Interaction between store-operated and arachidonate-activated calcium entry

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Abstract

A ubiquitous pathway for cellular Ca\textsuperscript{2+} influx involves ‘store-operated channels’ that respond to depletion of intracellular Ca\textsuperscript{2+} pools via an as yet unknown mechanism. Due to its wide-spread expression, store-operated Ca\textsuperscript{2+} entry (SOCE) has been considered a principal route for Ca\textsuperscript{2+} influx. However, recent evidence has suggested that alternative pathways, activated for example by lipid metabolites, are responsible for physiological Ca\textsuperscript{2+} influx. It is not clear if these messenger-activated Ca\textsuperscript{2+} entry routes exist in all cells and what interaction they have with SOCE. In the present study we demonstrate that HEK-293 cells and Saos-2 cells express an arachidonic acid (AA)-activated Ca\textsuperscript{2+} influx pathway that is distinct from SOCE on the basis of sensitivity to pharmacological blockers and depletion of cellular cholesterol. We examined the functional interaction between SOCE and the arachidonate-triggered Ca\textsuperscript{2+} influx (denoted non-SOCE). Both Ca\textsuperscript{2+} entry routes could underlie substantial long-lasting Ca\textsuperscript{2+} elevations. However, the two pathways could not operate simultaneously. With cells that had an on-going SOCE response, addition of arachidonate gave two profound effects. Firstly, it rapidly inhibited SOCE. Secondly, the mode of Ca\textsuperscript{2+} influx switched to the non-SOCE mechanism. Addition of arachidonate to naïve cells resulted in rapid activation of the non-SOCE pathway. However, this Ca\textsuperscript{2+} entry route was very slowly engaged if the SOCE pathway was already operative. These data indicate that the SOCE and arachidonate-activated non-SOCE pathways interact in an inhibitory manner. We probed the plausible mechanisms by which these two pathways may communicate.

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1. Introduction

Calcium (Ca\textsuperscript{2+}) is a ubiquitous intracellular messenger that controls a diverse range of cellular processes [1]. Cells have access to two sources of Ca\textsuperscript{2+}—finite stores located in intracellular organelles and a more substantial pool of extracellular Ca\textsuperscript{2+}. In many cells types, release of Ca\textsuperscript{2+} from intracellular stores leads to the activation of a Ca\textsuperscript{2+} influx pathway denoted ‘store-operated Ca\textsuperscript{2+} entry’ (SOCE) [2]. This well-characterised Ca\textsuperscript{2+} entry mechanism is responsible for replenishing depleted intracellular stores and prolonging cellular Ca\textsuperscript{2+} signals [3]. Although the molecular identity of the channels responsible for SOCE and their precise mechanism of activation is unclear, plausible candidates are members of the canonical transient receptor potential (TRPC) family [3,4] with the involvement of stromal interaction molecule (STIM) [5]. This hypothesis is supported by several studies, which have demonstrated that ablation or heterologous expression of various TRPC isoforms/STIM can alter SOCE responses (e.g. [5–8]).

SOCE has been demonstrated using many different cell types during physiological stimulation or following deplection of intracellular Ca\textsuperscript{2+} stores using SERCA inhibitors such as thapsigargin [2,3,9,10]. The wide-spread expression of SOCE highlights its importance in replenishing intracellular Ca\textsuperscript{2+} pools after they have been discharged. However, it is becoming increasingly apparent that stimulation of cells...
with hormones that invoke the production of Ca\(^{2+}\)-releasing messengers not only activates SOCE, but can also promote additional Ca\(^{2+}\) entry pathways [11,12].

The intracellular messenger inositol 1,4,5-trisphosphate (InsP\(_3\)) is produced following hydrolysis of the minor membrane phospholipid phosphatidylinositol 4,5-bisphosphate (PIP\(_2\)). It is firmly established that InsP\(_3\) can release Ca\(^{2+}\) from intracellular stores and thereby trigger SOCE. InsP\(_3\) may also activate Ca\(^{2+}\) entry by binding to InsP\(_3\)Rs located in the plasma membrane [13]. The other product of PIP\(_2\) hydrolysis is diacylglycerol (DAG). Unlike the water soluble InsP\(_3\), DAG stays in the plane of the plasma membrane where it can activate protein kinase C (PKC), or be metabolised further. Both PKC and DAG (or membrane permissive analogues) have been demonstrated to cause Ca\(^{2+}\) influx distinct from SOCE in some cell types [14,15]. Furthermore, other messengers resulting from DAG metabolism, including AA and leukotrienes, activate non-store-operated Ca\(^{2+}\) influx [16–19].

Probably the best characterised non-SOCE activator is AA. Studies from several labs, including those of Shuttleworth [17,20], Taylor [16,21] and Putney [22], in addition to our own work [19], have demonstrated pathways for AA-stimulated Ca\(^{2+}\) entry that are clearly distinct from SOCE. As with SOCE, the molecular targets of these messengers are not established, though TRPC isoforms have again been implicated. Although there are an increasing number of reports demonstrating significant Ca\(^{2+}\) influx via non-SOCE, it is presently unclear whether this is a general mechanism for Ca\(^{2+}\) entry. Furthermore, it is not fully established how cells utilise these distinct Ca\(^{2+}\) entry pathways during physiological signalling or how they interact.

In the present study, we sought to characterise the SOCE and arachidonate-activated non-SOCE pathways in HEK-293 and Saos-2 cells. Our data indicate that these Ca\(^{2+}\) influx mechanisms are pharmacologically distinct, suggesting that they are separate entities. However, they functionally interact with each other. Essentially, SOCE and non-SOCE cannot operate simultaneously due to mutual antagonism, as proposed by other groups [16,17,22]. In contrast to other studies [17,21], however, we observed that the reciprocal inhibition of the two Ca\(^{2+}\) entry pathways was not due to either elevated Ca\(^{2+}\) or nitric oxide (NO). Our data are consistent with a scheme in which arachidonate directly inhibits SOCE, and prior activation of SOCE substantially reduces the ability of the non-SOCE pathway to be engaged. Whenever AA is present, non-SOCE appears to be the dominant Ca\(^{2+}\) entry pathway.

2. Materials and methods

2.1. Cell culture

HEK-293 and Saos-2 cells were grown at 37°C in Dulbecco’s modified Eagle’s medium supplemented with 10% heat-inactivated fetal bovine serum, 100 U/ml penicillin, 100 µg/ml streptomycin and 2 mM l-glutamine in a humidified 95% air, 5% CO\(_2\) incubator. All experimental procedures were carried out at room temperature (20–22°C). Prior to imaging, the culture medium was replaced with an extracellular medium (EM) containing (mM): NaCl, 121; KCl, 5.4; MgCl\(_2\), 0.8; CaCl\(_2\), 1.8; NaHCO\(_3\), 6; D-glucose, 5.5; Hepes, 25; pH 7.3.

2.2. Video imaging

Measurement of cytosolic Ca\(^{2+}\) in HEK-293 and Saos-2 cells was performed by monitoring fura-2 fluorescence of cells adhered to glass coverslips using a PerkinElmer Life Sciences Ltd. (Seer Green, Beaconsfield, UK) imaging system. Fura-2 was loaded into the cells by incubation with 2 µM fura-2 acetoxyethyl ester (30 min incubation followed by a 30-min period for de-esterification). A single glass coverslip with adherent cells was mounted on the stage of a Nikon Diaphot TE300 inverted epi-fluorescence microscope coupled to a xenon arc lamp. Fluorescence images were obtained using alternate excitation at 340 and 380 nm. The excitation light was controlled by a Sutter filter changer (340HT15 and 380HT15; Sutter Industries). Emitted light was filtered at 510 nm and collected by a cooled Astrocam digital camera. The acquired images were stored and subsequently processed off-line with Ultraview software (PerkinElmer Life Sciences Ltd., UK).

2.3. Manganese quench imaging

Manganese (Mn\(^{2+}\)) entry was measured indirectly by recording the quench of fura-2 fluorescence when excited at 360 nm. To minimize the effect of contaminating Ca\(^{2+}\), Mn\(^{2+}\)-containing hapes buffer was supplemented with 1 mM EGTA and the Mn\(^{2+}\) concentration was adjusted to avoid chelation of this ion. This was achieved using the web based programme MaxChelator (http://www.stanford.edu/~cpatton/maxc.html). 2 mM Mn\(^{2+}\) in the presence of 1 mM EGTA equated to a free Mn\(^{2+}\) of ~1 mM.

2.4. Preparation and storage of AA

Porcine liver AA (Na\(^+\) salt; Calbiochem) was dissolved in MilliQ water as a concentrated stock solution before being dispensed in aliquots then frozen and stored at −20°C in a light-resistant container. Prior to use, an aliquot of AA was diluted to the required concentration by addition of EM containing 0.1% DMSO (to aid membrane permeability). All samples of AA were kept in the dark and on ice before use to suppress oxidation.

2.5. Materials

Thapsigargin, 2-aminoethoxydiphenyl borate (2-APB), gadolinium, SNP, l-NAME and calmidazolium were purchased from Sigma. LOE-908 ((R,S)-(3,4-dihydro-6,7-dime-
thoxy-isochinolin-1yl)-2-phenyl-N,N-di[2-(2,3,4-trimethoxy phenyl) ethyl] acetamid mesylate) was generously given by Boehringer Ingelheim (Biberach an der Riss, Germany). Fura-2 was obtained from Invitrogen Molecular Probes. DAF-FM and NOC-18 were purchased from Calbiochem.

2.6. Statistics

Quantitative data are expressed as mean ± S.E.M. For Ca²⁺ imaging experiments, representative traces are shown. These traces depict the most consistent pattern of response observed in multiple cells (>30) imaged in at least three independent experiments. All experiments were repeated on different days. Statistical significance was determined by Student’s t-test (calculated using GraphPad Prism, San Diego, USA).

3. Results

3.1. Thapsigargin and AA activate pharmacologically distinct Ca²⁺ influx pathways

Stimulation of HEK-293 or Saos-2 cells with 2 μM thapsigargin in Ca²⁺-free medium evoked a rapid transient mobilization of Ca²⁺ from intracellular stores (Fig. 1A). Subsequent superfusion of cells with Ca²⁺-containing medium led to a sustained cytosolic Ca²⁺ increase, indicative of SOCE (Fig. 1A). A similar pattern of response was observed if AA (30 μM) was used instead of thapsigargin (Fig. 1B).

Although the patterns of Ca²⁺ responses evoked by thapsigargin and AA were alike in both HEK-293 and Saos-2 cells, the Ca²⁺ signals were pharmacologically clearly distinct. In both cell types, the Ca²⁺ entry stimulated by thapsigargin was substantially inhibited by 100 μM 2-APB or 1 μM Gd³⁺ (Fig. 1C and D), whilst the non-selective cation channel inhibitor LOE-908 (100 μM) produced only a modest reduction in SOCE. With AA-stimulated Ca²⁺ entry, the sensitivity to these blockers was reversed; LOE-908 evoked a profound inhibition, whilst 2-APB and Gd³⁺ were relatively ineffective (Fig. 1C and D). These data indicate that AA activated a Ca²⁺ entry pathway that was pharmacologically distinct from SOCE. The responses to all concentrations of AA used in this study were reversible upon washout of the fatty acid. There was no significant loss of fura-2 in cells incubated with 5–30 μM AA, suggesting that plasma membranes were not severely compromised by addition of the lipid.

To examine whether SOCE and AA-mediated Ca²⁺ entry could be simultaneously activated, cells were stimulated first with thapsigargin and then subsequently with AA. As
Fig. 2. AA inhibits thapsigargin-mediated SOCE and activates a pharmacologically distinct non-SOCE. HEK-293 cells (Ai) and Saos-2 cells (Bi) were treated with thapsigargin in the presence of extracellular Ca$^{2+}$. This resulted in Ca$^{2+}$ release followed by a sustained plateau of intracellular Ca$^{2+}$ due to SOCE. Subsequent addition of AA evoked a rapid inhibition of SOCE and the gradual activation of an alternative form of Ca$^{2+}$ entry. The AA-induced Ca$^{2+}$ entry was sensitive to LOE-908 in both HEK-293 (Aii) and Saos-2 (Bii) cells. (C) Thapsigargin-mediated Mn$^{2+}$ entry was monitored in HEK-293 cells. The traces depict representative responses of single cells treated with thapsigargin alone (black trace) or thapsigargin + 30 μM AA (grey trace). Quantitative analysis of the initial rate of fura-2 quench from experiments such as that illustrated in (C) is depicted in (D). The data represent the mean ± S.E.M. of at least 40 cells analyzed in three independent experiments.

depicted in Fig. 2Ai and Bi for HEK-293 and Saos-2 cells, respectively, both agents evoked a similar pattern of response. Thapsigargin caused a rapid initial increase in cytosolic Ca$^{2+}$ (due to release from stores), followed by a prolonged phase of Ca$^{2+}$ entry. With the Saos-2 cells, but not the HEK-293 cells, the amplitude of the Ca$^{2+}$ entry signal was substantially greater than the peak of the Ca$^{2+}$ release. For both cell types, the addition of AA immediately reduced cytosolic Ca$^{2+}$, consistent with a rapid inhibition of SOCE. Following this decline in Ca$^{2+}$, there was a gradual re-emergence of Ca$^{2+}$ influx, which could be inhibited by LOE-908 (Fig. 2Aii and Bii), demonstrating the activation of a distinct Ca$^{2+}$ influx pathway and not simply a transient inhibition of SOCE. These data suggest that the Ca$^{2+}$ entry pathways activated by thapsigargin and AA cannot be simultaneously activated; addition of AA to cells treated with thapsigargin rapidly curtails SOCE and slowly switches the pathway of entry to an LOE-908-sensitive route.

To examine the unidirectional rate of cation entry more precisely, we supplemented the extracellular medium with Mn$^{2+}$ and monitored the quench of fura-2 as this ion entered cells. When HEK cells were stimulated with thapsigargin for 3 min, the subsequent addition of Mn$^{2+}$ caused an almost instantaneous loss of fura-2 fluorescence (black trace in Fig. 2C). If AA was added simultaneously with thapsigargin, the initial rate of fura-2 quench was dramatically slowed (Fig. 2D and grey trace in Fig. 2C). The rate of fura-2 quench progressively increased after several minutes of incu-
bation with thapsigargin and AA. These observations are consistent with the rapid inhibition of SOCE by AA followed by the slow development of an alternative cation entry pathway.

A further distinction between the SOCE and AA-mediated Ca\(^{2+}\) entry pathways was obvious following treatment of HEK-293 cells with methyl-\(\beta\)-cyclodextrin (M\(\beta\)CD), an agent commonly used to extract cholesterol from plasma membranes and thereby disrupt lipid rafts [23]. Cells were preincubated with M\(\beta\)CD (10 mM for 30 min) before examining its effects on Ca\(^{2+}\) release and entry. Both thapsigargin and AA evoked higher amplitude Ca\(^{2+}\) release transients in cells treated with M\(\beta\)CD (Fig. 3Ai and Bi). In contrast, Ca\(^{2+}\) entry was significantly reduced by M\(\beta\)CD. The peak amplitude of thapsigargin-evoked SOCE in control cells was 189 ± 9 nM (n = 60 cells), this declined to 135 ± 7 nM (n = 108 cells) following M\(\beta\)CD incubation (Fig. 3Aii). However, the consequence of M\(\beta\)CD incubation on AA-mediated Ca\(^{2+}\) entry was even more dramatic. The robust Ca\(^{2+}\) entry evoked by AA in control cells (peak amplitude 335 ± 45 nM; n = 37 cells) was almost abolished in the presence of M\(\beta\)CD (peak amplitude 18 ± 3 nM; n = 61 cells).

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Fig. 3. Cholesterol depletion inhibits Ca\(^{2+}\) entry in HEK-293 cells. For depletion of cellular cholesterol, cells were incubated in 10 mM M\(\beta\)CD for 30 min prior to imaging. Panels A and B show quantitative analysis of the effect of cholesterol depletion on thapsigargin-mediated (A) and AA-mediated (B) Ca\(^{2+}\) changes. The responses from control cells and M\(\beta\)CD-treated cells are indicated by black and grey bars, respectively. The data represent the mean ± S.E.M. Statistical significance is denoted by *\(p < 0.05\). Panel C depicts a typical Ca\(^{2+}\) signal within an individual HEK-293 cell following M\(\beta\)CD treatment (grey trace). A control response (black trace; taken from Fig. 2Ai) is superimposed for comparison.
The lack of effect of AA on cells incubated with MβCD is illustrated in Fig. 3C. Whereas addition of AA to cells displaying thapsigargin-evoked SOCE usually caused rapid inhibition of Ca\(^{2+}\) entry (Fig. 2), following MβCD treatment there was no effect (Fig. 3C). Furthermore, the substantial LOE-sensitive Ca\(^{2+}\) entry triggered by AA addition (Fig. 2) was completely prevented (Fig. 3C). These data suggest that depletion of cholesterol from the plasma membrane partially inhibits SOCE, but completely curtails the effects of AA.

### 3.2. Prior activation of SOCE impacts on subsequent activation of AA-mediated Ca\(^{2+}\) entry

As described above, application of 30 \(\mu\)M AA to cells with an on-going SOCE response caused both a fast inhibition of SOCE and a slow switch to LOE-908-sensitive Ca\(^{2+}\) entry (e.g. Fig. 2). When applied to naïve cells, both thapsigargin and AA evoked a rapid Mn\(^{2+}\) entry response (Fig. 4A and B). The delayed development of AA-mediated Ca\(^{2+}\) entry following SOCE activation (Fig. 2) therefore contrasts with the ability of AA to trigger rapid cation influx in naïve cells.

We examined the effect of prior SOCE activation on Ca\(^{2+}\) influx triggered by a range of AA concentrations (5, 10 or 30 \(\mu\)M). As illustrated in Fig. 5Ai, cells were initially stimulated with AA in Ca\(^{2+}\)-free medium and extracellular Ca\(^{2+}\) was replenished after the decline of the Ca\(^{2+}\) release transient. The concentration-dependent effect of AA on Ca\(^{2+}\) entry is evident from the individual traces (Fig. 5Ai), the averaged rate of Ca\(^{2+}\) rise (Fig. 5B) and the peak amplitude of response (Fig. 5C).

When the same concentrations of AA were applied to cells in which SOCE had been previously activated by thapsigargin there was a significant reduction in the rate of AA-mediated Ca\(^{2+}\) entry (Fig. 5Aii and B). With 30 \(\mu\)M AA, the development of the Ca\(^{2+}\) entry signal was slowed to the same rate as observed using 5 \(\mu\)M AA on naïve cells (Fig. 5B). More striking, however, was the response to 5 \(\mu\)M AA. Whereas this concentration of AA evoked a robust Ca\(^{2+}\) entry when given to cells without activated SOCE (Fig. 5Ai), when it was applied to cells with an on-going SOCE response there was a very modest AA-mediated Ca\(^{2+}\) entry (Fig. 5Aii and C). It therefore appears that when AA is given as the first stimulus it can trigger a rapid influx of Ca\(^{2+}\). Prior stimulation of SOCE inhibits subsequent activation of the AA-mediated pathway. Both the rate of increase of the Ca\(^{2+}\) rise and amplitude of the Ca\(^{2+}\) signal through the AA-evoked entry pathway are diminished by SOCE (Fig. 5B and C).

Although the Ca\(^{2+}\) influx evoked by AA was concentration dependent, its effect in inhibiting SOCE was not. Each of the AA concentrations used in this study (5, 10 or 30 \(\mu\)M), appeared to fully inhibit SOCE since the cytosolic Ca\(^{2+}\) concentration always returned to basal levels before the LOE-908-sensitive Ca\(^{2+}\) entry developed (Figs. 2 and 5Aii).

### 3.3. Elevation of cytosolic Ca\(^{2+}\) is not responsible for slowing non-SOCE

Mignen and Shuttleworth [24] have proposed that Ca\(^{2+}\) influx through arachidonate-regulated Ca\(^{2+}\) (ARC) channels is inhibited by elevated intracellular Ca\(^{2+}\) levels. It is plausible therefore, that the Ca\(^{2+}\) signal arising from SOCE inhibited the subsequent development of AA-mediated Ca\(^{2+}\) entry. To examine this possibility, intracellular Ca\(^{2+}\) stores were passively depleted by incubating HEK-293 cells in Ca\(^{2+}\)-free medium (nominally Ca\(^{2+}\) free supplemented with 1 mM EGTA) for 2 h prior to imaging. This protocol was found to completely deplete the intracellular Ca\(^{2+}\) stores without a rise in cytosolic Ca\(^{2+}\) and consequently activate SOCE (data not shown). Subsequent addition of 30 \(\mu\)M AA (simultaneous with restoration of extracellular Ca\(^{2+}\)) resulted in a slowly rising Ca\(^{2+}\) entry signal (black trace in Fig. 6A), which was LOE-908 sensitive (grey trace in Fig. 6A). The rate of rise of the AA-induced Ca\(^{2+}\) signal in cells that had been preincubated in Ca\(^{2+}\)-free medium for 2 h was 0.08 \(\pm\) 0.01 nM s\(^{-1}\) (\(n = 17\) cells). This is similar to the sluggish development of Ca\(^{2+}\) influx when AA was applied following thapsigargin treatment (0.156 \(\pm\) 0.01 nM s\(^{-1}\); \(n = 71\)), and contrasts with the rapid activation of cation entry when the lipid was applied to naïve cells (Fig. 4). It therefore appears that activation of
Fig. 5. Prior activation of SOCE impacts on subsequent activation of AA-mediated Ca\textsuperscript{2+} entry in HEK-293 cells. (Ai) Naïve cells were treated with 5 \textmu M (grey trace) or 10 \textmu M (black trace) AA. Both AA concentrations evoked a transient Ca\textsuperscript{2+} release signal of comparable amplitude and a substantial Ca\textsuperscript{2+} entry response when extracellular Ca\textsuperscript{2+} was restored. (Aii) The effects of 5 \textmu M (grey trace) and 10 \textmu M (black trace) AA were substantially reduced when added to cells with on-going SOCE. The traces depict representative responses of single cells. Quantitative analysis of the initial rate (B) and peak amplitude (C) of AA-mediated Ca\textsuperscript{2+} influx in naïve cells (grey bars) and cells pre-treated with thapsigargin (black bars) is shown. The data represent mean ± S.E.M. of at least 30 cells analyzed in three independent experiments. Statistical significance is denoted by when *p < 0.01.

Fig. 6. Factors affecting AA-mediated Ca\textsuperscript{2+} entry. (A) Intracellular Ca\textsuperscript{2+} stores were passively depleted by incubation in Ca\textsuperscript{2+}-free medium for 2 h. Subsequently AA was added (simultaneous with the restoration of extracellular Ca\textsuperscript{2+}). Panels B and C depict AA-mediated Ca\textsuperscript{2+} influx following SOCE activation in HEK-293 (B) and Saos-2 (C) cells in the presence of 2-APB (grey traces) or Gd\textsuperscript{3+} (black traces). The SOCE blockers were added 2 min prior to thapsigargin and were present throughout the remainder of the experiment. The traces depict representative responses of single cells. (D) Modulation of AA-induced Ca\textsuperscript{2+} entry by the order of addition of SOCE channel blockers. The data represent the mean ± S.E.M. of at least 30 cells analyzed in three independent experiments.
SOCE either by thapsigargin or passive depletion of stores has the same effect of slowing Ca\(^{2+}\) entry triggered by AA.

LOE-908 blocks AA-mediated Ca\(^{2+}\) entry, but does not affect Ca\(^{2+}\) release by AA (data not shown). Therefore, the lack of Ca\(^{2+}\) signal in cells challenged with AA + LOE-908 in Fig. 6A indicates that the Ca\(^{2+}\) stores were depleted by incubation in Ca\(^{2+}\)-free medium for 2 h. If this were not the case, a Ca\(^{2+}\) release transient would have been observed.

The data described above suggest that activation of SOCE per se, but not the flux of Ca\(^{2+}\) through the store-operated channels inhibits the development of AA-stimulated Ca\(^{2+}\) entry. To establish this more fully, we examined the effect of inhibitors of SOCE. HEK-293 cells were incubated with either 100 \(\mu\)M 2-APB or 1 \(\mu\)M Gd\(^{3+}\) for 2 min prior to addition of 2 \(\mu\)M thapsigargin. Due to the presence of the SOCE antagonists, the response to thapsigargin was reduced to an acute Ca\(^{2+}\) release transient (Fig. 6B). For cells stimulated with thapsigargin in the presence of 2-APB, the subsequent addition of AA led to a slowly developing LOE-908-sensitive Ca\(^{2+}\) entry signal (grey trace in Fig. 6B). This response was similar to the slow on-set of AA-mediated Ca\(^{2+}\) influx in cells where SOCE had been stimulated by thapsigargin or following passive Ca\(^{2+}\) store depletion. In contrast, for cells stimulated with thapsigargin in the presence of 1 \(\mu\)M Gd\(^{3+}\) the subsequent addition of AA was without effect; there was no further change in cytosolic Ca\(^{2+}\) (black trace in Fig. 6B).

A similar pattern of response was observed using 2-APB and Gd\(^{3+}\) on either HEK-293 cells (Fig. 6B) or Saos-2 cells (Fig. 6C). These data concur with the results presented earlier, in that prior activation of SOCE impacts on the subsequent induction of AA-mediated Ca\(^{2+}\) influx. Since 2-APB substantially inhibits Ca\(^{2+}\) entry via SOCE (Fig. 1), the results obtained using this antagonist support our contention that Ca\(^{2+}\) influx through store-operated channels is not responsible for the subsequent slow development of AA-stimulated Ca\(^{2+}\) entry.

The effect of Gd\(^{3+}\) appeared to be even more extreme in that inhibition of SOCE using this cation completely prevented activation of Ca\(^{2+}\) entry by AA. It is important to note that Ca\(^{2+}\) entry evoked by addition of AA to naive cells was not substantially altered by Gd\(^{3+}\) (Fig. 1). Therefore, the complete block of AA-mediated Ca\(^{2+}\) entry by Gd\(^{3+}\) must be related to its effect on SOCE.

We examined whether 2-APB and Gd\(^{3+}\) had any overlap in their action by adding both inhibitors during the activation of SOCE, but changing the order in which they were applied. The protocol for these experiments was essentially to stimulate SOCE by application of 2 \(\mu\)M thapsigargin for 10 min. This was followed by the addition of 2-APB and/or Gd\(^{3+}\), and finally by superfusing cells with AA. When 2-APB was applied for 1 min before Gd\(^{3+}\), subsequent application of AA evoked the same slowly developing Ca\(^{2+}\) signal as observed when 2-APB was applied on its own (e.g. Fig. 6B). If Gd\(^{3+}\) was given simultaneously with 2-APB, the amplitude of the AA-mediated Ca\(^{2+}\) influx was significantly reduced (Fig. 6D). When Gd\(^{3+}\) was applied 1 min before 2-APB, the Ca\(^{2+}\) rise evoked by AA was dramatically reduced. These results concur with an earlier study [22], and suggest that 2-APB and Gd\(^{3+}\) have a mutually exclusive effect on the stimulation of AA-mediated Ca\(^{2+}\) influx. Both agents effectively block SOCE. However, only 2-APB allows the mode of Ca\(^{2+}\) entry to be switched from SOCE to the LOE-sensitive AA-stimulated pathway. Gd\(^{3+}\) somehow restrains the activation of this mechanism. The profound effect of changing the order of addition of 2-APB and Gd\(^{3+}\) suggests that whichever antagonist reached its target first determined the subsequent activation status of the AA-mediated Ca\(^{2+}\) influx pathway.

3.4. Calmidazolium rapidly switches SOCE to the LOE-908-sensitive Ca\(^{2+}\) entry pathway

We have previously demonstrated that high concentrations (\(\geq 10 \mu\)M) of calmidazolium activate the same Ca\(^{2+}\) entry pathway as AA [19]. The pharmacological profile of the Ca\(^{2+}\) entry pathway stimulated by calmidazolium [19] is the same as that shown for AA in the present study (Fig. 1). A typical response to calmidazolium is presented in Fig. 7A. When superfused over HEK-293 cells in Ca\(^{2+}\)-free medium, 10 \(\mu\)M calmidazolium evoked a rapid transient increase in intracellular Ca\(^{2+}\), which was followed by a robust Ca\(^{2+}\) entry signal when extracellular Ca\(^{2+}\) was

![Fig. 7. Calmidazolium-activated Ca\(^{2+}\) influx. (A) Ca\(^{2+}\) mobilization and entry induced by 10 \(\mu\)M calmidazolium. (B) Calmidazolium rapidly activated Ca\(^{2+}\) entry pathway following run down of SOCE (black trace) which was LOE-908 sensitive (grey trace). The traces depict representative responses of single cells.](image-url)
replaced (Fig. 7A). All cells responded in a similar manner ($n = 28$). Consistent with calmidazolium and AA stimulating the same $Ca^{2+}$ influx pathway, preincubation of cells with MβCD (10 mM for 30 min) significantly reduced the amplitude of calmidazolium-induced $Ca^{2+}$ influx. The peak $Ca^{2+}$ signal evoked by calmidazolium in control cells was $704.62 \pm 81.49$ nM (mean ± S.E.M., $n = 28$ cells). This was reduced to $90.05 \pm 11.33$ nM ($n = 83$ cells) following MβCD treatment.

With cells that had been treated with thapsigargin and therefore had an on-going SOCE response, application of calmidazolium caused a rapid $Ca^{2+}$ entry signal (black trace in Fig. 7B). This $Ca^{2+}$ influx was completely inhibited by 50 μM LOE-908 (grey trace in Fig. 7B), indicating that calmidazolium had activated the pathway that AA can also stimulate. However, unlike the slow development of the LOE-908-sensitive $Ca^{2+}$ entry signal when AA was applied during SOCE (e.g. Fig. 2), calmidazolium caused an almost immediate changeover. A rapid switch from SOCE to non-SOCE was also observed in experiments where AA and calmidazolium were added together ($n = 78$ cells). These data indicate that the non-SOCE pathway was potentially functional following SOCE activation, but AA is not capable of rapidly engaging it whereas calmidazolium can.

### 3.5. Effect of NO on SOCE and AA-mediated non-SOCE

It has been suggested that NO may act to couple activation of AA-evoked non-SOCE with the simultaneous inhibition of SOCE [21]. We therefore examined whether this messenger was responsible for any aspects of the inhibitory communication observed between SOCE and non-SOCE in this study.

Production of NO resulting from AA or thapsigargin application was monitored using the fluorescent indicator DAF-FM. This molecule is essentially non-fluorescent until it reacts with the nitrosonium cation (produced by spontaneous oxidation of nitric oxide). The resulting fluorescent compound is trapped in the cytoplasm, so that DAF-FM fluorescence summates with continual NO production. Perfusion of cells with 30 μM AA in $Ca^{2+}$-containing media...
resulted in a steady increase in DAF-FM fluorescence signal. In the presence of l-NAME (100 μM), which inhibits the NO-producing enzyme nitric oxide synthase (NOS) by competing with the natural substrate l-arginine, AA did not increase the DAF-FM signal (Fig. 8A and B). In contrast, thapsigargin did not provoke a change in DAF-FM fluorescence (Fig. 8C). AA also stimulated an increase in DAF-FM fluorescence in cells that had experienced prior treatment with thapsigargin (Fig. 8D). These data indicate that application of AA stimulated the production of NO. Whereas, Ca2+ store depletion and SOCE did not.

The NO donors sodium nitroprusside (SNP; 100 μM) and 2,2′-(hydroxynitrosohydrazino)bis-ethanamine (NOC-18; 1 mM) were utilised to examine whether NO affected Ca2+ influx through SOCE or non-SOCE. Cells were treated with thapsigargin or AA in Ca2+-free medium to activate SOCE or non-SOCE, respectively. Following the decline of the Ca2+ release signal, the NO donors were added and 3 min later Ca2+ was readmitted to the external solution. Typical responses to these treatments are depicted in Fig. 9. Both SNP and NOC-18 caused a modest decrease in SOCE (Fig. 9B). With non-SOCE, SNP caused an increase in the Ca2+ influx signal, whereas NOC-18 had no discernable effect (Fig. 9D).

We examined whether NO contributed in any part to the effect of AA on SOCE, or the slow development of non-SOCE in cells where SOCE had been activated. An experimental protocol similar to that depicted in Fig. 2Ai was used, with the difference that l-NAME or NO donor (SNP or NOC-18) were applied 3 min prior to AA addition. When l-NAME was used 11 out of 30 cells displayed a faster on-set of AA-evoked Ca2+ entry. Conversely, if SNP or NOC-18 were applied before AA addition, a number of cells (SNP, 1 out of 30; NOC-18, 4 out of 30) had a slower on-set of non-SOCE. These data could suggest that NO had a modest controlling influence on the inhibition of SOCE and triggering of non-SOCE in response to AA addition. However, the majority of cells were not affected by either l-NAME or NO addition, and the Ca2+ changes resembled the response shown in Fig. 2Ai.

4. Discussion

Consistent with other reports [16,17,22,25], the data in this study demonstrate that electrically non-excitable cells (HEK and Saos-2) express distinct SOCE and AA-activated non-SOCE pathways, which interact in a mutually antagonistic
manner. The SOCE and non-SOCE pathways were distinguished on the basis of the sensitivity to pharmacological blockers (Gd3+, 2-APB, LOE-908; Fig. 1) and depletion of cellular cholesterol using MβCD (Fig. 3). Both Ca2+ influx pathways could be rapidly activated in naïve cells (Fig. 4), and either route could give rise to long-lasting substantial Ca2+ entry signals (Fig. 1). The SOCE and non-SOCE pathways could not be activated simultaneously. Addition of AA to an on-going thapsigargin-mediated SOCE signal rapidly curtailed Ca2+ influx, and switched the mode of Ca2+ entry to the LOE-908-sensitive pathway (Fig. 2). Even with 5 μM AA, the inhibition of SOCE was rapid and complete (Fig. 5).

The mechanism underlying the inhibition of SOCE by AA is unclear. It has been suggested that NO, either directly or via stimulation of guanylate cyclase and protein kinase G, acts as a messenger downstream of AA to inhibit SOCE [21,25]. Consistent with that scheme, we observed that AA stimulated the production of NO (Fig. 8). However, in the present study exogenous application of substantial concentrations of NO donors did not fully inhibit SOCE (Fig. 9).

The SOCE of AA by BA is unlikely to involve Ca2+, since AA caused the cytoplasmic Ca2+ concentration to return to basal levels before the non-SOCE pathway developed. This suggests that immediately following AA addition there was a period with no Ca2+ influx at all. The simplest mechanism to explain our observations is that AA inhibits SOCE channels by direct interaction or altering their lipid environment, or that the non-SOCE channels are differentially sensitive to cholesterol depletion (Fig. 3A), non-SOCE was almost complete when incubated with 5-30 μM AA (Figs. 1 and 5). In contrast, LOE-sensitive Ca2+ entry developed slowly in cells where SOCE was already activated (Fig. 5). With 5 μM AA, the prior activation of SOCE reduced the subsequent non-SOCE signal to almost negligible levels (Fig. 5Aii).

It has been suggested that AA-stimulated Ca2+ entry is substantially inhibited by cytosolic Ca2+ concentrations >100 nM [17], and that this effect is mediated by activation of Ca2+-sensitive phosphatase calcineurin [28]. We therefore considered that the slow development of non-SOCE in cells where SOCE was active could have been due to the elevation of cytosolic Ca2+. However, passive depletion of intracellular Ca2+ stores by incubating cells in Ca2+-free medium had the same effect. It therefore appears that even without a corresponding Ca2+ rise, prior activation of SOCE significantly slows the ability of the non-SOCE mechanism to develop.

Calmidazolium caused the rapid activation of LOE-sensitive Ca2+ entry, even in cells where SOCE was already activated (Fig. 7). It is unclear why calmidazolium can override the effect of SOCE and rapidly engage the non-SOCE pathway, whereas AA cannot. Our previous work demonstrated that calmidazolium-induced Ca2+ entry was dependent on AA production via iPLA2, and that metabolism of AA was not required to activate the LOE-908-sensitive Ca2+ entry mechanism [19]. It therefore appears that AA and calmidazolium operate via exactly the same non-SOCE Ca2+ influx mechanism. However, calmidazolium clearly has some action, perhaps related to its effect as a calmodulin antagonist, which enables it to rapidly recruit the non-SOCE pathway, and is not shared by AA.

The specific cellular localization of SOCE and non-SOCE channels is unknown, and cannot be fully analyzed until the molecular entities that form these pathways are established. However, several TRPC isoforms have been localized within cholesterol-rich microdomains in the plasma membrane termed lipid rafts [29–31]. Depletion of these cholesterol-rich domains results in a reduction of Ca2+ entry in a number of cell types [32–34]. We observed that sequencing cholesterol reduced both SOCE and non-SOCE. Interestingly, although SOCE was significantly reduced following cholesterol depletion (Fig. 3A), non-SOCE was almost completely inhibited (Fig. 3B). These data could suggest that the SOCE and non-SOCE channels are differentially sensitive to cholesterol in their lipid environment, or that the non-SOCE channels are more abundantly expressed in lipid rafts and are thereby particularly sensitive to their disruption.

In summary, our data agree with previous studies that electrically non-excitable cells express both SOCE and non-SOCE, and that these influx pathways cannot be simultaneously activated [16,17,22]. However, our results do not concur with observations that NO or Ca2+ underlie the communication between the pathways. We envisage a scheme whereby cells express channels that are independently sensitive to store depletion or AA, but which share components that are critical for activation. In naïve cells, the apparatus for either Ca2+ influx pathway can be rapidly recruited. However, once the SOCE or non-SOCE mechanism is engaged, the proteins are effectively trapped in that particular channel configuration. Addition of AA to a cell with an on-going SOCE response immediately terminates Ca2+ influx due to an effect on SOCE channels [26,27], and then slowly recruits the protein(s) required for the non-SOCE pathway. We suggest that adding Gd3+ stabilizes the SOCE apparatus such that AA can no longer recruit the necessary components to form a viable Ca2+ entry channel. In contrast, although 2-APB blocks the SOCE pathway to the same extent as Gd3+, it does not stabilize SOCE in the same manner. This putative mechanism fits our data, but remains speculative until the molecular components of SOCE and AA-mediated non-SOCE are established in the future.
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References


