Crosstalk between N-terminal and C-terminal domains in stromal interaction molecule 2 (STIM2) determines enhanced STIM2 sensitivity

By

Scott M. Emrich¹,#, Ryan E. Yoast¹,#, Ping Xin¹, Xuexin Zhang¹, Trayambak Pathak¹, Robert Nwokonko¹, Maxime F. Gueguinou¹, Krishna P. Subedi³, Yandong Zhou¹, Indu S. Ambudkar³, Nadine Hempel², Khaled Machaca⁴, Donald L. Gill¹, and Mohamed Trebak¹,*

From

¹Department of Cellular and Molecular Physiology, and ²Department of Pharmacology, the Pennsylvania State University College of Medicine, Hershey, PA, 17033 USA; ³Secretory Physiology Section, NIDCR, NIH, Bethesda, MD 20892, US and ⁴Department of Physiology and Biophysics, Weill Cornell Medical College in Qatar, Education City, Qatar Foundation, P.O. Box 24144, Doha, Qatar.

#SME and REY contributed equally to this work

*Correspondence to: Mohamed Trebak (mtrebak@psu.edu)

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ABSTRACT

Store-operated Ca\(^{2+}\) entry (SOCE) is a ubiquitous pathway for Ca\(^{2+}\) influx across the plasma membrane (PM). SOCE is mediated by the endoplasmic reticulum (ER)-associated Ca\(^{2+}\)-sensing proteins stromal interaction molecule 1 (STIM1) and STIM2, which transition into an active conformation in response to ER Ca\(^{2+}\) store depletion, thereby interacting with and gating PM-associated ORAI1 channels. Although structurally homologous, STIM1 and STIM2 generate distinct Ca\(^{2+}\) signatures in response to varying strengths of agonist stimulation. The physiological functions of these Ca\(^{2+}\) signatures, particularly under native conditions, remain unclear. To investigate the structural properties distinguishing STIM1 and STIM2 activation of ORAI1 channels under native conditions, here we used CRISPR/Cas9 to generate STIM1\(-/-\), STIM2\(-/-\), and STIM1/2\(-/-\) knockouts in HEK293 and colorectal HCT116 cells. We show that depending on cell type, STIM2 can significantly sustain SOCE in response to maximal store depletion. Utilizing the SOCE modifier 2-aminoethoxydiphenylborate (2-APB), we demonstrate that 2-APB–activated store-independent Ca\(^{2+}\) entry is mediated exclusively by endogenous STIM2. Using variants that either stabilize or disrupt intramolecular interactions of STIM C termini, we show that the increased flexibility of the STIM2 C terminus contributes to its selective store-independent activation by 2-APB. However, STIM1 variants with enhanced flexibility in C terminus failed to support its store-independent activation. STIM1/STIM2 chimeric constructs indicated that coordination between N-terminal sensitivity and C-terminal flexibility is required for specific store-independent STIM2 activation. Our results clarify the structural determinants underlying activation of specific STIM isoforms, insights that are potentially useful for isoform-selective drug targeting.

INTRODUCTION

In non-excitable cells, cytosolic Ca\(^{2+}\) concentrations are controlled through release from intracellular Ca\(^{2+}\) stores of the endoplasmic reticulum (ER) and Ca\(^{2+}\) influx across the plasma membrane (PM) through the ubiquitous store-operated Ca\(^{2+}\) entry (SOCE) pathway (1). SOCE is required for the acute refilling of ER Ca\(^{2+}\) stores and sustaining downstream signaling to gene transcription in most cell types (2-5). SOCE is mediated by the ER Ca\(^{2+}\) sensors stromal interaction molecules (STIM1 and STIM2), which directly bind, crosslink, and gate plasma membrane ORAI1 channels in response to store depletion. Extensive investigation into the molecular interactions between STIM1 and ORAI1 has revealed STIM1 as a potent activator of ORAI1 channels through a strategically positioned phenylalanine (F394) located within the STIM-ORAI-activating region (SOAR1) in STIM1 C-terminus (6-8). Furthermore, the N-terminal EF-hand domain of STIM1 located within the ER lumen has a high affinity for Ca\(^{2+}\), requiring significant depletion of ER Ca\(^{2+}\) stores for the transition of STIM1 into an active conformation (9,10). In contrast, the structurally homologous STIM2 is a relatively weak activator of ORAI1, in which the residue equivalent to F394 of STIM1 is a leucine (L485) within the SOAR domain of STIM2 (SOAR2) that renders it a partial agonist (6). The luminal N-terminal EF-hand domain of STIM2 located within the ER lumen has a high affinity for Ca\(^{2+}\), requiring significant depletion of ER Ca\(^{2+}\) stores for the transition of STIM1 into an active conformation (9,10).
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maintain basal cytosolic and ER Ca²⁺ concentrations (11,12). Interestingly, a previous study proposed that the short N-terminal peptide before the EF-hand domain in STIM2 functions as "brake" to ensure negative regulation of STIM2 constitutive activity and prevent significant enhancement of basal cytosolic Ca²⁺ concentration (13). A recent report further showed that E470G mutation within the SOAR2 domain of STIM2 (E470 is equivalent to G379 in STIM1) renders STIM2 a fast and robust activator of ORAI1, while the opposite mutation in STIM1 only weakly activates ORAI1 (14). The same group showed that the transmembrane (TM) domain of STIM2 contributes to its slow activation kinetics compared to STIM1 (14). While STIM1 has been shown to drive the majority of SOCE in a variety of cell types in response to agonist-induced store depletion (9,15-17), the physiological function of STIM2 in this process is not clear. Further adding to the complexity of this system is recent evidence demonstrating that STIM2 plays a role in inducing the active conformation of STIM1 and regulating its coupling with ORAI1 channels under conditions of replete ER Ca²⁺ stores (18,19). The physiological implications of this mechanism, particularly how endogenous STIM proteins utilize this molecular coordination to drive cell function, are unknown. Additionally, the structural differences between STIM1 and STIM2 that determine their differential response to varying strengths of agonist and the distinct roles of STIM1 and STIM2 in cell physiology, especially under native conditions, are largely unknown.

Multiple classes of pharmacological compounds which either enhance or inhibit SOCE activity have been identified (20,21). Among these SOCE channel modifiers, the compound 2-aminoethoxydiphenyl borate (2-APB) has been well characterized and widely utilized (22-24). 2-APB displays a unique biphasic effect on ORAI1 activity, potentiating it at low concentrations (1-10 μM) and inhibiting it at high concentrations (~30-50 μM) (25). High concentrations of 2-APB have been demonstrated to: 1) inhibit several constitutively active STIM-independent ORAI1 channel mutants while potentiating wild type ORAI3, suggesting that 2-APB acts directly on ORAI1 channel pore; 2) strengthen intramolecular interaction within the STIM1 C-terminus, between the coiled-coil 1 (CC1) domain and SOAR1, thus preventing STIM1 unfolding in response to store-depletion. This would result in a reduction in STIM1 oligomerization and puncta formation which would reduce functional interactions with ORAI1 (22,26-31). However, the ability of 2-APB at low concentrations to selectively increase Ca²⁺ influx through ORAI1 channels remains enigmatic.

While 2-APB is relatively nonspecific and can affect a diverse array of ion channels, a clear understanding of how 2-APB interacts with and regulates ORAI and STIM proteins under native levels of expression would be crucial for the future development of potent and selective SOCE modifying compounds (32-34). Here, we generated multiple cell lines using CRISPR/Cas9 technology that are devoid of individual STIM proteins (STIM1⁻, STIM2⁻) as well as double STIM1 and STIM2 knockout cells (STIM1/2⁻). Our results demonstrate that store-independent activation of SOCE with low concentrations of 2-APB (10 μM) occurs exclusively through endogenous STIM2. Through the use of chimeric STIM proteins and STIM mutant constructs, we clearly demonstrate that the effects of 2-APB on STIM2 requires the combination of increased flexibility of the STIM2 C-terminus and increased sensitivity of its N-terminal EF hand. Thus, the use of low concentrations of 2-APB as a pharmacological tool to selectively drive
native STIM2-mediated Ca\textsuperscript{2+} entry provides a novel strategy to dissect the physiological functions of STIM2 under homeostatic and endogenous conditions. Further, development of 2-APB-derived analogues with increased specificity and potency for STIM2 will selectively target cellular pathways that rely on STIM2 in both healthy and disease conditions.

RESULTS

Generation and characterization of STIM1\textsuperscript{-/-}, STIM2\textsuperscript{-/-}, and STIM1/2\textsuperscript{-/-} knockout cell lines

We utilized CRISPR/Cas9 technology to delete each STIM protein individually and in combination in HEK293 cells. Western blot analysis of HEK293 STIM1\textsuperscript{-/-} and STIM2\textsuperscript{-/-} cell lines demonstrated no compensatory upregulation of STIM1 or STIM2 proteins in response to individual STIM knockout, while STIM1/2\textsuperscript{-/-} cells were devoid of both proteins (Fig. 1A, B). Fura2 Ca\textsuperscript{2+} imaging was performed in each HEK293 knockout cell line to measure SOCE in response to store depletion with the sarco/endoplasmic reticulum Ca\textsuperscript{2+} ATPase (SERCA) pump inhibitor thapsigargin (Fig. 1C, D). HEK293 STIM1\textsuperscript{-/-} cells exhibit substantial inhibition in peak SOCE (to \textasciitilde10\% of WT), while STIM2\textsuperscript{-/-} cells showed only minor reduction in peak SOCE (\textasciitilde84\% of WT). Knockout of both STIM1 and STIM2 (HEK293 STIM1/2\textsuperscript{-/-} cells) results in essentially complete abrogation of SOCE (\textasciitilde3\% of WT) upon re-addition of 2 mM Ca\textsuperscript{2+} to the extracellular medium. To investigate cell type-dependent contributions of STIM1 versus STIM2 to Ca\textsuperscript{2+} signals, we also generated individual STIM1 and STIM2 knockout of the colorectal cancer cell line HCT116 and documented knockouts with Western blots (Fig. 1E, F). Ca\textsuperscript{2+} imaging experiments demonstrated a significant reduction in peak SOCE in HCT116 STIM1\textsuperscript{-/-} cells (\textasciitilde24\% of WT) (Fig. 1G, H). Interestingly, HCT116 STIM2\textsuperscript{-/-} cells also showed substantial reduction in SOCE (\textasciitilde44\% of WT) despite having STIM1 protein present. These results suggest that STIM2 plays differential, cell type-specific role in supporting SOCE in response to maximal ER store depletion.

2-APB activates store-independent Ca\textsuperscript{2+} entry exclusively through STIM2

Using our newly generated HEK293 and HCT116 STIM knockout cell lines, we investigated the effects of low (10 \textmu M) and high (50 \textmu M) 2-APB under conditions where internal Ca\textsuperscript{2+} stores were replete. To address potential off target effects of CRISPR/Cas9, we generated additional STIM1 and STIM2 knockout clones in both cell lines using multiple guide RNA sequences (Fig. 2A, F). Our results presented in detail below show that STIM2, but not STIM1, is crucial for 2-APB-activated Ca\textsuperscript{2+} entry. To strengthen this argument, we present data obtained from 4 STIM2\textsuperscript{-/-} HCT116 cell clones representing 2 independent clones each obtained from two independent guide RNA (e.g. STIM2\textsuperscript{-/-}g1.1 corresponds to clone 1 from guide RNA 1 etc; Fig. 2F). Stimulation of wild-type HEK293 and HEK STIM1\textsuperscript{-/-} cells with 10 \textmu M 2-APB demonstrated a small yet sustained Ca\textsuperscript{2+} entry, which was not observed in HCT116 STIM2\textsuperscript{-/-} cells (Fig. 2G, H). Addition of 50 \textmu M 2-APB to different variants of HEK293 cells showed a marginal and transient
increase in cytosolic Ca\(^{2+}\) (Fig. 2D, E). Surprisingly, stimulation of wild-type HCT116 cells with 50 µM 2-APB showed a more pronounced, rapid and transient increase in cytosolic Ca\(^{2+}\), and this effect was further enhanced in HCT116 STIM1\(^{-/-}\) cells (Fig. 2I, J). This rapid potentiation with 50 µM 2-APB was not observed in HCT116 STIM2\(^{-/-}\) cells (Fig. 2I, J). These results suggest that 2-APB activates endogenous STIM2, but not STIM1, in both cell lines in a store-independent manner and that the more potent activation of Ca\(^{2+}\) entry by 50 µM 2-APB in HCT116 cells results from the more prominent role STIM2 plays in SOCE of HCT116 cells.

Previous studies showed that overexpression of wild-type (WT) STIM2 in wild-type HEK293 cells results in pre-clustered STIM2 puncta located at junctions between the ER and plasma membrane in the absence of store-depletion (11). Here, we utilized HEK293 STIM1/2\(^{-/-}\) cells with a clean genetic background lacking endogenous STIM proteins. When expressed in STIM1/2\(^{-/-}\) HEK293 cells, YFP-tagged STIM2 displayed significant puncta formation throughout the entirety of the cell under basal conditions (Fig. 3A). While treatment with 10 µM 2-APB caused a reduction in cell size and reorganization of STIM2 puncta, it did not lead to significant enhancement of puncta size from baseline (Fig. 3B). In contrast, overexpressed STIM1 protein displayed a diffuse, tubular distribution (Fig. 3A), and STIM1 did not form puncta in response to 10 µM 2-APB (Fig. 3A). To compare the behavior of native vs overexpressed STIMs, Ca\(^{2+}\) measurements were performed comparatively in four different groups of HEK293 cells: 1) STIM1/2\(^{-/-}\) cells overexpressing STIM2; 2) STIM1/2\(^{-/-}\) cells overexpressing STIM1; 3) STIM1\(^{-/-}\) cells; and 4) STIM2\(^{-/-}\) cells. Direct addition of 10 µM 2-APB to STIM1/2\(^{-/-}\) cells overexpressing STIM2 induced a large and sustained store-independent increase in cytosolic Ca\(^{2+}\) concentrations (Fig. 3C, D). Addition of 10 µM 2-APB to STIM1/2\(^{-/-}\) cells overexpressing STIM1 caused store-independent Ca\(^{2+}\) entry comparable in magnitude to that observed in HEK293 STIM1\(^{-/-}\) cells with endogenous STIM2 (Fig. 3C, D). Similar results were observed in response to 50 µM 2-APB stimulation, with STIM1/2\(^{-/-}\) cells overexpressing STIM2 demonstrating a rapid and transient store-independent increase in cytosolic Ca\(^{2+}\) (Fig. 3E, F). Overexpression of STIM1 in STIM1/2\(^{-/-}\) cells yielded a marginal increase in cytosolic Ca\(^{2+}\) with 50 µM 2-APB stimulation.

Depletion of intracellular Ca\(^{2+}\) stores induces rapid puncta formation of STIM1 and induces clustering with ORAI1 channels (7,35). High concentrations of 2-APB (50 µM) inhibit the formation of STIM1 puncta by clamping the STIM1 C-terminus in an inactive conformation (28). Interestingly, these inhibitory effects on STIM1 puncta formation are abolished in cells overexpressing both STIM1 and ORAI1 (24,28,31), suggesting that enhanced STIM1/ORAI1 expression and coupling can overcome 2-APB mediated inhibition of the STIM1 C-terminus. Therefore, we tested whether overexpression of ORAI1 could rescue store-independent Ca\(^{2+}\) entry driven by STIM1 in response to 2-APB treatment. Stimulation with 10µM 2-APB of STIM1/2\(^{-/-}\) cells overexpressing STIM2 and ORAI1 showed significantly enhanced Ca\(^{2+}\) entry compared to STIM1/2\(^{-/-}\) cells overexpressing STIM2 alone (Fig. 4A, B). This store-independent Ca\(^{2+}\) entry with STIM2/ORAI1 overexpression was even bigger, albeit transient, with 50 µM 2-APB (Fig. 4E, F). In contrast, overexpression of STIM1 with ORAI1 marginally enhanced Ca\(^{2+}\) entry in response to 10 µM 2-APB compared to STIM1/2\(^{-/-}\) cells overexpressing STIM1 alone (Fig. 4A, B). Stimulation of STIM1/2\(^{-/-}\) cells
overexpressing STIM1/ORAI1 with 50 μM 2-APB significantly enhanced Ca^{2+} entry compared to cells overexpressing STIM1 alone, although the magnitude of this effect was drastically less than with STIM2/ORAI1 expression (Fig. 4E, F).

Stimulation of STIM1/2^-/- cells overexpressing STIM2/ORAI1 with 10 μM 2-APB further enhanced the size of pre-clustered STIM2 puncta (Fig. 4C, D), while 2-APB stimulation did not induce puncta formation in STIM1/ORAI1 overexpressing cells (Fig 4G). Taken together, these results suggest that low and high concentrations of 2-APB induce store-independent Ca^{2+} entry through STIM2. Moderate 2-APB-mediated activation of STIM1 is observed only in cells overexpressing STIM1 and ORAI1, and stimulated with high concentrations of 2-APB (50 μM; Fig. 4E, F), a finding that is in agreement with previous reports (24).

Low concentrations of 2-APB potentiate SOCE exclusively through STIM2

We next examined the effects of 2-APB on the potentiation and inhibition of SOCE activated through agonist stimulation. For reasons that will become clear below, we used 100μM carbachol with 2 mM extracellular Ca^{2+} or 150μM ATP with 10 mM extracellular Ca^{2+} for HEK293 cells and 300 μM ATP with 10 mM extracellular Ca^{2+} for HCT116 cells. Stimulation with carbachol followed by re-addition of Ca^{2+} to the extracellular medium induced SOCE in WT HEK293 cells. While HEK293 STIM2^-/- cells showed only partial inhibition of SOCE, minimal SOCE was detected in STIM1^-/-, and essentially no SOCE was observed in STIM1/2^-/- cells (Fig. 5A). The same protocol using stimulation with ATP showed marginal Ca^{2+} entry in all three knockout cell lines, STIM1^-/-, STIM2^-/- and STIM1/2^-/- (Fig. 5B), suggesting that ATP is a weak agonist that relies on both STIM1 and STIM2 for SOCE activation. Indeed, 10 mM external Ca^{2+} was used to enhance the driving force and detect a Ca^{2+} signal in WT HEK293 cells. In all conditions however, subsequent addition of 10 μM 2-APB consistently induced rapid potentiation of SOCE in STIM1^-/- cells, while no potentiation was observed in STIM2^-/- or STIM1/2^-/- cells (Fig. 5A, B). SOCE was inhibited by the sequential addition of 50 μM 2-APB in all cell lines (Fig. 5A, B). WT HEK293 cells showed marginal potentiation with 10 μM 2-APB when SOCE was stimulated with carbachol (Fig. 5A), likely because carbachol causes maximal depletion of ER stores and maximal activation of SOCE compared to ATP (Fig. 5B). Similar results were observed with ATP in HCT116 cells. Supramaximal ATP concentrations (300 μM with 10 mM Ca^{2+} extracellular; 150 μM ATP did not yield consistent SOCE activation) activated moderate SOCE in WT HCT116 cells, with marginal SOCE detected in STIM1^-/- and STIM2^-/- cells (Fig. 5C). Addition of 10 μM 2-APB potentiated SOCE in WT HCT116 and STIM1^-/- cells, but not in STIM2^-/- cells. This potentiation was rapidly inhibited following addition of 50 μM 2-APB (Fig. 5C). Despite repeated attempts with various protocols, HCT116 cell lines did not respond to carbachol stimulation.

Flexibility of the STIM2 C-terminus is critical to 2-APB Selectivity

Under conditions of high ER Ca^{2+} concentrations, the C-terminus of STIM1 and STIM2 are maintained in an inactive and clamped confirmation characterized by strong molecular interactions between the CC1 and CC3 domains, which shield SOAR domains from ORAI1 (14,18,36-39). Depletion of ER stores and loss of Ca^{2+} binding to the luminal EF hand domains of STIM1 and STIM2 triggers rearrangement of the N- and C-terminus, opening the CC1/CC3 clamp and exposing either SOAR1 or...
SOAR2 for gating and activation of ORAI1 channels (2,4,6,37,38,40-42). Accordingly, 2-APB induced store-independent Ca\(^{2+}\) entry is mediated through proper interaction and gating of SOAR2 with ORAI1 channels as the dominant negative splice variant STIM2.1, containing an eight amino acid insertion within SOAR2(43,44), demonstrated no increase in cytosolic Ca\(^{2+}\) concentrations in response to 10 μM 2-APB (Fig. 6A, B). In conjunction with the SOAR domains (Fig. 6C), studies have described key residues within the C-terminal CC1 and CC3 domains that contribute to the induction of the active conformation of STIM1 and STIM2 (37,40). Mutation of residues located within the ORAI1 Activating Small Fragment (OASF1 (37,40); Fig. 6C) region of STIM1 have been identified to either stabilize (R426L) or disrupt (L251S) CC1/CC3 interactions (Fig. 6C). Comparison of corresponding mutations in STIM2 that stabilize (R517L) or disrupt (L342S/L507S/L514S, CCmt) CC1/CC3 interactions suggest that the OASF region of STIM2 (OASF2) adopts a weaker CC1/CC3 interaction generating a more flexible conformation of OASF2 compared to OASF1 (19). Utilizing intramolecular FRET-based OASF1 and OASF2 sensors expressed in HEK293 STIM1/2-/- cells, we confirm previous findings of significantly reduced intramolecular FRET signal of the OASF2 region of STIM2 compared to OASF1 of STIM1 (Fig. 6D). We next evaluated the effects of CC1/CC3 stabilization or disruption mutations on 2-APB-mediated Ca\(^{2+}\) entry by expressing these mutations within the context of full length STIM1 and STIM2 proteins in HEK293 STIM1/2\(^{-/}\) cells. Similar to overexpression of ORAI1, even high concentrations (75 μM) of 2-APB caused a marginal decrease in OASF1 intramolecular FRET (37).

OASF2 mutant displayed significantly decreased FRET signal compared to OASF1 (Fig. 6E), suggesting even further flexibility of this mutant. The stabilized OASF2-R517L mutant demonstrated significantly higher FRET signal than OASF2, which was still substantially less than OASF1 (Fig. 6E). We next tested the ability of 10 μM 2-APB to induce changes in intramolecular FRET of OASF1 and OASF2 constructs and observed marginal FRET changes with all different OASF variants (Fig. 6F, G; see Fig. 6H, I for zoom-in normalized FRET), suggesting that any rearrangements of C-terminal STIM caused by 2-APB are likely very subtle. These results are consistent with data from the Romanin group showing that, without overexpression of ORAI1, even high concentrations (75 μM) of 2-APB caused a marginal decrease in OASF1 intramolecular FRET (37).

We next evaluated the effects of CC1/CC3 stabilization or disruption mutations on 2-APB-mediated Ca\(^{2+}\) entry by expressing these mutations within the context of full length STIM1 and STIM2 proteins in HEK293 STIM1/2\(^{-/}\) cells. Similar to overexpression of STIM2, both the flexible CCmt-STIM2 and stabilized R517L-STIM2 mutants displayed extensive preformed puncta (Fig. 7A). While all STIM2 mutants showed reorganization of baseline puncta in response to the change in shape of cells, 2-APB did not lead to significant enhancement of puncta size (Fig. 7B). Stimulation of STIM2-expressing cells with 10 μM 2-APB induced rapid and sustained store-independent Ca\(^{2+}\) entry (Fig. 7C, D), as shown in Fig. 3C, D. Interestingly, the flexible CCmt-STIM2 displayed substantially increased Ca\(^{2+}\) entry compared to STIM2, while the stabilized R517L-STIM2 supported Ca\(^{2+}\) entry that was slightly reduced compared to STIM2 (Fig. 7C, D). Similar results were obtained when STIM2 mutants were stimulated with 50 μM 2-APB.
However, the stabilized R517L-STIM2 mutant displayed nearly identical Ca\(^{2+}\) entry compared to STIM2 (Fig. 7E, F).

STIM1 and the stabilized R426L-STIM1 mutant expressed in STIM1/2\(^{-/-}\) cells displayed tubular localization throughout the cell, while the flexible L251S-STIM1 had extensive preformed puncta (Fig. 8A). Stimulation of STIM1 with 10 \(\mu\)M 2-APB induced minimal store-independent Ca\(^{2+}\) entry, which was completely abolished in the stabilized R426L-STIM1 mutant (Fig. 8C, D). The flexible L251S-STIM1 mutant showed preformed puncta and constitutive Ca\(^{2+}\) activity (Fig. 8B) as previously reported (37). Unexpectedly however, it supported minimal Ca\(^{2+}\) entry in response to 10 \(\mu\)M 2-APB that was comparable to STIM1 (Fig. 8B). In response to 50 \(\mu\)M 2-APB, STIM1 and the flexible L251S-STIM1 mutants showed similar patterns of small initial Ca\(^{2+}\) entry to those seen on stimulation with 10 \(\mu\)M 2-APB, while the stabilized R426L-STIM1 showed essentially no response (Fig. 8E, F). Furthermore, the basal activity of L251S-STIM1 mutant was subsequently inhibited by 50 \(\mu\)M 2-APB (Fig. 8E). Our results thus far suggest that while the increased flexibility of the STIM2 C-terminus plays a key role in driving Ca\(^{2+}\) entry in response to 2-APB and weak agonists, enhanced C-terminal flexibility alone cannot rescue the activation of STIM1 by 2-APB.

Recent reports have demonstrated that STIM2 remodels the STIM1 C-terminus under conditions of high ER Ca\(^{2+}\) concentrations to allow interactions with ORAI1 under low stimulus intensities (18,19). We examined this effect by co-expression of STIM1 and STIM2 in STIM1/2\(^{-/-}\) cells. Interestingly, the expression of both STIM1 and STIM2 led to significant constitutive activity by comparison to either STIM expressed alone (Fig. 8G). Stimulation with either 10 \(\mu\)M (Fig. 8H, I) or 50 \(\mu\)M (Fig. 8J, K) of 2-APB activated significant Ca\(^{2+}\) entry in cells expressing STIM2 or co-expressing both STIM proteins but not in cell expressing STIM1 alone.

**Coordination between STIM2 N- and C-termini is required for 2-APB sensitivity**

We next determined the contribution of the STIM2 N-terminus in mediating store-independent Ca\(^{2+}\) entry by constructing two chimeric STIM constructs: a chimera of STIM1 N-terminus with the STIM2 TM and C-terminus (S1N-S2C) and another of STIM2 N-terminus with the STIM1 TM and C-terminus (S2N-S1C; Fig. 9A). Interestingly, expression of S1N-S2C chimera in HEK293 STIM1/2\(^{-/-}\) cells showed preformed puncta throughout the cell (Fig. 9B, top), while expression of S2N-S1C chimera displayed diffuse ER localization similar to STIM1 (Fig. 9B, bottom), further confirming that STIM2 preformed puncta are driven by its C-terminus and not by the enhanced sensitivity of its N-terminus to store depletion(6). Each STIM chimera was overexpressed in STIM1/2\(^{-/-}\) cells to measure SOCE in response to store depletion with thapsigargin (Fig. 9C, D). Stimulation of S1N-S2C-expressing cells displayed similar release of ER Ca\(^{2+}\) compared to S2N-S1C-expressing cells and non-transfected STIM1/2\(^{-/-}\) cells. However, S2N-S1C overexpression resulted in increased constitutive activity (Fig. 9F, G). SOCE was significantly rescued with expression of either S1N-S2C or S2N-S1C chimeras. However, S2N-S1C expression supported bigger SOCE (Fig. 9C, D), likely the result of its enhanced constitutive activity. Interestingly, addition of 50 \(\mu\)M 2-APB strongly inhibited SOCE mediated by S2N-S1C chimera with less inhibitory effect on SOCE supported by the S1N-S2C chimera (Fig. 9C-E), suggesting the STIM2 C-terminus protects against 2-APB-mediated
inhibition. These findings are consistent with the reported constitutive activity of a similar STIM1 chimera containing the EF-SAM domain of STIM2 co-expressed with ORAI1 (45). We then determined the ability of the STIM chimeras to mediate store-independent Ca\(^{2+}\) entry in response to 2-APB stimulation when expressed in STIM1/2\(^{-/-}\) cells. As expected, overexpression of STIM2 supported a sustained Ca\(^{2+}\) entry in response to 10 μM 2-APB, while STIM1 overexpression mediated marginal Ca\(^{2+}\) entry under the same conditions (Fig. 9H, I). Unexpectedly, both S1N-S2C and S2N-S1C chimeras mediated minimal Ca\(^{2+}\) entry comparable to STIM1 in response to 10 μM 2-APB (Fig. 9H, I). Similar results were observed when cells were stimulated with 50 μM 2-APB with each STIM chimera displaying minimal store-independent Ca\(^{2+}\) entry (Fig. 9J, K). These results suggest that both the high sensitivity of the STIM2 N-terminus and the communication with the flexible conformation of its C-terminus are key determinants for activation of store-independent Ca\(^{2+}\) entry in response to 2-APB, and likely in response to weak agonists or low concentrations of strong agonists.

**DISCUSSION**

Recent studies have defined unique functions for STIM2 in regulating cytosolic Ca\(^{2+}\) signals under conditions of minimal ER Ca\(^{2+}\) depletion (18,19). Upon stimulation with low concentrations of agonist that minimally deplete intracellular stores, STIM2 has been demonstrated to recruit STIM1 into ER-PM junctions to interact with ORAI1 channels and drive SOCE under conditions where STIM1 would normally be unable to oligomerize and form puncta (18). Mechanistically, the increased flexibility of the STIM2 C-terminal OASF2 region along with higher PIP2 binding affinity of the C-terminal polybasic domain are suggested to account for constitutive activation and preformed puncta of STIM2 under resting conditions (19,46). The flexibility of OASF2 triggers remodeling of the STIM1 C-terminus for interaction and gating of ORAI1 under high ER Ca\(^{2+}\) concentrations, possibly through heteromeric interactions between STIM1 and STIM2 (18). Previous studies investigated the effects of high concentrations of 2-APB on STIM and ORAI using overexpression and showed that 2-APB induces avid binding of cytosolic C-termini of both STIM1 and STIM2 to ORAI1 causing increased channel activity (24). Functional binding of expressed STIM1 C-terminus to ORAI1 also occurred in DT40 chicken B cells lacking endogenous STIM1 and STIM2, indicating that C-terminal fragments work independently of native STIMs and that STIM C-terminus is critical for 2-APB action (24). Here we show that under native levels of expression, both the N-terminus and C-terminus of STIM2 are necessary and sufficient for enhanced STIM2 sensitivity, suggesting that weak agonists or low concentrations of strong agonists use STIM2 to enhance the diversity of physiological Ca\(^{2+}\) signals. Through the use of multiple CRISPR/Cas9 cell lines with STIM1 and STIM2 knocked out individually or in combination, we conclude that 2-APB-activated store-independent Ca\(^{2+}\) entry is mediated exclusively by endogenous STIM2. This store-independent entry was significantly enhanced with STIM2 overexpression in HEK293 STIM1/2\(^{-/-}\) cells and augmented further when STIM2 was expressed in combination with ORAI1. Importantly, STIM2-dependent 2-APB-activated Ca\(^{2+}\) entry could not be rescued with overexpression of STIM1 alone or with STIM1 and ORAI1 overexpression. Only high concentrations of 2-APB (50μM) could induce a moderate level of store-independent Ca\(^{2+}\) entry in STIM1/ORAI1 overexpressing cells that was ~1/3 of that observed in cells
overexpressing STIM2/ORAI1 and stimulated by the same concentration of 2-APB. By testing full length STIM1 and STIM2 variants with mutations predicted to stabilize or disrupt CC1/CC3 interactions, we determined that increased flexibility of the STIM2 C-terminus plays a significant role in response to 2-APB mediated Ca\(^{2+}\) entry. Introduction of a stabilization mutation (R517L) shown to restrict STIM2 C-terminal flexibility and strengthen its CC1/CC3 interactions slightly reduced Ca\(^{2+}\) entry in response to 10μM 2-APB compared to STIM2. Importantly, introduction of a mutation into the C-terminus of STIM1 (L251S) making it more flexible caused preformed STIM1 puncta and induced constitutive Ca\(^{2+}\) entry, but could not support store-independent Ca\(^{2+}\) entry in response to low or high concentrations of 2-APB. These data suggest that while increased flexibility of the STIM2 C-terminus is a critical factor in the induction of store-independent Ca\(^{2+}\) entry, it is not sufficient.

In addition to differences in CC1/CC3 interactions and differences in key residues, phenylalanine (F394) in SOAR1 is a leucine in SOAR2 and glycine (G379) in SOAR1 is a glutamate in SOAR2, functional differences within the STIM2 N-terminus also distinguish its activity from STIM1. Substitution of three amino acids within the N-terminal EF-hand domain of STIM2 confer reduced Ca\(^{2+}\) binding affinity, lowering the threshold of STIM2 to induce an active conformation in response to minor reductions in ER Ca\(^{2+}\) concentrations (11). While mutation of these residues in the STIM1 EF-hand to mimic STIM2 increases the Ca\(^{2+}\) sensitivity of STIM1, conversion of the STIM2 EF-hand into the corresponding STIM1 residues failed to mimic the activation kinetics of STIM1 (11). Our chimeric STIM constructs demonstrated that substitution of the entirety of STIM1 N-terminus with that of STIM2 failed to activate store-independent Ca\(^{2+}\) entry in response to 2-APB stimulation. Similarly, replacement of the STIM2 N-terminus with that of STIM1 reduced the response to 2-APB to levels similar to STIM1. These data suggest that enhanced sensitivity of the STIM2 N-terminus in concert with the increased flexibility of its C-terminus are responsible for the unique store-independent activation of STIM2.

Conventional models of SOCE and the regulation of cytosolic Ca\(^{2+}\) signals have long placed STIM1 as the primary activator of ORAI channels in response to substantial ER depletion, while STIM2 is believed to act as a homeostatic housekeeping sensor that maintains basal Ca\(^{2+}\) concentrations (11,12). Unique structural features of STIM2 N- and C-termini dampen its interaction with ORAI channels to prevent Ca\(^{2+}\) overload as a result of its constitutive activity in ER-PM junctions (6,11). Our data utilizing multiple STIM knockout cell lines show that depending on the cell type, native STIM2 can play a significant role in sustaining SOCE when ER Ca\(^{2+}\) is maximally depleted. Knockout of STIM2 in HEK293 cells resulted in a minor reduction (~16% inhibition) of peak SOCE compared to wild type cells while STIM2 knockout in HCT116 drastically reduced SOCE (~56% inhibition) without obvious compensatory effect on STIM1 protein levels in either cell line. Further, we noted no differences in STIM1 and STIM2 protein expression between HEK293 and HCT116 cell lines. It is tempting to speculate that in HCT116 cells, STIM2 interacts with and remolds the C-terminus of STIM1 into an active conformation despite ER Ca\(^{2+}\) being significantly depleted. In support of this idea, knockout of STIM1 in HCT116 cells augmented store-independent Ca\(^{2+}\) entry in response to low and high concentrations of 2-
APB, suggesting that STIM1 and STIM2 might dynamically regulate the function of one another in a cell type-specific manner. Our results in HCT116 cells are in line with recent data in NIH 3T3 fibroblasts and αT3 cells where knockout of STIM2 resulted in a 90% reduction in SOCE (47). In addition to these cell types, a multitude of studies have identified STIM2 as the predominant STIM isoform regulating SOCE in primary neurons (48-50). STIM2-mediated Ca\textsuperscript{2+} entry plays a critical role in maintaining mushroom dendritic spine development \textit{in vitro} and \textit{in vivo} as well as regulating neuronal apoptosis in response to environmental stress (48,50-52). The use of 2-APB and newly characterized 2-APB analogues (53-55) will be critical in understanding the endogenous interactions between STIM1 and STIM2 and how these ER sensors positively and negatively regulate one another in various physiological systems. Our findings elucidating the differential effects of 2-APB on STIM1 and STIM2 offers readily exploitable approaches to understand the endogenous interactions between STIM1 and STIM2 and might lead to drugs that specifically distinguish STIM2-mediated from STIM1-driven (patho)physiological functions.

**EXPERIMENTAL PROCEDURES**

**Cell culture and transfection**

HEK293 cells were maintained in high glucose (4.5 g/L) Dulbecco’s modified Eagle’s medium (DMEM) media supplemented with 10% heat-inactivated fetal bovine serum, penicillin (100 U/mL), streptomycin (100 mg/mL) and cultured in a 5% CO\textsubscript{2} incubator at 37°C. HCT116 cells were maintained in McCoy’s 5A medium supplemented with 10% heat-inactivated fetal bovine serum, penicillin (100 U/mL), streptomycin (100 mg/mL), and cultured in the same incubation conditions. Cells were transfected with either a Nucleofector II Device (Amaxa Biosystems) and the Nucleofector Kit V (VCA-1003) following the manufacturer’s protocol or with Lipofectamine 2000 (Invitrogen). After transfection, cells were plated onto 25 mm round glass coverslips and incubated for 24 hours before use in experiments (56,57).

**DNA Constructs**

For use in confocal and Ca\textsuperscript{2+} imaging experiments, pEX-CMV-SP-YFP-STIM2(15-746) (Addgene plasmid # 18862) and pEX-SP-YFP-STIM1(23-685) (Addgene plasmid # 18857) were gifts from Tobias Meyer. pECPF-C1-Orai1 and YFP-STIM2.1 were from Donald Gill. STIM1 CC1/CC3 stabilization and disruption mutants (STIM1-L251S, STIM1-R426L) and OASF FRET sensors were gifts from Christoph Romanin. STIM2 CC1/CC3 stabilization and disruption mutants (STIM2-L342S/L507S/L514S (CCmt), STIM2-R517L) and OASF2 FRET sensors were gifts from Indu S. Ambudkar. STIM chimeras were cloned using the In-Fusion system (Takara). Briefly, STIM2N-STIM1C was generated by mixing two primers (CFP-R and S2-S1TM F) with pEX-CMV-SP-YFP-STIM1 to generate a vector lacking amino acids 21-213 of STIM1. The STIM2 insert for STIM2N-STIM1C was generated by mixing two primers (Vector-F and S1TM-S2 R) with pEX-CMV-SP-YFP-STIM2. The above vector and insert were combined according to the In-Fusion protocol to generate the final STIM1N-STIM2C. Using the same method as above to generate STIM2N-STIM1C, two primers (CFP-R and S1-S2TM F) were combined with pEX-CMV-SP-YFP-STIM2 to generate a STIM2 vector lacking aa1-208. This was then mixed with the STIM1 N-terminal fragment (aa21-213) generated by mixing primers Vector-F with S2TM-S1R with pEX-
SP-YFP-STIM1 to generate the STIM1N-STIM2C construct. Primer sequences utilized for cloning of STIM chimeras were as follows: S2-S1TM F: AAGATTTTTTATATGCTTTGTTGTTCTATC, S1TM-S2 R: GCATGAAAAAACATCTCCTCAGTGAG, S1-S2TM F: CAAGGACATCCTCACAATTATCTATAAGTG, S2TM-S1R: GTGAGGATGTCCTTGAGGTGATTATGTC, CFP-R: TCTCGGCATGGAGACAGCTGT, Vector-F: TCTCGGCATGGAGACAGCTGT.

**Generation of STIM1, STIM2, and STIM1/2 double knockout cell lines**

Guide RNAs with sequences specific for STIM1 or STIM2 were designed and inserted into the BsmBI restriction site of the lentiCRISPR v2 vector (Addgene Plasmid #52961). The guide sequences used were the following: STIM1g1: 5’-TGATGAGCTTATCCTCACCA-3’, STIM2g1: 5’-AGATGGTGGAATTGAAGTAG-3’, and STIM2g2: 5’-AGAAGAAGACAGATTTAGTC-3’. HEK293 cells were transfected with the cloned lentiCRISPR v2 vectors using a Nucleofector II Device (Amaxa Biosystems) and HCT116 were transfected using Lipofectamine 2000 (Invitrogen). 48 hours after transfection, HEK293 and HCT116 cells were cultured with their respective media containing puromycin (2 μg/mL) (Gemini Bio Products) and selected for 6 days. After puromycin selection, cells were plated at a density of one cell per well into 96-well plates to isolate individual clones. Disruption of the STIM1 or STIM2 gene was confirmed in individual clones through Sanger Sequencing, Western Blotting analysis, and functional Ca²⁺ imaging experiments.

**Single cell Ca²⁺ imaging**

HEK293 and HCT116 cells were seeded on 25 mm glass coverslips overnight, mounted in Attofluor cell chambers (Thermo Fisher Scientific), and incubated in DMEM or McCoys media containing 2 μM Fura-2AM (Molecular Probes) at 37°C for 30 minutes as described previously (56,58). Following loading with Fura-2AM, cells were washed three times and incubated for 10 minutes in a Heps-buffered saline solution containing the following components: 140 mM NaCl, 1.13 mM MgCl₂, 4.7 mM KCl, 2 mM CaCl₂, 10 mM D-glucose, and 10 mM HEPES with pH adjusted to 7.4. Chambers were mounted onto a Nikon TS100 inverted microscope equipped with a 20X Fluor objective and fluorescence images were recorded with a digital fluorescence imaging system (InCytlm2, Intracellular Imaging Inc.). Ca²⁺ imaging was also performed on a Leica DMi8 fluorescence microscope controlled by Leica Application Suite X (Leica). Fura-2AM fluorescence was measured every two seconds by alternative excitation at 340 nm (F340) and 380 nm (F380) and emission fluorescence was collected at 510 nm. Cytosolic Ca²⁺ concentrations are represented as the ratio of F340/F380. All Ca²⁺ imaging experiments were performed at room temperature and traces are shown as mean ± SEM from at least three independent experiments.

**Western blotting**

Cell lines were harvested, washed once with chilled PBS, and lysed in pre-chilled RIPA buffer (Sigma) containing Halt Protease and Phosphatase Inhibitor Cocktail (Thermo Fisher Scientific) for 10 minutes on ice. Samples were centrifuged at 14,000 g, 4°C for 10 minutes and clarified protein supernatants were quantified with the Pierce...
BCA Protein Assay Kit (Thermo Fisher Scientific). 25 μg of protein extract was loaded on a 4-12% gel NuPAGE Bis-Tris gel (Life Technologies) and transferred to a PVDF membrane using a Trans-Blot Turbo Transfer System (Bio-Rad). Membranes were blocked in Odyssey Blocking Buffer (LI-COR) for 1 hour at room temperature and incubated overnight at 4°C with one of the following primary antibodies: STIM1 antibody (1:2000; #4916, Cell Signaling Technology), STIM2 (1:1000; #4917, Cell Signaling Technology), GAPDH (1:4000; MAB374, Sigma). Membranes were washed 3 times in 0.1% TBST and incubated for 1 hour at room temperature with the following secondary antibodies: IRDye 680RD Goat Anti-Mouse (1:10,000 LI-COR), IRDye 800RD Donkey Anti-Rabbit (1:10,000 LI-COR). Following 3 washes in 0.1% TBST, membranes were imaged on an Odyssey CLx Imaging System (LI-COR). Western blot image analysis was performed in Image Studio Ver 5.2 (LI-COR) and ImageJ.

**Förster Resonance Energy Transfer (FRET) Measurements**

Measurement of OASF FRET sensors was performed on a Leica DMI 6000B inverted automated fluorescence microscope with CFP (438Ex/483Em), YFP (500Ex/542Em), and FRET (438Ex/542Em) filter cubes. Images were acquired every 20 seconds with each filter cube with a 40x oil objective and analyzed using Slidebook 6.0 software (Intelligent Imaging Innovations). Exposure times for each channel were the following: 1000 ms (CFP), 250 ms (YFP), and 1000 ms (FRET). Three channel corrected FRET was calculated with the below formula:

\[ F_C = I_{DA} - F_D/D_D \cdot I_{DD} - F_A/D_A \cdot I_{AA} \]

For this formula, \( I_{DD}, I_{AA}, \) and \( I_{DA} \) are the intensity of the background-subtracted CFP, YFP, and FRET images, respectively. For FRET analysis of OASF sensors, YFP-OASF-YFP and CFP-OASF-CFP were used for calculation of correction images and bleed through of CFP and YFP through the FRET filter. For analysis of OASF2 sensors, YFP-OASF2-YFP and CFP-OASF2-CFP were used for correction images. The E-FRET method to analyze 3-cube FRET images as initially described by Zal and Gascoigne (59) was calculated with the following formula:

\[ E_{app} = \frac{F_C}{F_C + G \cdot I_{DD}} \]

All FRET experiments were performed by transient transfection of OASF/OASF2 constructs into HEK293 STIM1/STIM2 double knockout cells.

**Statistics**

All data analyses were performed with GraphPad Prism 8 (GraphPad Software) and data is presented as mean ± SEM. An unpaired, nonparametric Mann-Whitney test was used for statistical comparisons between two groups and the Kruskal-Wallis test with Dunn’s multiple comparisons was used for comparison between multiple groups.

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**CONFLICT OF INTEREST STATEMENT**
The authors declare that they have no conflicts of interest with the contents of this article.

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**ABBREVIATIONS**
STIM, Stromal interaction molecule; SOCE, store-operated Ca\(^{2+}\) entry; SOAR, STIM-ORAI activating region; OASF, ORAI activating small fragment; ER, endoplasmic reticulum; 2-APB, 2-Aminoethoxydiphenyl borate.

**FIGURE LEGENDS**

**Figure 1:** Generation and characterization of STIM1, STIM2, and STIM1/STIM2 knockout cell lines.

(A) Western blot analysis of STIM1, STIM2, and the loading control GAPDH in HEK293 wild-type (WT) cells, STIM1\(^{-/-}\) cells, STIM2\(^{-/-}\) cells, and STIM1/2\(^{-/-}\) cells. Blots are representative of 3 independent experiments and intensity of STIM bands normalized to GAPDH are quantified in (B).

(C and D) Representative cytosolic Ca\(^{2+}\) traces in different HEK293 cells as measured by Fura-2 in response to store depletion with 2μM thapsigargin (Tg) in nominally Ca\(^{2+}\) free external solution and subsequent re-introduction of 2mM Ca\(^{2+}\) to the extracellular medium. All traces are represented as mean (dark symbols) ± SEM (light background shading). (D) Peak SOCE calculated as the baseline-subtracted maximal values of Fura2 ratio units. Each scatter plot displays distribution of peak SOCE values for n=100 cells from a total of 3 independent experiments.

(E) Western blot analysis of STIM1, STIM2, and the loading control GAPDH in WT HCT116 cells, STIM1\(^{-/-}\) cells, and STIM2\(^{-/-}\) cells. Blots are representative of 3 independent experiments and densitometry of STIM bands normalized to GAPDH are quantified in (F).

(G and H) Representative Ca\(^{2+}\) imaging traces in different HCT116 cells using the same protocol as in (C). (H) Peak SOCE calculated as in (D). Each scatter plot displays distribution of peak SOCE values for n=100 cells from a total of 3 independent experiments. ****p<0.0001, Kruskal-Wallis test with Dunn’s multiple comparisons to WT parental line.

**Figure 2:** 2-APB activates store-independent Ca\(^{2+}\) entry exclusively through endogenous STIM2.

(A) Western blot analysis of STIM1, STIM2, and the loading control GAPDH in additional HEK293 STIM2\(^{-/-}\) clones.

(B-E) Ca\(^{2+}\) entry was measured using Fura2 upon addition of 10 μM 2-APB in the presence of 2mM Ca\(^{2+}\) in WT HEK293 and each STIM CRISPR cell line. Ca\(^{2+}\) imaging traces are average data from n= 145-154 individual cells per condition. (C) Scatter plots show mean ± SEM of baseline-subtracted maximal values of Fura2 ratio units. (D) Ca\(^{2+}\) entry measured upon addition of 50 μM 2-APB. Ca\(^{2+}\) imaging traces are average data from n= 131-150 individual cells per condition. (E) Scatter plots show mean ± SEM of baseline-subtracted maximal value of Fura2 ratio units.

(F) Western blot analysis of STIM1, STIM2, and the loading control GAPDH in additional HCT116 STIM1\(^{-/-}\) and STIM2\(^{-/-}\) clones. For STIM2\(^{-/-}\), clones were generated with 2 distinct guide RNAs (see methods), with 2 independent clones per guide RNA.
(G-J) Same experimental conditions as in (B-E) except that WT HCT116 and its STIM CRISPR cell line variants were used. (G) 10 μM 2-APB was used for stimulation and Ca\(^{2+}\) imaging traces are average data from n= 119-125 cells per condition. (H) Scatter plots show mean ± SEM of baseline-subtracted maximal values of Fura2 ratio units. (I) 50 μM 2-APB was used for stimulation and Ca\(^{2+}\) imaging traces are average data from n= 114-120 individual cells per condition. (J) Scatter plots show mean ± SEM of baseline-subtracted maximal values of Fura2 ratio units. All traces are averaged from 3 independent experiments. ****p<0.0001, ***p<0.001, **p<0.01, *p<0.05, Kruskal-Wallis test with Dunn’s multiple comparisons to WT parental line.

**Figure 3: STIM2 is sufficient to rescue store-independent Ca\(^{2+}\) entry.**

(A) Representative confocal images of HEK293 STIM1/2\(^{-/-}\) cells expressing either YFP-STIM2 or YFP-STIM1 at baseline and after stimulation with 10 μM 2-APB for 5 minutes. (B) Quantification of average STIM2 puncta size at baseline and after 10 μM 2-APB stimulation. (C) HEK293 STIM1/2\(^{-/-}\) cells overexpressing YFP-STIM2 (STIM2) or YFP-STIM1 (STIM1) and stimulated with 10 μM 2-APB in the presence of 2mM Ca\(^{2+}\). Dotted traces represent 10 μM 2-APB-induced Ca\(^{2+}\) entry in HEK293 STIM1\(^{-/-}\) cells (STIM1\(^{-/-}\)) or HEK293 STIM2\(^{-/-}\) cells (STIM2\(^{-/-}\)). Ca\(^{2+}\) imaging traces are average data from n= 100-118 transfected cells per condition from 3 independent experiments. (D) Scatter plots show mean ± SEM of baseline-subtracted maximal values of Fura2 ratio units. (E) Same experimental conditions as in (C) but with 50 μM 2-APB stimulation. Ca\(^{2+}\) imaging traces are average data from n= 99-125 transfected cells per condition from 3 independent experiments. (F) Scatter plots show mean ± SEM of baseline-subtracted maximal values of Fura2 ratio units. ****p<0.0001, *p<0.05, nonparametric Mann-Whitney test.

**Figure 4: ORAI1 overexpression enhances STIM2, but not STIM1, mediated store-independent Ca\(^{2+}\) entry.**

(A) HEK293 STIM1/2\(^{-/-}\) cells overexpressing ORAI1-CFP and YFP-STIM2 (STIM2 + ORAI1) or ORAI1-CFP and YFP-STIM1 (STIM1 + ORAI1) and stimulated with 10 μM 2-APB in the presence of 2mM Ca\(^{2+}\). Dotted traces represent addition of 10 μM 2-APB to cells overexpressing either STIM1 alone (STIM1) or STIM2 alone (STIM2). Ca\(^{2+}\) imaging traces are average data from n= 165-176 transfected cells per condition from 3 independent experiments. (B) Scatter plots show mean ± SEM of baseline-subtracted maximal values of Fura2 ratio units. (E) Same experimental conditions as in (A) but with 50 μM 2-APB stimulation. Ca\(^{2+}\) imaging traces are average data from n= 151-172 transfected cells per condition from 3 independent experiments. (F) Scatter plots show mean ± SEM of baseline-subtracted maximal values of Fura2 ratio units. ****p<0.0001, **p<0.01, *p<0.05, nonparametric Mann-Whitney test. (C, D and G) (C) Representative confocal images of HEK293 STIM1/2\(^{-/-}\) cells overexpressing ORAI1-CFP and YFP-STIM2 at baseline and after stimulation with 10 μM 2-APB for 5 minutes. (D) Quantification of average STIM2 puncta size in STIM2 + ORAI1 expressing cells (from C) at baseline and after 10 μM 2-APB stimulation, nonparametric Mann-Whitney test (G) Representative confocal images of HEK293 STIM1/2\(^{-/-}\) cells overexpressing ORAI1-CFP and YFP-STIM1 at baseline and after stimulation with 10 μM 2-APB for 5 minutes.
Figure 5: Low concentrations of 2-APB potentiate SOCE through STIM2.

(A) Measurement of carbachol-induced SOCE in WT HEK293 and STIM CRISPR cell lines. Cells were stimulated with 100 μM carbachol in a nominally Ca\(^{2+}\) free solution followed by re-introduction of 2 mM Ca\(^{2+}\) to the extracellular medium. 10 μM and 50 μM 2-APB were added in the presence of extracellular Ca\(^{2+}\) at the indicated times. Ca\(^{2+}\) imaging traces are average data from n= 101-140 cells per condition from 3 independent experiments.

(B) Same experimental conditions as in (A) but with 150 μM ATP in a nominally Ca\(^{2+}\) free solution, followed by re-introduction of 10mM Ca\(^{2+}\) (instead of 2 mM) to the extracellular medium. 10 mM was used to enhance the driving force and reveal responses to ATP, which were undetectable in 2 mM external Ca\(^{2+}\). Ca\(^{2+}\) imaging traces are average data from n= 116-135 cells per condition from 3 independent experiments.

(C) Measurement of ATP-induced SOCE in WT HCT116 and STIM CRISPR cell lines. Cells were stimulated with 300 μM ATP in a nominally Ca\(^{2+}\) free solution, followed by re-introduction of 10mM Ca\(^{2+}\) to the extracellular medium. The use of 150 μM ATP in HCT116 cells generated small and inconsistent SOCE. 10 μM and 50 μM 2-APB were added in the presence of extracellular Ca\(^{2+}\) at the indicated times. Ca\(^{2+}\) imaging traces are average data from n= 60-100 cells per condition from 3 independent experiments.

Figure 6: Differential CC1/CC3 interactions contribute to increased flexibility of the STIM2 C-terminus

(A) Representative confocal images of HEK293 STIM1/2\(^{-/-}\) cells expressing either YFP-STIM2 or YFP-STIM2.1.

(B) HEK293 STIM1/2\(^{-/-}\) cells overexpressing YFP-STIM2 or YFP-STIM2.1 and stimulated with 10 μM 2-APB in the presence of 2 mM Ca\(^{2+}\). Ca\(^{2+}\) imaging traces are average data from n= 122-159 transfected cells per condition from 3 independent experiments. Scatter plots show mean ± SEM of baseline-subtracted maximal values of Fura2 ratio units. ****p<0.0001, nonparametric Mann-Whitney test.

(C) Diagram of STIM1 and STIM2 structure showing CC1 and CC3 stabilization/disruption mutations.

(D) OASF FRET sensors for STIM1, YFP-OASF1-CFP (Y-OASF1-C) and for STIM2, YFP-OASF2-CFP (Y-OASF2-C) were expressed in HEK293 STIM1/2\(^{-/-}\) cells and baseline E-FRET signal was measured from Y-OASF1-C and Y-OASF2-C. ****p<0.0001, nonparametric Mann-Whitney test.

(E) Scatter plots demonstrating the distribution of intramolecular FRET signal from STIM1/2\(^{-/-}\) cells overexpressing YFP-OASF1-CFP or YFP-OASF2-CFP (WT) or their stabilization and disruption mutants. ****p<0.0001, ***p<0.001, mutants were compared to WT constructs using nonparametric Mann-Whitney test.

(F and G) Intramolecular FRET in HEK293 STIM1/2\(^{-/-}\) cells expressing either OASF2 (F) or OASF1 (G) WT and mutant variants under basal conditions and after addition of 10 μM 2-APB.

(H, I) Zoomed-in and normalized FRET data from (F, G) respectively.

Figure 7: Flexibility of the STIM2 C-terminus contributes to 2-APB Selectivity
(A) Representative confocal images of HEK293 STIM1/2−/− cells expressing YFP-STIM2 or YFP-tagged mutants, including the relaxed CCmt-STIM2 (L342S/L507S/L514S) mutant, or the stabilized R517L-STIM2 mutant at baseline and after stimulation with 10 μM 2-APB.

(B) Quantification of average puncta size in STIM2, CCmt-STIM2 and R517L-STIM2 mutants at baseline and after 10 μM 2-APB stimulation.

(C and D) (C) HEK293 STIM1/2−/− cells overexpressing YFP-tagged versions of either STIM2, CCmt-STIM2, or R517L-STIM2 and stimulated with 10 μM 2-APB in the presence of 2 mM Ca2+. Ca2+ imaging traces are average data from n= 113-120 transfected cells per condition from 3 independent experiments. (D) Scatter plots show mean ± SEM of baseline-subtracted maximal values of Fura2 ratio units. ****p<0.0001, Kruskal-Wallis test with Dunn’s multiple comparisons of STIM2 mutants to wildtype STIM2.

(E and F) (E) Same as in (C) but stimulation with 50 μM 2-APB in the presence of 2 mM Ca2+. Ca2+ imaging traces are average data from n= 119-127 transfected cells per condition from 3 independent experiments. (F) Scatter plots show mean ± SEM of baseline-subtracted maximal values of Fura2 ratio units. ****p<0.0001, Kruskal-Wallis test with Dunn’s multiple comparisons of STIM2 mutants to wildtype STIM2.

Figure 8: Increased flexibility of the STIM1 C-terminus is not sufficient to recapitulate store-independent Ca2+ entry.

(A) Representative confocal images of HEK293 STIM1/2−/− cells expressing YFP-tagged versions of either STIM1, the relaxed L251S-STIM1 mutant, or the stabilized R426L-STIM1 mutant at baseline and after stimulation with 10 μM 2-APB.

(B) Quantification of basal Ca2+ levels in cells expressing STIM1, L251S-STIM1, or R426L-STIM1 mutants prior to 2-APB stimulation.

(C) HEK293 STIM1/2−/− cells overexpressing YFP-tagged versions of either STIM1, L251S-STIM1, or R426L-STIM1 and stimulated with 10 μM 2-APB in the presence of 2 mM Ca2+. Ca2+ imaging traces are average data from n= 95-100 transfected cells per condition from 3 independent experiments. (D) Scatter plots show mean ± SEM of baseline-subtracted maximal values of Fura2 ratio units.

(E) Same experimental conditions as in (C) but with 50 μM 2-APB stimulation. Ca2+ imaging traces are average data from n= 50-99 transfected cells per condition from 3 independent experiments. (F) Scatter plots show mean ± SEM of baseline-subtracted maximal values of Fura2 ratio units. ****p<0.0001, Kruskal-Wallis test with Dunn’s multiple comparisons of STIM1 mutants to wildtype STIM1.

(G) Quantification of basal Ca2+ levels in cells expressing STIM1, STIM2, or STIM1+STIM2 prior to 2-APB stimulation.

(H, I) (H) Same as in (C) but with overexpression of YFP-tagged versions of either STIM2, STIM1, or STIM1+STIM2 and stimulated with 10 μM 2-APB in the presence of 2mM Ca2+. (I) Scatter plots show mean ± SEM of baseline-subtracted maximal values of Fura2 ratio units.

(J, K) (J) Same experimental conditions as in (E) but with overexpression of YFP-tagged versions of either STIM2, STIM1, or STIM1+STIM2 and stimulated with 50 μM 2-APB in the presence of 2 mM Ca2+. (K) Scatter plots show mean ± SEM of baseline-subtracted maximal values of Fura2 ratio units.

Figure 9: Coordination between STIM2 N- and C-terminus underlies 2-APB sensitivity
(A) Schematic of YFP-STIM1, YFP-STIM2, and STIM chimeric constructs used for Ca\textsuperscript{2+} imaging experiments.
(B) Representative confocal images of several HEK293 STIM1/2\textsuperscript{-/-} cells expressing either chimeric YFP-S1N-S2C (top) or YFP-S2N-S1C (bottom) constructs.
(C) Measurement of thapsigargin-activated SOCE in control HEK293 STIM1/2\textsuperscript{-/-} cells or cells overexpressing YFP-S1N-S2C or YFP-S2N-S1C. Cells were stimulated with 2 \textmu M thapsigargin (Tg) in a nominally Ca\textsuperscript{2+} free solution, followed by re-introduction of 2 mM Ca\textsuperscript{2+} to the extracellular medium. 50 \textmu M 2-APB was added in the presence of extracellular Ca\textsuperscript{2+} at the indicated time. Ca\textsuperscript{2+} imaging traces are average data from n= 112-151 cells per condition from 3 independent experiments. (D) Peak SOCE was calculated as baseline-subtracted maximal values of Fura2 ratio units.
(E) Quantification of the slope of inhibition Ca\textsuperscript{2+} entry by 50 \textmu M 2-APB in cells expressing YFP-S1N-S2C or YFP-S2N-S1C chimeras.
(F) Quantification of basal Ca\textsuperscript{2+} levels in cells expressing YFP-S1N-S2C or YFP-S2N-S1C.
(G) Constitutive activity of HEK293 STIM1/2\textsuperscript{-/-} cells overexpressing YFP-tagged versions of either S1N-S2C or S2N-S1C chimeras. Cells were maintained in a nominally Ca\textsuperscript{2+} free solution for 5 min followed by restoration of 2 mM Ca\textsuperscript{2+} to extracellular milieu to determine basal unregulated activity. Data are average from n= 164-166 cells per condition from 3 independent experiments.
(H) HEK293 STIM1/2\textsuperscript{-/-} cells overexpressing YFP-tagged versions of either STIM1, STIM2, S1N-S2C, or S2N-S1C and stimulated with 10 \textmu M 2-APB in the presence of 2 mM Ca\textsuperscript{2+}. Ca\textsuperscript{2+} imaging traces are average data from n= 122-180 transfected cells per condition from 3 independent experiments. (I) Scatter plots show mean ± SEM of baseline-subtracted maximal values of Fura2 ratio units.
(J) Similar protocol to (H) but stimulation with 50 \textmu M 2-APB. Ca\textsuperscript{2+} imaging traces are average data from n= 143-191 transfected cells per condition from 3 independent experiments. (K) Scatter plots show mean ± SEM of baseline-subtracted maximal values of Fura2 ratio units.
**HEK293 cells**

A

100 kDa
75 kDa
100 kDa
37 kDa

HEK293
STIM1
STIM2
GAPDH

B

[Graph showing F340/F380 ratio over time for HEK293 cells with different STIM and 2-APB conditions.]

C

[Graph showing F340/F380 ratio over time for HEK293 cells with different STIM and 2-APB conditions.]

**HCT116 cells**

F

100 kDa
75 kDa
100 kDa
37 kDa

HCT116
STIM1
STIM2
GAPDH

G

[Graph showing F340/F380 ratio over time for HCT116 cells with different STIM and 2-APB conditions.]

H

[Graph showing F340/F380 ratio over time for HCT116 cells with different STIM and 2-APB conditions.]

**FIG. 2**
YFP-STIM1 and YFP-STIM2 expressed in HEK293 STIM1/2−/− cells

**FIG. 3**

(A) Baseline 10μM 2-APB

(B) STIM1 STIM2 baseline 10μM 2-APB

(C) STIM2 STIM1 STIM1−/− STIM2−/−

(D) F/F0 STIM2 STIM1 STIM1−/− STIM2−/−

(E) F/F0 50μM 2-APB STIM2 STIM1 STIM1−/− STIM2−/−

(F) ns **** ns **** ns ns
YFP-STIM1 and YFP-STIM2 co-expressed with ORAI1-CFP in HEK293 STIM1/2−/− cells

**FIG. 4**

A

| 2-APB | Time (min) |
|-------|-------------|
| 0.2   | 0           |
| 0.4   | 2           |
| 0.6   | 4           |
| 0.8   | 6           |
| 1.0   | 8           |
| 1.2   | 10          |

B

| 2-APB | Time (min) |
|-------|-------------|
| 0.2   | 0           |
| 0.4   | 2           |
| 0.6   | 4           |
| 0.8   | 6           |
| 1.0   | 8           |
| 1.2   | 10          |

C

| 2-APB | Time (min) |
|-------|-------------|
| 0.2   | 0           |
| 0.4   | 2           |
| 0.6   | 4           |
| 0.8   | 6           |
| 1.0   | 8           |
| 1.2   | 10          |

D

**Puncta Size (μm²)**

E

**F340/F380**

F

**F340/F380**

G

**F340/F380**
**HEK293 cells**

(A) F340/F380 vs. Time (min)

- HEK293
- STIM1<sup>−/−</sup>
- STIM2<sup>−/−</sup>
- STIM1/2<sup>−/−</sup>

- 100 μM CCh
- 2mM Ca<sup>2+</sup>
- 10μM 2-APB
- 50μM 2-APB

(B) F340/F380 vs. Time (min)

- HEK293
- STIM1<sup>−/−</sup>
- STIM2<sup>−/−</sup>
- STIM1/2<sup>−/−</sup>

- 150 μM ATP
- 10μM Ca<sup>2+</sup>
- 10μM 2-APB
- 50μM 2-APB

**HCT116 cells**

(C) F340/F380 vs. Time (min)

- HCT116
- STIM1<sup>−/−</sup>
- STIM2<sup>−/−</sup>

- 300 μM ATP
- 10μM Ca<sup>2+</sup>
- 5μM 2-APB
- 50μM 2-APB
YFP-STIM2 and YFP-STIM2.1 Constructs expressed in HEK293 STIM1/2−/− cells

OASF1 and OASF2 FRET Sensors expressed in HEK293 STIM1/2−/− cells

FIG. 6

E-FRET

OASF1 and OASF2 FRET Sensors expressed in HEK293 STIM1/2−/− cells

YFP-STIM2 and YFP-STIM2.1 Constructs expressed in HEK293 STIM1/2−/− cells

A

B

C

D

E

F

G

H

I

OASF1 (233-474)

SOAR1 (344-442)

OASF2 (324-565)

SOAR2 (435-534)

STIM1

STIM2

YFP-STIM2

YFP-STIM2.1

10μM 2-APB

10μM 2-APB

10μM 2-APB
FIG. 7

YFP-STIM2 mutants expressed in HEK293 STIM1/2−/− cells

A  Baseline  10μM 2-APB

STIM2

CCmt-STIM2

R517L-STIM2

B  C  D

F340/F380

Time (min)

0 2 4 6 8 10

0 0.2 0.4 0.6 0.8

STIM2  CCmt-STIM2  R517L-STIM2

E  F

Puncta Size (μm²)

0 0.5 1.0 1.5 2.0 2.5 3.0

R517 Baseline  10μM 2-APB

****

ns

****

ns

****

ns

0.0 0.1 0.2 0.3 0.4 0.5 0.6 0.7 0.8 0.9 1.0

F340/F380

Time (min)

0 2 4 6 8 10

0 0.2 0.4 0.6 0.8 1.0

STIM2  CCmt-STIM2  R517L-STIM2
YFP-STIM1 mutants expressed in HEK293 STIM1/2-/- cells

FIG. 8

A  Baseline  10μM 2-APB

B  STIM1 Mutants Basal Ca^{2+}

C  10μM 2-APB

D  STIM1 L251S STIM1 R426L STIM1

E  50μM 2-APB

F  STIM2 STIM1+ STIM2

G  Basal Ca^{2+}

H  STIM2 STIM1+ STIM2

I  STIM1

J  STIM2 STIM1+ STIM2

K  STIM1

30
YFP-STIM and YFP-STIM chimeric constructs expressed in HEK293 STIM1/2−/− cells

FIG. 9

A

STIM1

[Diagram showing STIM1 constructs]

STIM2

[Diagram showing STIM2 constructs]

S1N-S2C

[Diagram showing S1N-S2C constructs]

S2N-S1C

[Diagram showing S2N-S1C constructs]

B

[Images of YFP-STIM constructs in HEK293 cells]

C

F340/F380

0.2

0.4

0.6

0.8

1.0

Time (min)

D

F-F0

0

0.1

0.2

0.3

0.4

0.5

S1N-S2C

S2N-S1C

STIM1

STIM2

F-F0

0

S1N-S2C

S2N-S1C

STIM1

STIM2

F340/F380

Slope F340/F380 (15-30 min)

E

****

****

****

****

Basal Ca2+

F

G

H

I

J

K

[Graphs and charts showing F340/F380 ratios and F-F0 changes with 10 μM 2-APB]

[Graphs and charts showing F340/F380 ratios and F-F0 changes with 50 μM Ca2+]

[Graphs and charts showing F340/F380 ratios and F-F0 changes with 2 μM Ca2+]

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Crosstalk between N-terminal and C-terminal domains in stromal interaction molecule 2 (STIM2) determines enhanced STIM2 sensitivity
Scott M. Emrich, Ryan E. Yoast, Ping Xin, Xuexin Zhang, Trayambak Pathak, Robert Nwokonko, Maxime Gueguinou, Krishna P. Subedi, Yandong Zhou, Indu S. Ambudkar, Nadine Hempel, Khaled Machaca, Donald L. Gill and Mohamed Trebak

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