Pore properties of Orai1 calcium channel dimers and their activation by the STIM1 ER calcium sensor

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Running title: Function of Orail dimers

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Keywords: Calcium, calcium channel, ion channel, cell signaling, Orai1, STIM1, store-operated channel, channel gating, selectivity filter

ABSTRACT

Store-operated Ca²⁺ entry signals are mediated by plasma membrane Orai channels activated through intermembrane coupling with Ca²⁺-sensing STIM proteins in the endoplasmic reticulum. The nature of this elaborate Orai gating mechanism has remained enigmatic. Based on the Drosophila Orai structure, mammalian Orai1 channels are hexamers comprising three dimeric subunit pairs. We utilized concatenated Orai1 dimers to probe the function of key domains within the channel pore and gating regions. The Orai1-E106Q selectivity filter mutant, widely considered a dominant pore blocker, was surprisingly non-dominant within concatenated heterodimers with Orai1-WT. The Orai1-E106Q/WT heterodimer formed STIM1activated nonselective cation channels with significantly enlarged apparent pore-diameter. Other Glu-106 substitutions entirely blocked the function of heterodimers with Orai1-WT. The hydrophobic pore-lining mutation, V102C that constitutively opens channels, was suppressed by Orai1-WT in the heterodimer. In contrast, the naturally occurring R91W pore-lining mutation associated with human immunodeficiency, was a completely dominantnegative over Orai-WT in heterodimers. Heterodimers containing the inhibitory K85E mutation extending outward from the pore-helix, gave an interesting partial effect on both channel-activation and STIM1-binding, indicating an important allosteric link between the cytosolic Orai1 domains. The Orai1 C-terminal STIM1-binding domain mutation, L273D, powerfully blocked STIM1induced channel activation. The Orai1-L273D/WT heterodimer had drastically impaired STIM1induced channel-gating, but, unexpectedly, retained full STIM1-binding. This reveals the critical role of Leu-273 in transducing the STIM1 binding signal into the allosteric conformational change that initiates channel gating. Overall, our results provide important new insights into the role of key functional domains that mediate STIM1-induced gating of the Orai1 channel.

The ubiquitously expressed Orai channels are an evolutionarily conserved family of Ca^{2+} entry channels mediating "store-operated" Ca^{2+} signals. These signals play a crucial role in regulating key cellular responses including gene expression, growth, secretion, and motility (1-4). Orai channels function in the plasma membrane (PM) but are activated through a dynamic intermembrane coupling process between the ER and PM, mediated by STIM proteins. These ER membrane proteins are sensors of ER luminal Ca²⁺ changes and undergo an intricate unfolding process when Ca²⁺ stored in the ER becomes depleted (1-3,5-7). The unfolded STIM1 protein extends from the ER membrane surface and becomes trapped in ER-PM junctions by attaching to the PM where it tethers and activates PM Orail channels (1,2,5-7). A considerable number of immunological, muscular and inflammatory disease states are related to changes in the operation of STIM proteins and Orai channels (1,8-10). Hence, our understanding of the molecular coupling and function of the two proteins has major translational significance. Although we have recently determined much on the function and organization of STIM and Orai proteins (11-14), there is still considerable uncertainty about how the proteins associate within junctions and how these interactions mediate opening of Orai channels and the generation of Ca^{2+} signals (1-4,15,16).

The highly Ca²⁺-selective mammalian Orai1 channel bears close sequence homology with the Drosophila Orai (dOrai) channel from which the crystal structure was recently resolved (17). The dOrai channel protein has four transmembranehelices and is organized as a hexameric assembly of subunits (see Fig. 1A). Although earlier studies were interpreted to indicate a tetrameric structure (18-24), recent evidence indicates the Orai1 channel functions as a hexamer (25,26). Based on the dOrai crystal structure, most of the hexameric channel has 6-fold symmetry. The central pore is created from the six pore-forming N-terminal transmembrane helices (M1). Surrounding this are the tightly packed M2 and M3 helices, and at the periphery of the channel, the C-terminal M4 helices form STIM1-binding sites on the cytoplasmic face of the channel. The six M4 helices each have a small cytoplasmic extension (M4-ext; 39 amino acids in Orai1) that folds in two different configurations - one is almost straight from the M4-helix, the other is bent almost 180° back on itself. Each adjacent pair of M4-ext helices closely interact with one another in an antiparallel configuration. Thus, the STIM1-binding domain made up of the pair of M4-ext helices has three-fold symmetry (see Fig. 1A). Unclear is whether these two M4-ext helices dissociate to form separate STIM1-binding sites (17,27).

Because of the C-terminal interactions between adjacent pairs of M4-ext helices, the Orai channel comprises three dimers. Our studies therefore focused on the function of concatenated versions of these Orai1 dimers. Using such dimers, we were able to mutate either one or both of the separate Orai1 monomers, allowing us to gain important new information on the functional properties of the N-terminal pore-forming helix and the C-terminal STIM1-binding domain.

Results and Discussion

Orail dimers – reliable probes for Orail structure/function. In a recent study, we compared the function of concatenated Orail channels of varying length (25). Surprisingly we observed that concatemers containing 2 to 6 Orai1 subunits, all give rise to authentic CRAC currents, similar to the expressed Orai1 monomer. Although Orai1 exists and functions as a hexamer, we concluded that the activity of concatemers containing 3-5 Orail subunits, results from hexamers assembling from only the first two N-terminal subunits in the concatemer. This results in the formation of a hexameric "trimer of dimers", with the additional C-terminal Orail subunits in each concatemer (that is, subunits beyond the initial N-terminal dimer) extending out from the hexamer and not contributing to function. The exceptions to this anomalous behavior, were concatenated Orail dimers and hexamers. The hexamer appeared to largely form hexameric rings. The Orail dimers reliably assembled into hexameric Orai1 channels. We therefore constructed a series of Orai1 dimers (Fig. 1B) containing point mutations at functionally key residues, placed within either one or both subunits in the dimer (see Fig. 1C). For each construct, we assessed both the expression level (Fig. S1) and cellular location of expression (Fig. S2). In all cases, mutations did not alter either the level or the PM-localization of Orai1 dimers. We expressed each mutant in HEK cells in which endogenous Orail was eliminated through CRISPR/Cas9 gene editing, and STIM1-YFP was stably expressed (referred to as HEK-O1^{ko}S1⁺ cells) (25). The functional consequences of hetero- and homodimer

mutations provided important new mechanistic information on Orail channel operation.

Dominance of selectivity filter mutations in **Orail dimers.** The E106 residue is well established to function as the critical selectivity filter in the Orai1 channel (28-30). We initially examined the expression of Orail monomers containing the E106D and E106Q mutations, measuring their function either expressed alone or co-expressed with Orai1-WT (Fig. 2). The Orai1-E106D mutation has a shortened side chain which results in altered pore geometry and decrease channel selectivity (28-30). As expected, this mutation is still able to mediate Ca^{2+} entry (Fig. 2A) and coexpressed with Orai1-WT the mutant has little effect (Fig. 2B). In contrast, removing the charge on the E106 by replacing with Gln (but retaining side-chain length) gives a non-conducting pore (Fig. 2C). In this case, co-expression of Orai1-WT with Orai1-E106Q, clearly causes inhibition of Ca²⁺ entry (Fig. 2D). This result was expected since the E106Q mutant has been recognized earlier as a "pore-dead" mutant (19,29-32).

In such co-expression studies, the stoichiometry of subunits in hexamers is of course uncertain. Therefore, in further experiments, we compared the function of the E106D and E106Q subunits when incorporated into Orai1 dimer constructs. Initially we combined Orai1-WT and Orai1-E106D subunits (Fig. 3A-C). The Orai1-E106D homodimer (DD) gave rise to Ca^{2+} entry that was little different to the Orai1-WT homodimer, consistent with the data in Fig. 2 for monomer expression. The heterodimers of Orail-WT and Orai1-E106D (OD, DO) also behaved similarly for Ca²⁺ entry. Using the Orai1-E106O homodimer (OO), there was no detectable Ca^{2+} entry (Fig. 3D), again consistent with the monomer in Fig. 2C. However, the heterodimers of Orail-WT and Orai1-E106Q (OQ, QO) unexpectedly revealed strong Ca²⁺ entry, similar to that mediated by Orai1-WT (Fig. 3E,F). This result is in stark contrast to heterodimers comprising Orai1-WT together with the Orai1-E106A mutant. For this mutant, the homodimer (AA) is clearly pore-dead (Fig. 3G), just like the Orai1-E106Q mutant (QQ). However, the heterodimers of Orai1-WT and Orai1-E106A (OA, AO) were also almost without function (Fig. 3H,I).

pore-dead mutation and one that is a dominantnegative over Orai1-WT subunits (1,19,29-32). Indeed, the dominance of E106Q has been described in Orai1 competition studies as providing evidence to support a purported tetrameric assembly of Orail channels (19). The dominance of E106Q was also the basis for its conditional transgenic expression in animal studies to assess the role of Orai1 in muscle tissue (33). Based on our studies here, although E106A and E106Q are clearly both pore-inactive mutations, their relative dominance in blocking Orai1-WT channels is very different. Clearly, therefore, such dominance studies would have been more conclusive using E106A instead of E106Q. The descriptions in earlier papers of the dominance of E106Q over Orai-WT (1,19,29-32) was in each case based on coexpression studies in which the relative expression of the Orai1 subunits could not be controlled. Only in concatemer studies can the ratio of expression be defined. Indeed, we used similar concatemeric constructs to reveal that E106L, E106V, E106G, and E106K all have complete dominance over Orai1-WT (Fig. S3).

This is important information since the action of E106Q has been widely described as both a

Probing the anomalous functional characteristics of E106Q. The unexpected finding that OQ and OO heterodimers are functional prompted us to undertake an electrophysiological characterization to assess their selectivity and conductance. Important was to compare their function with constructs containing E106D since this mutation is known to convert the highly Ca²⁺-selective Orail channel into a non-selective cation channel (28-30). As expected, the Orai1-WT homodimer (OO) gave typical CRAC current with full inward rectification and a reversal potential slightly above 50 mV (Fig. 4A,B). In contrast, the Orai1-E106D homodimer (DD) gave rise to a non-selective current with a strong outward component and much reduced reversal potential (approximately 5 mV; Fig. 4B). The reduction of a single methylene group by replacing the wildtype Glu-106 residue with Asp, is reported to cause widening of the pore sufficient to make it cation nonselective (34). Surprisingly, the presence of just a single aspartate in the E106D heterodimers (OD, DO) gave an I/V curve very similar to DD, with reversal potentials

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of approximately 10 mV (Fig 4B). We would have expected that the properties of the heterodimeric channels (OD and DO) would be somewhere between the two homodimers. Thus our results indicate that replacement of just three Glu residues with Asp in the selectivity filter of the hexameric channel, is still sufficient to render it a nonselective channel.

The effects of the E106Q mutation are also surprising (Fig. 4C,D). Clearly, the homodimeric replacement in QQ results in a channel with no conductance, as expected. However, the large but non-selective current seen with both the OQ and QO heterodimers, was unexpected. In both cases the reversal potential was close to zero mV. Assuming the channel operates as a hexamer, this reveals that the full configuration of 6 charges in the selectivity filter is not required for cation conductance. Thus, the presence of just 3 negative charges in the pore filter is sufficient for strong cation permeation, albeit non-selectively.

We also examined the function of a concatenated hexameric Orail construct containing three pairs of OQ or QO dimers (Fig. 4E-G). The function of both of these two hexamers was very comparable to that of their respective contributing heterodimers. For Ca^{2+} entry, both the OQOQOQ and QOQOQO hexamers gave similar function as the wildtype homohexamer (OOOOOO) (Fig. 4E). Current measurements with the hexamers revealed that both heterohexamers behaved as nonselective cation channels, similar to heterodimers. Reversal potential for hexamers were slightly elevated (Fig. 4G), and currents mediated by the hexamers were slightly smaller perhaps due to less efficient expression compared to dimers. The similar function of both hexamers and dimers, supports the view that the Orail channel is hexameric and comprises three pairs of dimers. In our previous paper, we concluded that a hexamer-concatemer could create a functional channel either as a single hexameric ring or as a "trimer of dimers" in which three concatemeric hexamers each contribute one dimer to form a single functional hexamer (25). The results we show here for the two heterohexamers would be consistent with either scheme.

Pore properties of the non-selective OQ and QO concatemer dimers. The surprising effect of

in the Orail selectivity filter prompted us to investigate whether this altered the channel pore diameter. For this we measured the permeation of methylated ammonium derivatives of increasing size. We compared permeation of the OQ or QO Orail heterodimers with that of the OO wildtype homodimer expressed in HEK-O1^{ko}S1⁺ cells (Fig. 5). Consistent with previous studies expressing wildtype monomeric Orai1 (34,35), we observed that the OO dimer had almost no current after switching from 20 mM Ca²⁺ to 150 mM methylammonium (Fig. 5A,B). Increasing the methyl groups on methylammonium to two, three and four, also revealed no measurable currents (Fig. 5C,D). These results are similar to earlier results from which the Orail pore size was calculated to be \sim 3.9Å (34,35). In contrast, using cells expressing the OQ heterodimer, switching from Ca²⁺ to methylammonium resulted in a massive increase in current (Fig. 5E,F). Moreover, large currents were still observed following consecutive switching from Ca²⁺ to dimethylammonium or even trimethylammonium. The same current measurements using the OO construct provided very similar results (Fig. 5G,H). In both cases, the decreased currents and increasingly negative reversal potentials seen with increasing number of methyl groups on ammonium ion, allowed us to estimate an apparent pore diameter for the OQ and QO constructs considerably greater the WT pore (see Experimental Procedures and Fig. S4; this estimate is based on the assumption that permeability is limited only by the steric properties of the pore). We also constructed equivalent Orai1 heterodimers containing E106 residues mutated to asparagine instead of glutamine (ON, NO and NN). Ca²⁺ measurements revealed a similar functional pattern; NN gave no Ca2+ entry whereas ON and NO gave Ca^{2+} entry identical to the wildtype, OO (Fig. S5A,B). Current measurements revealed the ON and NO dimers had similar ion non-selectivity (Fig. S5C-E), and currents measured with methylated ammonium ions (Fig. S5F-M) indicated similarly enlarged estimated pore size.

replacing three glutamate residues with glutamines

These surprising results clearly reveal that reducing the number of negatively charged residues in the selectivity filter of Orail from 6 (in the wildtype hexamer formed from OO) to 3 (formed by the OQ, QO, ON, or NO dimers), causes a

substantial increase in apparent pore diameter and a major alteration in ion channel selectivity. This change in pore properties was even larger than that resulting from substitution of all six E106 residues within the hexamer with Asp residues, indicating that both charge density and steric influence of head-group size contribute to the geometry of the pore. The results with the OQ and ON heterodimers also gives support to a recent hypothesis on the ionselectivity mechanism for the Orai channel. Thus, in an interesting new crystallography study on dOrai channel structure, Hou et al (36) suggest that glutamates comprising the selectivity filter, have sidechain flexibility that allows two Ca²⁺ ions to be positioned one above the other. Repulsion between the two Ca^{2+} ions allows one Ca^{2+} to be displaced and move through the pore and be replaced by a new Ca^{2+} ion. Thus in the presence of external Ca^{2+} , the channel is highly Ca^{2+} selective. In the case that three glutamates are replaced with three similarly sized but uncharged glutamines, we would suggest that the selectivity filter loses the ability to bind two Ca²⁺ ions, Ca²⁺ selectivity would be lost, and monovalent cations would permeate in the presence of external Ca²⁺.

Orail dimers mutated elsewhere in the Orail M1 pore-forming helix: In further experiments, we assessed the role of substituting other pore-lining Orai1 residues within the heterodimers. Immediately below the E106 selectivity filter, the hexameric pore is lined with three rings of hydrophobic residues, thought to act as a hydrophobic gate (37). Mutating the first of these residues (V102; see Fig. 1C) to Cys causes the channel to be constitutively active although with altered cation selectivity (37). Using an Orai1 V102C homodimer (CC), we observed substantial constitutive Ca2+ entry in HEK-O1ko cells (not over-expressing STIM1) (Fig. 6A, Fig. S6A). Patch-clamp measurements also revealed a substantial current (Fig. 6B, Fig. S6B) with a reversal potential of approximately +25 mV (Fig. 6C, Fig. S6C). These results all agree closely with studies expressing monomer Orai1-V102C (13, 37). In contrast, the heterodimers containing one Orai1-WT and one Orai1-V102C subunit (OC or CO) displayed no constitutive Ca²⁺ entry or current (Fig. 6A-C). Using HEK-O1^{ko}S1⁺ we measured currents after depletion of stores following breakin with 20 mM BAPTA in the pipette. Cells

expressing the CC homodimer revealed a typical CRAC-like store-dependent current with reversal potential greater than +50 mV (Fig. 6D, Fig. S6D), in close agreement with the earlier results with the Orai1-V102C monomer (13,37). In similar experiments, the two Orai1 heterodimers, OC and CO, gave current properties almost identical to those seen with CC. The results are interesting in revealing that the wildtype Val at position 102 is dominant over the Cys mutant in the heterodimer. We might have expected that the heterodimer would have some constitutive function, but instead the hydrophobic gating function of this residue appears to be maintained with just three hydrophobic Val residues lining the pore. Conversely, the constitutive activation of Ca²⁺ entry by V102C appears to require mutation of all six residues.

This result is an interesting contrast to experiments to assess the function of a different pore-lining residue, R91, residing below the hydrophobic pore region (Fig. 1C). The R91 residue lies within the highly positively charged region of the pore toward the cytosolic side of the channel. The naturally occurring R91W mutation has been reported to cause severe combined imunodeficiency in homozygous patients (28). We constructed Orail mutant heterodimers with one R91W mutation (OW and WO). Expressed in HEK-O1^{ko}S1⁺ cells, we observed that the Orai1 heterodimers, OW and WO, each gave no storedependent Ca²⁺ entry (Fig. 6E,F). Indeed, their action was virtually the same as non-transfected cells. This result is quite different to that earlier reported by (38) in which co-expression of Orai1-WT and Orai1-R91W resulted in only a modest slowing of the activation of I_{CRAC}. In that study, it is likely that expression of WT and mutant R91W were not equal, in contrast to the OW and WO heterodimers. Another earlier study examined tetramer-concatemers of Orai1 in which 1, 2, 3, or all 4 WT Orai1 subunits were substituted with Orai1-R91W subunits (39). In that study, a single R91W residue caused 50% loss of activity, and with 2 or 3 mutant subunits, there was almost 100% loss of activity. Our results would concur with this dominance observed with R91W. However, the use of the tetrameric construct is difficult to reconcile since the Orai1 channel functions as a hexamer (17,25,26). Likely the expressed tetramers form hexamers through trimers of dimers, as described above. Given the almost complete dominance of the R91W in our heterodimers, patients heterozygous for the R91W gene would be expected to have substantially reduced Orai1-mediated Ca^{2+} entry. Such heterozygous R91W carriers had T cells with approximately 50% of WT Ca^{2+} entry (28). It is possible that compensatory expression of other Orai subtypes may have occurred in such patients.

We next used Orai1 dimers to examine the effects of mutating one of the residues of the poreforming M1 helix which is not lining the pore (see Fig. 1C). The K85 residue has been the center of much attention since the K85E mutation completely blocks store-dependent channel activation (13,40-42). This residue faces outward from the pore-forming helix with its head group likely extending into the cytosol close to the cytosolic membrane surface. Using Orai1 homodimers with both K85 residues mutated to E, we determined that there was no Ca²⁺ entry function, consistent with the blocking action determined earlier (40) (Fig. 6G,I). Interestingly, we observed that the heterodimers with a single K85E mutation (OE and EO), each displayed Ca^{2+} entry that was approximately 50% of the OO-WT homodimer (Fig. 6H). Current analysis confirmed that OE and EO had similarly diminished channel function and that the I/V relationship revealed each mediated otherwise normal CRAC current (Fig. 6J-K, Fig. S6E). Since the K85 site has been implicated in binding STIM1 to possibly gate the channel (40-42), we also examined FRET between CFP-tagged Orai1 dimers (C-terminally labeled) and STIM1-YFP. In these studies, we observed FRET with each heterodimer (OE or EO) was decreased approximately 30% compared to the WT homodimer (Fig. 6L). The EE mutant homodimer gave a slight further decrease in FRET.

The K85E mutation has been the subject of considerable speculation. It clearly has a profound effect on the store-operated activation of Orail channels. It was speculated that K85 mediates gating through possible interactions of STIM1 with the cytosolically-protruding extension of the pore-forming M1 helix (M1-ext) (40-42). However, recent studies suggest that K85 is necessary for maintaining the configuration of the channel rather than directly mediating channel gating (13).

Nevertheless, our current results do show that altering the cytosolic M1 extension (K85E) does affect binding of STIM1. Although the OE and EO heterodimers have reduced current, the poreproperties have not changed. The effects on STIM1-binding of mutating the K85 sites in the dimer, supports a model of close coupling between the M1-ext and the M4-ext, as suggested earlier (41). These two domains exist in a closely associated complex together with the 16 amino acid 2-3 loop joining the M2 and M3 helices (see Fig. 1C). Recent evidence indicates that the 2-3 loop mediates important communication with the M1-ext helix (43). Although the C-terminal M4-ext is the primary site for STIM1 binding to gate the channel, we would speculate that the entire cytosolic domain (M1-ext, 2-3 loop, and M4-ext) exists as a conformationally coupled entity, alteration of which affects both STIM1 binding and gating of the channel. However, even though K85E alters STIM1 binding, we do not consider that the N-terminus is a STIM1 binding site required for gating. Thus, we revealed earlier that a C-terminal mutation in Orai1 that mimics STIM1-induced channel activation without any STIM1 present, is similarly sensitive to the K85E mutation (13).

Probing the STIM1-Orail coupling interface at the C-terminus of Orail: We shifted our exploration of the function of Orail dimers to understanding more about the C-terminal M4-ext helix, the site at which STIM1 binds strongly and initiates activation of the channel. Unlike the Nterminus which has 6 identical helices, the dOrai crystal structure shows that the short 45-amino acid C-terminal M4-ext exists in two different configurations (Fig. 1A,C). Thus, each dimer of Orai contains one M4-ext that is in an extended conformation, and one in a bent conformation (17). Based on the dOrail structure, the two M4-ext helices in Orai1 are likely attached in an antiparallel configuration through inter-helical hydrophobic interactions between the L273 and L276 residues in each helix (see Fig. 7A). Mutation of L273 in Orai1 to either D or S is known to abolish binding of STIM1 and any store-dependent activetion of the channel (27,38,41,44-46). Likely this stems from disruption of the two bound M4-ext helices, although how STIM associates with these helices is still unclear (47,48). We constructed tdTomato-tagged Orai1 dimers in which either one

or both monomers contained the L273D mutation. We tested the function of these dimers in HEK- $O1^{ko}S1^+$ cells. As expected, the L273D mutant Orai1 homodimer (LL) gave no store-operated Ca²⁺ entry (Fig. 7B,C), nor any current through the channel (Fig. 7D,E). In contrast, the two Orai1 heterodimers (OL and LO) both gave very small Ca²⁺ entry (Fig. 7B,C). Current measurements also revealed that the conductance of the OL and LO heterodimers was extremely low (Fig. 7D,E). Summary data for both the Ca²⁺ and current measurements reveal that the OL and LO heterodimers have less than 10% of the activity of the wildtype OO homodimer (Fig. 7C,E).

We considered that the drastically decreased function of the OL and LO heterodimer channels might reflect a loss of the ability to bind STIM1. To address this question we undertook FRET studies using another set of Orai1-dimers (OO, OL, LO and LL) with C-terminal CFP-tags. These were transiently expressed in cells stably expressing STIM1-YFP. Surprisingly, we observed that the Orail OL and LO heterodimers did not have a defect in STIM1-binding activity. Thus, after storedepletion, the FRET values for OO, OL, and LO with STIM1-YFP were not significantly different (Fig. 7F). As expected, the FRET value for the LL homodimer was much decreased indicating that STIM1 binding was effectively abolished. In further experiments, we undertook direct visualization of cells co-expressing Orai-tdT dimers and STIM1-YFP (Fig. 7G). In these studies we utilized HEK-O1^{ko}S1⁺ cells and transiently transfected each of the Orai1-tdT-tagged dimers. The results are highly consistent with the FRET studies. In cells transfected with the wildtype Orai1-OO homodimer (Fig. 7G, top row, upper-left cell), there was almost complete co-localization of OOtdT with STIM1-YFP. The two molecules colocalized within the same clearly observable punctal regions in cells which represent the ER-PM junctional areas. An almost identical pattern of colocalization was also seen in cells expressing the OL and LO heterodimers (Fig. 7G, 2nd and 3rd rows). These results contrast dramatically with cells expressing the LL mutant homodimer (Fig. 7G, bottom row). In this case, there is clearly no co-localization of the LL dimer with the STIM1 protein at the PM junctions, even though STIM1 can still form puncta by interacting with PM

phospholipids in the ER-PM junctions (49,50). Indeed, as seen in Fig. 7G (top row) the lower right cell did not express Orai1, yet the STIM1 still entered puncta exactly the same as in the cell coexpressing Orai1.

These results provide important new information about the critical nature of the Orai1 Cterminal configuration in mediating gating of the Orai1 channel. The single L273D mutation in the OL or LO Orai1 heterodimers, drastically impedes the ability of Orai1 to be gated by STIM1, yet, the same heterodimers have not lost any significant ability to bind STIM1. Recent studies indicate that binding of STIM1 to the Orai1 C-terminal M4-ext is sufficient to gate the channel (13). This interaction causes an allosteric change in the Orai1 channel that is propagated from the M4-ext through the M3, M2 and M1 helices resulting in certain pore-lining M1 residues undergoing a structural alteration that constitutes gating of the channel (13,51). From the LL-Orai1 homodimer result, the two M4-ext helices have likely undergone unfolding sufficient to completely prevent STIM1 binding. With the single L273D mutation in the OL or LO heterodimers, the rearrangement must be much more subtle such that STIM1 binding is largely intact. Yet, the propagation of allosteric change across the protein to gate the channel, is almost completely lost. Thus, it seems that the M4ext may undergo only a small conformational change to initiate gating. This conclusion agrees with studies on cross-linking of the M4-ext helices by Tirado-Lee et al (27). Recent structural analysis has suggested that the interacting pairs of M4-ext helices act as latches on the outside of the channel that must be opened to allow the M1-ext helices to expand outward permitting the cytosolic mouth of the pore to open (36). This model was based on a crystalized dOrai structure that stabilized the unlatched channel with straightened M4-ext helices. However, this dOrai configuration was not based on any input from STIM proteins. Our results indicate that the initial STIM1-induced alteration of the M4-ext helices is extremely subtle and unlikely to be a large unlatching of the entire Orai outer structure.

Concluding remarks: The use of concatenated Orail dimers has proven highly useful in determining the role of key residues in the channel pore and STIM1 binding site of Orai1. Using a CRISPR/ Cas9-derived Orai1-free background cell system, the dimer concatemers are reliable probes for the function of the hexameric Orai1 channel. Two highlights of these findings are particularly prominent. First, our studies provide some important new understanding on the properties of the pore selectivity filter that normally functions with six charged Glu residues. We speculate that these residues may have sufficient flexibility to allow formation of two Ca²⁺ binding sites within the selectivity filter, as recently suggested (36). The symmetrical and somewhat conservative replacement of three Glu residues with three intervening Gln residues (formed with OO or OO dimers), may prevent formation of the putative second site and render the pore as a nonselective cation channel. Functionally this construct operates the same as six Asp residues in the selectivity filter, however, in that case the shortened head groups lack the dimension and flexibility to form two sites. Future structure and modeling studies might examine this possibility. Second, our studies also throw new light on the Orail C-terminal binding site for STIM1. The OL and LO Orai1 heterodimers appear to have little change in their STIM1-binding properties, yet they have almost entirely lost the ability to mediate gating of the channel. This result emphasizes the exquisitely precise nature of the initial interacting site for STIM1 on Orai1 (the M4ext) through which it induces channel activation. The two M4-ext sites likely propagate this allosteric gating of the channel across the tightly packed M3, M2 and M1 helices as recently suggested (13).

Experimental Procedures

DNA constructs – Concatenated Orai1 dimers were constructed as described earlier. The 36 aa linker sequence is TSSSTMLHPGLSPGLLPLH PASIGGSGGSGGSGGSGRAT (25). For Orai1 heterodimers, point mutations in the M1 or M4-ext helices (E106, V102, R91, K85, L273), were performed on Orai1-WT monomers using the QuickChange Lightning Site-Directed Mutagenesis Kit (Agilent; 210518), then inserted into the ptdTomato-N1-Orai1 plasmid with XhoI and BamHI restriction sites. For C terminal-CFP tagged constructs, the tdT-tag was changed to ECFP with BamHI and NotI restriction sites.

Cell culture and transfection – HEK-O1^{ko} cells and the HEK-O1^{ko}S1⁺ cell line were generated using the CRISPR-Cas9 nickase system as described earlier (25). Cells were cultured in DMEM (Corning Cellgro) supplemented with 10% fetal bovine serum, penicillin and streptomycin (Gemini Bioproducts, CA) containing puromycin (2 μ g/ml) at 37°C with 5% CO₂. All transfections were performed by electroporation at 180 V, 25 ms in 4 mm cuvettes (Molecular Bio-Products) using the Bio-Rad Gene Pulser Xcell system in OPTI-MEM medium as previously described (52). Experiments were all performed 18-24 h after transfection.

Cytosolic Ca²⁺ measurements – Cytosolic Ca²⁺ levels were measured as described earlier (53) by ratiometric imaging using fura-2 (Molecular Probes). Cells were used 18-24 hours after transfection, and incubated with 2 µM fura-2 in buffer containing: 107 mM NaCl, 7.2 mM KCl, 1.2 mM MgCl₂, 1 mM CaCl₂, 11.5 mM Glucose, 0.1% BSA, 20 mM HEPES pH 7.2 for 30 min at room temperature, followed by treatment with fura-2 free solution of another 30 mins. Fluorescence ratio imaging was measured utilizing the Leica DMI 6000B fluorescence microscope and Hamamatsu camera ORCA-Flash 4 controlled by Slidebook 6.0 software (Intelligent Imaging Innovations; Denver, CO) as described previously (54). Consecutive excitation at 340nm (F340) and 380nm (F380) was undertaken every 2 sec and emission fluorescence was collected at 505 nm. Intracellular Ca^{2+} levels are shown as F340/F380 ratios obtained from groups of 8-25 single cells on coverslips. For the Orail concatemer experiments, data is shown for cells expressing a narrow range of tdTomato fluorescence in order to maximize consistency of responses between cells. For each concatemer experiment, we simultaneously examined each set of mutants with the corresponding wildtype concatemer. The expression levels for wildtype and mutant concatemers were within a similar narrow range of fluorescence intensity. All Ca²⁺ imaging experiments were performed at room temperature and representative traces of at least three independent repeats are shown. Scatter plots with means \pm SEM are shown for all the cells recorded.

Deconvolved fluorescence image analysis -For imaging of cells, we used the inverted Leica DMI 6000B automated fluorescence microscope and Hamamatsu ORCA-Flash 4 camera controlled by Slidebook software to collect and analyze high resolution fluorescence images. For the PM localization studies of tdTomato-tagged Orai1 concatemers, stacks of 10-20 3-D Z-axis image-planes close to the cell-glass interface were collected at 0.35 um steps. The no-neighbor deconvolution function of the Slidebook 6.0 software was used to analyze images and derive enhanced deconvolved images with minimized fluorescence contamination from out-of-focus planes. The tdT-tagged Orail concatemer images shown were typical of at least three independent analyses.

Confocal Imaging – Cells transfected with tdT-tagged constructs were imaged 14-20 hr after transfection. Live cell images were collected on the inverted Leica TCS SP8 confocal microscope using a 63x/1.40 Oil objective. Images shown are typical of at least three independent experiments.

Electrophysiological measurements and pore size estimation – Whole-cell patch-clamp recording was performed on HEK-O1^{ko}S1⁺ cells transiently transfected with tdT-tagged Orai1 concatemer constructs, or not transfected, as described (11). To passively deplete ER Ca²⁺ stores, the pipette solution contained: 135 mM Csaspartate, 10 mM HEPES, 8 mM MgCl₂ and 20 mM BAPTA (pH 7.2 with CsOH). For measuring constitutively active currents, HEK-O1^{ko} cells were transiently transfected with different Orai1-V102C concatemers, and the pipette solution included: 135 mM Cs-aspartate, 10 mM HEPES, 5 mM MgCl₂, 10 mM EGTA and 3 mM CaCl₂. The bath solution contained: 130 mM NaCl, 4.5 mM KCl, 5 mM HEPES, 10 mM Dextrose, 10 mM TEA-Cl and 20 mM CaCl₂ (pH 7.2 with NaOH). Divalent free solution included: 150 mM NaCl, 5 mM HEPES, 10 mM TEA-Cl, 10 mM HEDTA and 1 mM EDTA (pH 7.4). When indicated, 150 mM NaCl was substituted by 150 mM methylamine HCl, dimethylamine HCl, trimethylamine HCl, or tetramethylamine HCl. ICRAC was recorded until steady state, then the external solution was switched to different methylammonium-DVF solutions indicated. Currents were recorded in the standard whole-cell configuration using the EPC-

10 amplifier (HEKA). Glass electrodes with a typical resistance of 2-4 M Ω were pulled using a P-97 pipette puller (Sutter Instruments). A 50-ms step to -100 mV from a holding potential of 0 mV, followed by a 50-ms ramp from -100 to 100 mV, was delivered every 2 seconds. Currents were filtered at 3.0 kHz and sampled at 20 kHz. A +10 mV junction potential compensation was applied to correct the liquid junction potential between the bath and pipette solutions. The current measure at -100 mV was used in I-V curves. All data were acquired with Patch Master and analyzed using FitMaster and Graph Pad Prism 6.

Using the Goldman-Hodgkin-Katz (GHK) voltage equation, relative permeabilities of the different methylated ammonium ions and Na^+ (P_x/P_{Na}) were calculated from changes in the reversal potential:

$$P^{x}/P_{Na} = \frac{[Na]_{0}}{[X]_{0}}e^{\frac{\Delta E_{rev}F}{RT}}$$

in which $[X]_o$ and $[Na]_o$ are the external ionic concentrations, ΔE_{rev} is the change in reversal potential when switched to the testing cation, R is the universal gas constant, T is the temperature in Kelvin, F is Faraday's constant.

We estimated pore dimensions using the relative ion permeability ratios, as described earlier (35), assuming that ion permeability is limited mainly by steric hindrance. The relative permeability was calculate from:

$$\frac{Px}{P_{Na}} = k \left(1 - \frac{d_{ion}}{d_{pore}} \right)^2$$

in which k is a proportionality constant, d_{ion} is the ion diameter, and d_{pore} is the pore diameter (55,56). An estimate for d_{pore} was derived by relating decrease in permeability ratio to the increase in organic cation size.

Western blot analyses – Cells were washed with ice-cold PBS and lysed on ice using lysis buffer (RIPA, Sigma and 1x Protease inhibitor cocktail) for 30 min, followed by centrifugation at 14,000 g for 10 min at 4°C. Supernatants were collected and protein quantified using Bio-Rad DC kits. Proteins were resolved on 4-15% NuPAGE Bis-Tris precast gels and transferred to Bio-Rad Immuno-Blot PVDF membranes. After blocking in 5% non-fat milk for 1 hr at room temperature the membrane was incubated with primary antibody at 4°C overnight. Membranes were washed 3 times in PBST and incubated with secondary antibody for 1 hr at room temperature. Subsequently, membranes were washed 3 times with PBST. Peroxidase activity was examined with Pierce ECL Plus Western Blotting Substrate (Thermo Scientific) and fluorescence was collected using the FluorChem M imager (ProteinSimple).

Förster resonance energy transfer measurements - To determine FRET signals between stably expressed STIM1-YFP and transiently expressed CFP-tagged Orail dimers, we used the Leica DMI 6000B inverted automated fluorescence microscope equipped with CFP (438Ex/483Em), YFP (500Ex/542Em) and FRET (438Ex/542Em) filter cubes. At each time point, three sets of images (CFP, YFP and FRET) were collected at room temperature using a x40 oil objective (numerical aperture 1.35; Leica) and processed using Slidebook 6.0 software (Intelligent Imaging Innovations). Images were captured at 20-s intervals. Exposure times for the CFP, YFP and FRET channels were 1,000, 250 and 1000 ms, respectively. The decreased YFP channel exposure time compensates for the greater fluorescence intensity of YFP compared with CFP. Three-channel-corrected FRET was calculated using the formula:

$$F_{\rm C} = I_{\rm DA} - Fd/Dd \times I_{\rm DD} - Fa/Da \times I_{\rm AA}$$

where I_{DD} , I_{AA} and I_{DA} represent the background subtracted CFP, YFP and FRET images, respecttively, FC represents the corrected energy transfer, Fd/D_d represents measured bleed-through of CFP through the FRET filter (0.457) and Fa/Da is the bleed-through of YFP through the FRET filter (0.19). We used the E-FRET method to analyze 3cube FRET images as describe by (57), using the formula:

$$E_{app} = F_c/(F_c + G \times I_{DD})$$

in which G is the instrument specific constant (44,57). The EYFP-ECFP construct made as described above, was used to determine the Gparameter for E-FRET calculations. The value of G was determined by measuring the CFP fluorescence increase after YFP acceptor photobleaching using HEK-WT cells transiently transfected with the pEYFP-ECFP construct. The value of G was calculated as 1.9 ± 0.1 (n=32 cells). For studies determining E-FRET between YFP and CFP constructs, cells with a narrow range of YFP/CFP ratios were selected to ensure comparability between measurements. In all our E-FRET summary data, a region close to the PM was selected, and E-FRET analysis was conducted on cells with similar YFP/CFP ratios.

Acknowledgements: This work was supported by NIH Grants GM120783 and GM109279 to DLG, Predoctoral Fellowship GM125376 to RMN, and a Penn State University Junior Faculty Development Program grant to YZ.

Conflict of interest: The authors declare that they have no conflict of interest with the contents of this article.

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FOOTNOTES

This work was supported in whole or part by National Institute of Health Grants R01 GM120783 and R01 GM109279, and GM125376

The abbreviations used are: STIM, stromal-interacting molecule; PM, plasma membrane; ER, endoplasmic reticulum; CRAC, Ca²⁺ release-activating Ca²⁺; SOCE, store-operated Ca²⁺ entry; tdT, tdTomato

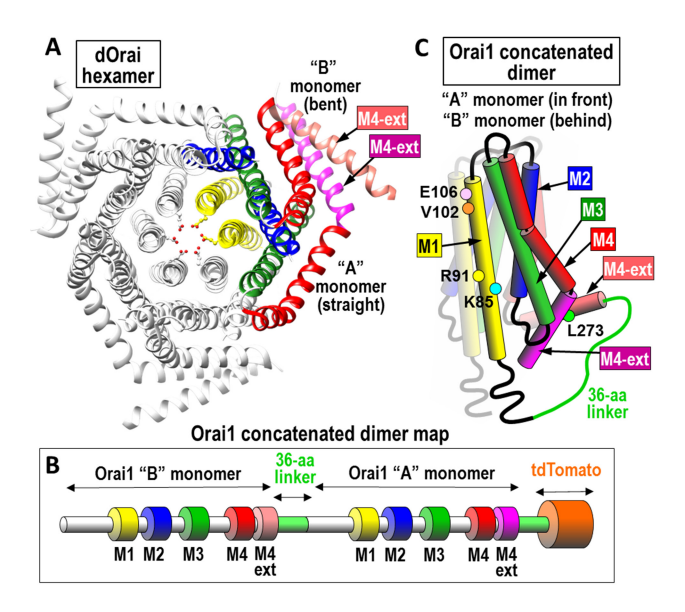


Figure 1. Structure of Orai channel and dimeric subunits. (A) Top view of *Drosophila* Orai channel hexamer. Two out of the six monomers are colored to show one of the three dimers that constitute the channel. The two subunits in each dimer are almost identical except the "A" monomer has a straight M4 extension helix (M4-ext; purple) and the "B" monomer has a bent M4-ext (pink). The two M4-ext helices are linked in an antiparallel configuration by hydrophobic interactions. (B) Linear sequence of the concatenated Orai1 dimer constructs used; the two monomers are joined through a 36-amino acid linker sequence, and tagged with tdTomato at the C-terminus. Note: the sequences of the "A" and "B" monomers are the same. (C) Schematic of the concatenated Orai1 dimer structure. The membrane-spanning domains (M1, M2, M3, M4, and M4-ext) are labeled as shown. Colored dots reveal each of the point mutations used in this study.

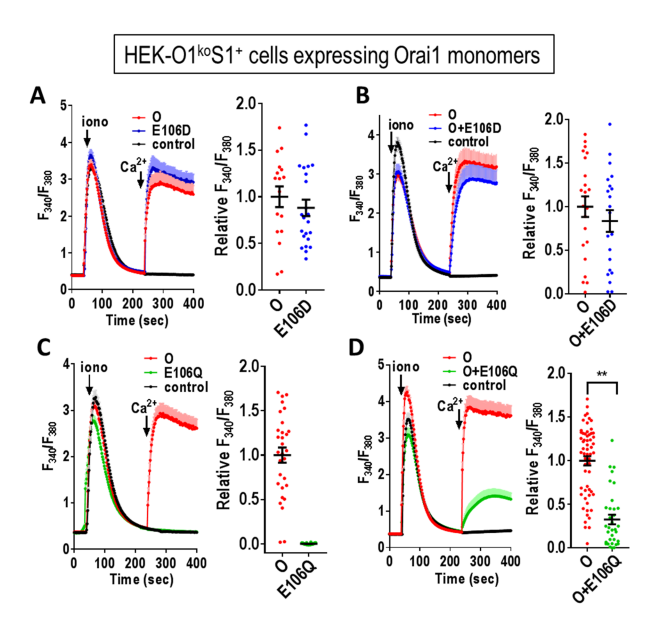


Figure 2. Consequences of mutating the selectivity filter (E106) within expressed Orai1 monomers. tdTomato-tagged Orai1 monomer constructs were transiently expressed in HEK-O1^{ko}S1⁺ cells which stably express STIM1-YFP. Fura-2 ratiometric Ca²⁺ measurements were conducted to reveal cytosolic Ca²⁺ after store-depletion with 2.5 μ M ionomycin in Ca²⁺-free medium followed by addition of 1 mM Ca²⁺ (arrows). (A) Cells were transfected with either Orai1-E106D (E106D) or Orai1-WT (O), or untransfected (control). (B) Cells were co-transfected with both Orai1-E106D-CFP and Orai1 WT, or untransfected (control). (C) Cells were transfected with either Orai1-E106Q (E106Q) or Orai1-WT (O), or untransfected (control). (D) Cells were co-transfected with both Orai1-E106Q-CFP and Orai1-WT, or untransfected (control). (D) Cells were co-transfected with both Orai1-E106Q-CFP and Orai1-WT, or untransfected (control). All the traces (means ± SE) are representative of three independent experiments. Summary scatter plots with means ± SEM for normalized peak Ca²⁺ entry are for all individual cells recorded in three independent experiments. (** P < 0.005)

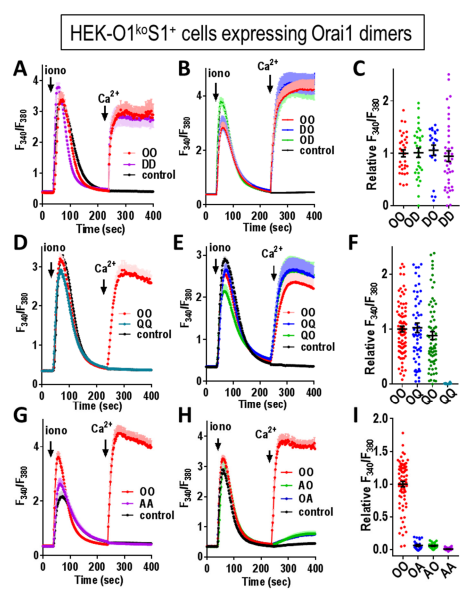


Figure 3. Function of concatenated Orai1 dimers containing the E106D and E106O selectivity filter mutations. Orai1-tdT dimer constructs were transiently expressed at similar levels in HEK-O1^{ko}S1⁺ cells and fura-2 ratiometric Ca²⁺ measurements were made to determine cytosolic Ca²⁺ after store-depletion with 2.5 μ M ionomycin in Ca²⁺-free medium, followed by addition of 1 mM Ca²⁺ (arrows). (A) Cells were transfected with either Orai1-WT homodimer (OO) or Orai1-E106D homodimer (DD), or untransfected (control). (B) Cells were transfected with either OO or Orai1-WT/Orai1-E106D (OD), Orai1-E106D/Orai1-WT (DO), heterodimers, or untransfected (control). (C) Summary scatter plots with means ± SEM for peak Ca^{2+} entry of all individual cells recorded in three independent experiments with the Orail concatemer constructs shown in (A,B). (D) Cells were transfected with either Orai1-WT homodimer (OO) or Orai1-E106Q mutant homodimer (QQ), or were untransfected (control). (E) Cells were transfected with either OO, or with Orai1-WT/Orai1-E106Q (OQ) or Orai1-E106Q/Orai1-WT (QO) dimer, or were untransfected (control). (F) Summary scatter plots with means \pm SEM for peak Ca²⁺ entry of all individual cells recorded in three independent experiments with the Orail concatemer constructs shown in (D,E). (G) Cells were transfected with either Orai1-WT dimer (OO) or Orai1-E106A homodimer (AA), or were untransfected (control). (H) Cells were transfected with either OO or Orai1-WT/Orai1E106A (OA) or Orai1-E106A/WT dimer (AO), or were untransfected (control). (F) Summary scatter plots with means \pm SEM for peak Ca²⁺ entry of all individual cells recorded in three independent experiments with the Orail concatemer constructs shown in (G,H). All the traces shown (means \pm SE) are representative of three independent experiments.

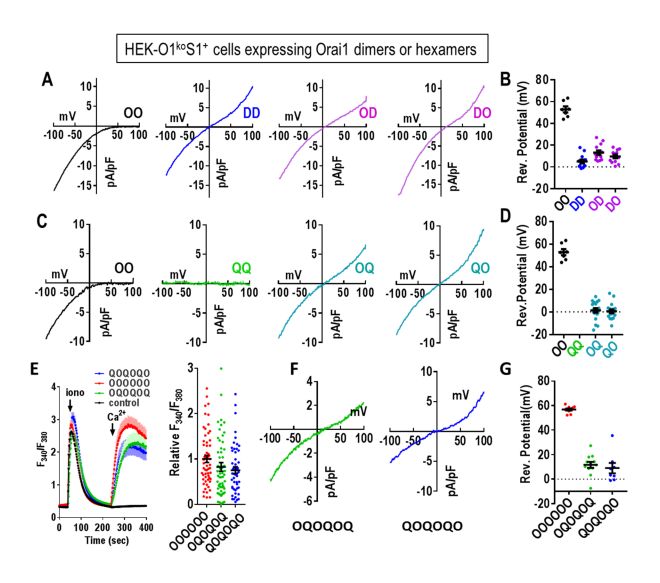


Figure 4. Electrophysiological properties of Orail E106D and E106Q concatenated dimers. (A) Representative I/V plots from whole cell recordings using HEK-O1^{ko}S1⁺ cells transiently expressing each of the tdT-tagged concatenated Orail dimer constructs (OO, DD, OD and DO) as described in Fig. 3. (B) Summary scatter plot (means \pm SEM) of reversal potential measurements taken from multiple I/V curves for the OO, DD, OD and DO dimers shown in (A). (C) Representative I/V plots from whole cell recordings from HEK-O1^{ko}S1⁺ cells transiently expressing each of the tdT-tagged concatenated Orai1 dimer constructs (OO, QQ, OQ and QO) as described in Fig. 3. (D) Summary scatter plot (means \pm SEM) of reversal potential measurements taken from multiple I/V curves for the QQ, OQ and QO as shown in (C). Current for the QQ construct was essentially zero. (E) Fura-2 ratiometric Ca^{2+} measurements reveal cytosolic Ca^{2+} after storedepletion with 2.5 μ M ionomycin in Ca²⁺-free medium followed by addition of 1 mM Ca²⁺ (arrows). In each case, traces for Orai1-WT hexamer (OOOOOO) were compared with traces for Orai1 heterohexamers (OQOQOQ, QOQOQO). Untransfected control cells are also shown. Traces shown (means \pm SE) are representative of three independent experiments. Summary scatter plots with means \pm SEM for peak Ca²⁺ entry of all individual cells recorded in three independent experiments with the Orail concatemer constructs shown. (F) Representative I/V relationship for the OQOQOQ and QOQOQO hexamers are shown. (G) Summary scatter plot (means \pm SEM) of reversal potential measurements taken from multiple I/V curves for the OOOOOO, OQOQOQ and QOQOQO hexamers as shown in (F).

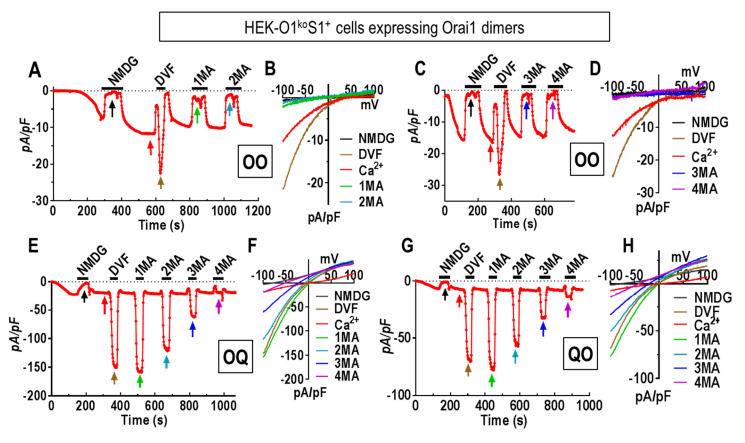


Figure 5. Altered pore properties of the non-selective Orail concatenated channel dimers, OQ and QO. (A-D) Electrophysiological measurements using HEK-O1^{ko}S1⁺ cells expressing the Orai1-WT dimer (OO). Current developed initially in the presence of 20 mM Ca²⁺ in the external solution. Thereafter, the external solution was periodically switched from 20 mM Ca²⁺ with 130 mM Na⁺ to divalent cation free solutions containing the following: 150 mM NMDG; 150 mM Na⁺ (DVF); 150 mM methylammonium (1MA); 150 mM dimethylammonium (2MA); 150 mM trimethylammonium (3MA); and finally 150 mM tetramethylammonium (4MA). In between each divalentfree addition, cells were switched back to 20 mM Ca²⁺ with 130 mM Na⁺. Current at -100 mV was plotted against time, and I/V relationships are shown (B, D) at the times indicated by arrows in A and C. (E) Using HEK-O1^{ko}S1⁺ cells expressing the OQ dimer, current developed in 20 mM external Ca²⁺. After switching to NMDG, divalent solutions were again added in the external as shown in A and C. For channel formed by OQ heterodimers, methylated ammonium cation still can produce significant currents. (F) I/V relationship of currents mediated by OQ collected in different external solutions as indicated by the arrows in (E). (G) HEK-O1^{ko}S1⁺ cells expressing the QO dimer were exposed to external solutions as shown in (E). (H) I/V relationship of currents mediated by QO collected in the external solutions indicated in (G).

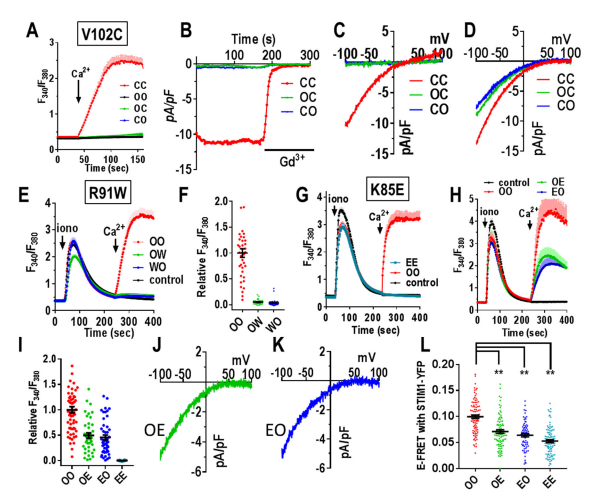


Figure 6. Mutation of key Orail channel pore residues in concatenated dimers. (A) Fura-2 ratiometric Ca²⁺ addback measurements for HEK-O1^{ko} cells expressing similar levels of the C-terminal tdT-tagged Orai1 dimers OO (Orai1-WT/Orai1-WT), OC (Orai1-WT/Orai1-V102C), (Orai1-V102C/Orai1-WT), or \widetilde{CC} (Orai1-V102C/Orai1-V102C). Constitutive Ca²⁺ entry was measured after addition of 1 mM Ca²⁺ (arrows) after Ca²⁺-free medium. Traces (means \pm SEM) are representative of three independent experiments. (B) Whole cell patch clamp recording of HEK- $O1^{ko}$ cells expressing CC, OC or CO; current at -100 mV was plotted against time. Cytosolic Ca²⁺ was maintained at 90 nM with BAPTA to prevent store-depletion. Constitutive current was measured in 20 mM Ca^{2+} external solution, followed by blockade with 10 μ M Gd³⁺. (C) Representative I/V curves for the constitutive currents of the HEK-O1^{ko} cells expressing OC, CO and CC shown in (B). (D) Representative I/V curves for Ca^{2+} currents following store depletion with 20 mM BAPTA in HEK-O1^{ko}S1⁺ cells transiently expressing OC, CO and CC. (E) Ca²⁺ add-back measurements in fura-2-loaded HEK-O1^{ko}S1⁺ cells expressing similar levels of the C-terminal tdT-tagged Orai1 dimers OO (Orai1-WT/Orai1-WT), OW (Orai1-WT/Orai1-R91W), or WO (Orai1-R91W/Orai1-WT). Ca²⁺ stores were released with 2.5 μ M ionomycin in Ca²⁺-free medium followed by 1 mM Ca²⁺ (arrows). Traces (means ± SEM) are the results for all cells in three independent experiments. (F) Summary scatter plots of peak Ca^{2+} entry normalized to Ca^{2+} entry with WT dimer (OO); results are means \pm SEM for all cells in three independent experiments represented by (E). (G,H) Ca²⁺ add-back measurements using fura-2-loaded HEK-O1^{ko}S1⁺ cells expressing similar levels of the Cterminal tdT-tagged Orai1-K85E homodimers (EE) or the K85E heterodimers (OE, EO). Ca²⁺ stores were released with 2.5 μ M ionomycin in Ca²⁺-free medium followed by 1 mM Ca²⁺ (arrows). Traces (means ± SE) are representative of all cells in three independent experiments. (I) Summary scatter plots of peak Ca^{2+} entry normalized to wildtype (OO) dimer Ca^{2+} entry ; results are means \pm SEM of three independent experiments represented in (G and H). Representative I/V relationship of Ca^{2+} currents after store depletion in HEK-O1^{ko}S1⁺ cells transiently expressing OE (J) or EO (K). (L) Scatter plots of E-FRET between stably expressed STIM1-YFP and transiently expressed C-terminally CFP-tagged Orai1 dimers (OO, OE, EO or EE, respectively) in HEK-O1^{ko}S1⁺ cells. E-FRET was measured at 5 mins after addition of 2.5 μ M ionomycin to deplete stores. Results are means \pm SEM of three independent experiments.

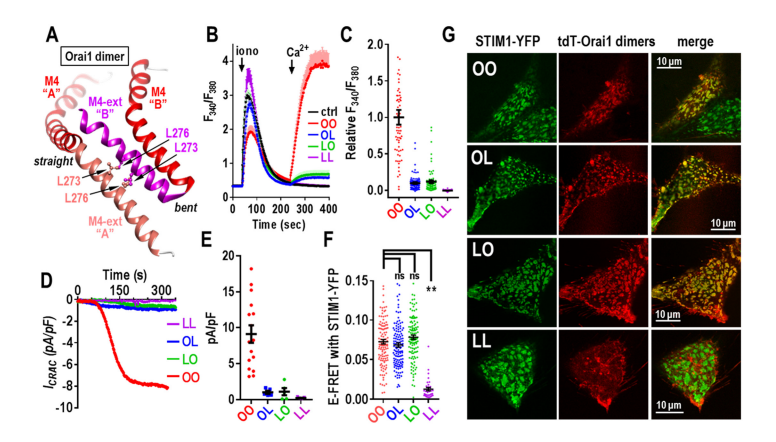


Figure 7. Effects of mutating the key STIM1-binding residue (L273) of the Orai1 C-terminus in concatenated **Orail dimers.** (A) Structural model Orail based on the dOrai M4 and M4 extensions (M4-ext), showing their configuration and interactions. The M4-ext from the straight monomer "A" and the M4-ext from the bent monomer "B" lie in a close antiparallel configuration stabilized by interactions between the L273 and L276 residues on each M4-ext. (B) Ca²⁺ entry in fura-2-loaded HEK-O1^{ko}S1⁺ cells expressing similar levels of the C-terminal tdT-tagged Orail dimer constructs: OO (Orail-WT/Orail-WT), OL (Orai-WT/Orail-L273D), LO (Orail-L273D/Orail-WT), LL (Orai1-L273D/Orai1-L273D), or untransfected control cells (ctrl). Ca²⁺ stores were released with 2.5 µM ionomycin in Ca²⁺-free medium followed by addition of 1 mM Ca²⁺ (arrows). Traces (means \pm SE) are representative of three independent experiments. (C) Summary scatter plots with means \pm SEM of peak Ca²⁺ entry normalized to peak Ca^{2+} entry for the OO dimer; all cells are shown from three independent experiments. (D) Whole cell patch clamp recording of HEK-O1^{ko}S1⁺ cells expressing OO, OL, LO or LL, currents at -100 mV plotted against time. (E) Summary scatter plots of means \pm SEM of peak Ca²⁺ current density for all individual cells recorded for each of the Orail dimer constructs shown in (D). (F) Scatter plots of E-FRET between stably expressed STIM1-YFP and transiently expressed C-terminally CFP-tagged Orai1 dimers (OO, OL, LO or LL, respectively) in HEK-O1^{ko}S1⁺ cells. E-FRET was measured at 5 mins after addition of 2.5 μ M ionomycin to deplete stores. Results are means \pm SEM of three independent experiments. (G) Confocal images showing the localization of stably expressed STIM1-YFP and transiently-transfected Orai1-tdTdimers (OO, OL, LO, and LL) expressed in HEK-O1^{ko}S1⁺ cells. Images were taken 5 mins after 2.5 µM ionomycin treatment to empty stores. In each case, confocal images show STIM1 and Orail-dimers localized only in the PM layer immediately adjacent to the coverslip. For OO (top row), there are two cells; the centered cell expresses both STIM1-YFP and OO-tdT, whereas the lower right cell expresses only STIM1-YFP. Note: there is near-perfect co-localization of STIM and with the OO, OL, and LO Orai1-dimers; in contrast, there is no co-localization of STIM1 with the LL Orai1-dimer.

Pore properties of Orai1 calcium channel dimers and their activation by the STIM1 ER calcium sensor

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J. Biol. Chem. published online June 28, 2018

Access the most updated version of this article at doi: 10.1074/jbc.RA118.003424

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