UV CD spectra were acquired in high and low Ca²⁺ to assess secondary structure. In the presence of excess Ca²⁺ and at 4 °C, the AALF and VVAA proteins showed a high level of α-helicity, as indicated by the strong negative ellipticity observed at ~208 and 222 nm. In low Ca²⁺, both the AALF and VVAA double mutant proteins showed a marked loss in negative ellipticity at ~208 and 222 nm, indicative of decreased α-helicity (Fig. 7A and B). However, the loss in negative ellipticity for these mutants was greater than observed for the wild-type cSTIM–Lfull protein, and the spectra showed a shift to lower wavelength for the ~208 nm minimum. These spectral differences imply that the VVAA and AALF mutations result in a less well-folded state in low Ca²⁺ compared with wild-type cSTIM–Lfull (Fig. S3A).

The thermal stability was also assessed by monitoring changes in CD ellipticity at 225 nm. In the presence of excess Ca²⁺ (i.e., 5 mM CaCl₂), the Tm of the AALF and VVAA double mutant proteins was decreased by ~15.5 and 19.5 °C, respectively, compared with wild-type (Table 1), consistent with the role of this C-terminal z10 region in packing into the EF-hand hydrophobic cleft. In low Ca²⁺, the VVAA mutant showed a Tm drop of ~13.5 °C, and the AALF double mutant Tm decreased an astonishing ~37.0 °C (Fig. 7C and D).

Then, we performed size-exclusion chromatography with in-line multiangle light scattering (SEC–MALS) to assess the quaternary structure of cSTIM1−Lfull at stabilizing temperatures (i.e., 4 °C). In high Ca²⁺, wild-type, VVAA and AALF versions of cSTIM–Lfull all eluted similarly as monomers (Fig. 7E). In low Ca²⁺, all proteins showed a shift to earlier elution volumes, indicating a change in particle size and/or shape. The wild-type and VVAA protein remained primarily monomeric despite the apparent conformational change. In contrast, cSTIM–Lfull−AALF formed dimers and larger oligomers in low Ca²⁺, readily detectable even at low concentration and temperature, consistent with the extreme destabilization observed for this mutant (Fig. 7E). Finally, we assessed the self-association kinetics of wild-type, VVAA and AALF using static light scattering at a higher temperature (i.e., 37 °C). As expected given the rank order of thermal stabilities, the AALF protein showed the fastest, and the wild-type protein showed the slowest self-association kinetics (Fig. 7F).
Fig. 6. Solution NMR characterization of cSTIM–L_{full}. (A) $^1$H–$^{15}$N-HSQC spectrum of cSTIM–L_{full} in low Ca$^{2+}$. Representative $^1$H($^{15}$N) crosspeaks showing peak doubling based on connectivities observed in 3D backbone NMR experiments are circled. The dashed black ellipse indicates the region of the EF–SAM structure that remains unassigned.

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Collectively, these biophysical data indicate that the inherent amino acid composition of the far C-terminal region of cSTIM \textsubscript{L}\textsubscript{full} can radically modulate the structure, stability, and self-association of the domain, which has important implications for triggering STIM and SOCE activation.

**Discussion**

The solution NMR structures of human STIM1 and STIM2 EF–SAM have been determined in high Ca\textsuperscript{2+} [27,29]. There is a high structural conservation between human STIM1 and STIM2 EF–SAM, as
both show a hydrophobic cleft formed by the EF-hand domain and the packing of a five-helix bundle SAM domain into the cleft, which is primarily mediated by the \( \alpha 10 \) helix. The intramolecular EF-hand:SAM–\( \alpha 10 \) interaction occurs akin to the manner that calmodulin recognizes myriad target peptides with hydrophobic anchor residues penetrating into the nonpolar pocket [27,44,52]. The sequence identity between human STIM1 and STIM2 EF–SAM domains is \( \sim 58% \) [53]. In comparison, the sequence identities between the \( C. \) elegans EF–SAM core and the human paralogs are 40% and 38% for STIM1 and STIM2, respectively [53]. Here, our crystal structure clearly reveals a high structural conservation between \( C. \) elegans EF–SAM and the human STIM1 and STIM2 solution structures, consistent with the high sequence conservation of EF–SAM from mammals to roundworms and indicative of a common Ca\(^{2+}\) sensing mechanism among STIM phylogeny. In \( C. \) elegans, EF–SAM the L176 and L180 of \( \alpha 10 \) represent the anchor residues that pack into the EF-hand cleft. The EF-hand:SAM domain interaction in \( C. \) elegans is also buttressed by an ionic interaction between the \( \alpha 3 \) R85 residue and the D177 on \( \alpha 10 \). Human STIM2 EF–SAM, but not STIM1, shows a similar EF-hand:SAM domain ionic interaction between K108 of \( \alpha 3 \) and D200 of \( \alpha 10 \) [29], K108 and D200 of human STIM2 align with R85 and D177 of \( C. \) elegans STIM. Given the similar locations of ionic and hydrophobic interactions in human STIM2 and \( C. \) elegans STIM EF–SAM domains, it is not unexpected that these two proteins share very similar EF-hand:SAM domain orientations (Fig. 2).

Our work also reveals that the nonconserved luminal region, upstream of EF–SAM plays a direct role in stabilizing the EF–SAM fold. Our crystal structure of cSTIM–L\(_\text{full}\) showed that an ancillary helix (\( \alpha 0 \)) forms in this nonconserved region; further, \( \alpha 0 \) interacts with both the EF-hand and \( \alpha 5 \) helix linker, acting as a brace that holds the immediately adjacent \( \alpha 1 \), \( \alpha 4 \), and \( \alpha 5 \) in place (Fig. 4). Despite sequence divergence between human STIM1, STIM2, and \( C. \) elegans STIM in this region upstream of EF–SAM [53], our solution experiments showed that each of these luminal domains devoid of this nonconserved segment is less thermally stable in both Ca\(^{2+}\)-bound and Ca\(^{2+}\)-free states (Table 1). Thus, it is tempting to speculate that the interaction of the nonconserved N-terminal region with the EF–SAM core protein is a conserved structural mechanism capable of stabilizing all STIM molecules. Indeed, it was recently shown that this nonconserved region of human STIM1 contacts W121 and K122 of the \( \alpha 4 \) helix of EF–SAM [42]. Remarkably, W121 of human STIM1 aligns with W102 of \( C. \) elegans STIM, an important contact residue for \( \alpha 0 \) elucidated in our crystal structure. Further, S-nitrosylation of Cys residues within this nonconserved region of STIM1 was found to further thermodynamically stabilize the full luminal domain of the protein, concomitant with an inhibition of STIM1 activation and SOCE [42,49]. In contrast, S-glutathionylation of these Cys residues (i.e., CS6) weakened Ca\(^{2+}\) binding affinity of the human STIM1 luminal domain, resulting in constitutive SOCE [50]. Therefore, sequence variability in the region upstream of EF–SAM may have evolved commensurate with an ability to undergo luminal-specific posttranslational modifications that can modulate EF–SAM stability and STIM function depending on specific environmental cues.

Interestingly, our solution NMR data reveal that two different conformations are sampled by cSTIM–L\(_\text{full}\) in low Ca\(^{2+}\). The SAM domain remains well folded, whereas the \( \beta 1 \), \( \alpha 2 \), \( \alpha 3 \), \( \beta 2 \), and a portion of \( \alpha 4 \) of the EF-hand domain is likely unfolded in both states. In contrast, \( \alpha 0 \) and \( \alpha 1 \) are well folded in form 1 and become unfolded in form 2 (Fig. 6 and Fig. S5). The two conformations provide a possible mechanistic explanation for the enhanced stability observed for the S-nitrosylated human STIM1 luminal domain in low Ca\(^{2+}\). Because \( \alpha 0 \), \( \alpha 1 \), a portion of \( \alpha 4 \) and \( \alpha 5 \) remain folded in form 1, stabilizing \( \alpha 0 \) contacts with \( \alpha 4 \) and \( \alpha 5 \) may be maintained in low Ca\(^{2+}\). In the case of human STIM1, the S-nitrosylation in the nonconserved region may favor these contacts compared with the unmodified state. Nevertheless, high-resolution structural information is needed to tease out the precise stabilization mechanisms of the full human STIM1 luminal domains.

Remarkably, we also found that both forms of cSTIM–L\(_\text{full}\) detected in low Ca\(^{2+}\) show an \( \alpha \)-helix to \( \beta \)-strand transition near the end of the \( \alpha 10 \) helix. Given that \( \beta \)-strands are not stable in the absence of H-bonding, we speculated that \( \beta \)-strands from two STIM luminal domains may pair as the mechanism driving EF–SAM dimerization in low Ca\(^{2+}\). Thus, we created VVAA and AALF mutations in \( \alpha 10 \), which decrease the predicted \( \beta \)-strand propensity based on PSIPRED analysis [54]. However, we found that these mutations markedly destabilize the cSTIM–L\(_\text{full}\) proteins in high and low Ca\(^{2+}\), and rather than suppressing potential \( \beta \)-sheet-mediated dimerization, increase the self-association kinetics (Fig. 7). Thus, \( \beta \)-sheet formation in this \( \alpha 10 \) region could be more involved in stabilizing the higher order quaternary structure after dimerization/oligomerization occurs, and dimerization is likely primarily driven by the unfolding events revealed in the EF-hand domain by our NMR work (see above).

Recent work using a chimera of human EF–SAM fused to \( T. \) thermophilus GroP-like gene E (GrpE), which restricts EF–SAM movement within strictly forming GrpE dimers, proposed that human EF–SAM binds up to \(-5\)–\(-6\) Ca\(^{2+}\) ions, not only to the canonical EF-hand site but also to multiple unconventional surface sites [55]. Here, our work
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provides the first direct atomic resolution insights into Ca\(^{2+}\) binding, elucidating a single Ca\(^{2+}\) binding site within each C. elegans EF–SAM protomer, even in the context of dimers making up the asymmetric unit and 10 mM CaCl\(_2\) (Fig. 2 and Fig. S3). No additional Ca\(^{2+}\) binding sites were observed in this study. Further, we show that the Ca\(^{2+}\) ion is coordinated with pentagonal bipyramidal geometry in the EF-hand loop, as observed with classical EF-hand proteins such as calmodulin. In addition, no areas of unaccounted-for electron density were observed in our Fo–Fc omit maps other than the canonical Ca\(^{2+}\) binding site, when the single Ca\(^{2+}\) ion was removed. Nevertheless, we cannot exclude the possibility of more Ca\(^{2+}\) ions transiently interacting with the highly electrostatically negative surface of the luminal domain and that transient Ca\(^{2+}\) binding events may play a role promoting the canonical Ca\(^{2+}\) binding revealed by our crystal structure. Interestingly, the work by Gudlur et al. also showed that large-scale unfolding does not occur within either the EF-hand or SAM domains in low Ca\(^{2+}\) [55]. Consistently, our solution NMR data of C. elegans EF–SAM in low Ca\(^{2+}\) shows that the SAM domain remains largely folded. In contrast, we find that regions of the EF-hand domain unequivocally become conformationally unstable, adopting unfolded states. Ultimately, high-resolution characterization of the structural dynamics of the STIM luminal domain in low Ca\(^{2+}\) remains an important milestone for fully understanding STIM activation.

Interestingly, knockdown of the ER Ca\(^{2+}\) pump causes the unfolded protein response in C. elegans, yet a similar response is not induced by STIM knockdown in these roundworms, consistent with the protein unfolding events revealed by our work [56]. In C. elegans, STIM knockdown causes sterility due to the importance of SOCE in the function of sheath cells and spermatheca required for ovulation [56,57]. Transgenic worms expressing constitutively active STIM show severe arrhythmia in the posterior body wall muscle contraction required for defection [56]. When C. elegans STIM is expressed in HEK293 cells, the protein appears preclustered at ER–PM junctions; however, SOCE is not activated until ER Ca\(^{2+}\) stores are depleted [58]. A series of human STIM1–C. elegans STIM chimeras revealed that the cytosolic domain of C. elegans STIM mediates the preclustering of C. elegans STIM [58]. Thus, structural changes in EF–SAM caused by low Ca\(^{2+}\) is key to inducing the conformational rearrangement of the cytosolic domain necessary for STIM activation and subsequent Orai1 channel gating. Here, our work has shed light on these luminal structural changes that include stable single Ca\(^{2+}\) coordination in the canonical EF-hand loop at high Ca\(^{2+}\), and conformational instability and unfolding of several motifs within the EF-hand domain as well as an \(\alpha\)-helix to \(\beta\)-strand transition in the SAM domain at low Ca\(^{2+}\).

Methods

Protein constructs, expression, and purification

C. elegans STIM, human STIM1, and human STIM2 cDNAs were from Origene Technologies, Inc. All STIM luminal constructs were subcloned into pET28a vectors (Novagen, Inc.). The residue numbers of the STIM luminal domain constructs used in this study are described in Table 1. Recombinant expression and purification of all luminal domain STIM proteins containing an N-terminal hexahistidine (His\(_{6}\)) tag were carried out as described previously [26,28,32]. All experiments except for crystal- lization were performed in 20 mM Tris–HCl (pH7.5) containing 100 mM NaCl and 0.5 mM EDTA or 5 mM CaCl\(_2\) for low and high Ca\(^{2+}\) samples, respectively. In addition, STIM1–N\(_{\text{ext}}\) and STIM2–N\(_{\text{ext}}\) protein buffers were supplemented with 1 mM DTT. Protein concentrations were calculated using \(\lambda\)\(_{280}\) nm = 1.182, 1.319, 1.342, and 1.181 (mg mL\(^{-1}\)) for cSTIM–L\(_{\text{full}}\), cSTIM–L\(_{\text{N}}\), STIM1–N\(_{\text{ext}}\), and STIM2–N\(_{\text{ext}}\), respectively.

Crystallization

For crystallization of cSTIM–L\(_{\text{full}}\), the \(\lambda\)-selenomethionine-labeled protein was prepared in 20 mM Tris–HCl, 300 mM NaCl, and 10 mM CaCl\(_2\). Crystals were grown at 4 °C in sitting and hanging drop plates (Hampton, HR3-114 and HR3-172) by mixing equal volumes of protein (10 mg mL\(^{-1}\)) and crystallization buffer [18% (v/v) PEG 3,350, 8% (v/v) tacsimate, pH 5.6]. Crystals appeared in approximately a week. Crystals were cryoprotected by immersion in well solution supplemented with 25% (v/v) glycerol before flash freezing in liquid nitrogen.

Data collection and crystal structure determination

Diffraction data were collected at the Canadian Light Source (CLS) beamline CMCF-BM (08B1-1). Data were processed with HKL-2000 (HKL Research Inc.). The initial phase and a model of the cSTIM–L\(_{\text{full}}\) were calculated using MAD in AutoSol [59]. The structure was refined using PHENIX [60]. The model was completed with iterative cycles of model building in COOT [61] and refinement with PHENIX_refine [60] and REFMAC [62]. The cSTIM–L\(_{\text{full}}\) structure contains two molecules in the asymmetric unit. Crystallographic data and refinement statistics are shown in Table S1.

Far-UV circular dichroism spectroscopy

Far-UV CD spectra were recorded on a Jasco J-815 CD spectrometer (Jasco, Inc.). Data were collected using 0.1-cm path length ES quartz cuvettes. Wavelength scan rates were 20 nm min\(^{-1}\) with a response time of 8 s and bandwidth of 1 nm. Thermal melts were acquired in 1 °C
increments at a scan rate of 1 °C min⁻¹. All CD experiments were performed using 0.4 mg mL⁻¹ protein.

Size exclusion chromatography with in-line multi-angle light scattering

SEC was performed on a Superdex 200 increase 10/300 GL column (GE Healthcare). 100 µL of protein at 2.0 mg mL⁻¹ was injected onto the column for each experiment. Multiangle light scattering (MALS) measurements were performed in-line using the three-angle (45, 90, and 135°) miniDawn static light-scattering instrument with a 690-nm laser (Wyatt Technologies, Inc.). Molecular weight was calculated using ASTRA software (Wyatt Technologies, Inc.) based on Zimm plot analysis using a refractive index increment, dn/dc = 0.185 L g⁻¹.

Static light scattering

Self-association kinetics of cSTIM−Lo full−WT, -VVAA, and -AALF was measured by using Unit (Unchained Labs). After 10 min of temperature equilibration, a final concentration of 25 mM EDTA was added to the EF−SAM proteins (0.1 mM) solubilized in 5 mM CaCl₂ to induce self-association. SLS intensity was collected at 266 and 437 nm every ~50 s for each time course.

45Ca²⁺ binding assay

45Ca²⁺ radioisotope binding assays were performed as described previously [63]. Briefly, cSTIM−Lo full (0.1 mM protein), which was dissolved in 0.45 mL of 20 mM Tris, 100 mM NaCl, pH 7.5, was placed in the sample compartment of an Amicon Ultra-0.5 mL concentrator (10 kDa molecular mass cutoff), and a 1 µCi bolus of 45CaCl₂ solution was added to the protein solution. The sample was subjected to centrifugation (at 5000 rpm) in a table top centrifuge (Eppendorf, 5415D) at ambient temperature for 30 s. Successive additions of nonradioactive Ca²⁺ were made to the protein solution, and the centrifugation procedure was repeated. The radioactivity in the filtrate (free Ca²⁺) and the radioactivity in an equal volume of the protein sample (total Ca²⁺) were determined by liquid scintillation counting.

Solution NMR spectroscopy

Solution NMR data were acquired using a Bruker Avance 800 MHz spectrometers equipped with a cryogenic, triple-resonance probe (Bruker Biospin). Assignments of backbone atom chemical shifts were made using standard 1H−15N−HSQC [64], 1H−15N−13C−HNCACB [67] experiments. Spectra were processed using NMRPipe [68], and sequential assignments were made using NMRView [69]. Backbone relaxation data were acquired using an 15N−(1H) heteronuclear pulse sequence [70] with and without a 3 s 1H saturation before 15N excitation. Secondary structure probability was determined from the assigned chemical shifts based on the chemical shift index [71]. The NMR buffer was 20 mM Tris, 100 mM NaCl, pH 7.5, and 7% (v/v) deuterium oxide (D₂O).

Accession numbers

The Protein Data Bank (PDB) accession number for the C. elegans STIM luminal domain (i.e., residues 23−192; cSTIM−Lo full) X-ray crystal structure determined in the presence of Ca²⁺ is 6PW7. The Biological Magnetic Resonance Data Bank accession numbers for the backbone atom chemical shifts of the C. elegans luminal domain (i.e., residues 23−192; cSTIM−Lo full) in the absence (form 1), absence (form 2), and presence of Ca²⁺ is 27981, 27982 and 27983, respectively.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.jmb.2019.10.003.

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References

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