REVIEWS

CALCIUM SIGNALLING: DYNAMICS, HOMEOSTASIS AND REMODELLING

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Ca²⁺ is a highly versatile intracellular signal that operates over a wide temporal range to regulate many different cellular processes. An extensive Ca²⁺-signalling toolkit is used to assemble signalling systems with very different spatial and temporal dynamics. Rapid highly localized Ca²⁺ spikes regulate fast responses, whereas slower responses are controlled by repetitive global Ca²⁺ transients or intracellular Ca²⁺ waves. Ca²⁺ has a direct role in controlling the expression patterns of its signalling systems that are constantly being remodelled in both health and disease.

CALCIUM (

Ca²⁺ is a highly versatile intracellular signal that can regulate many different cellular functions^{1,2}. To achieve this versatility, the Ca²⁺-signalling system operates in many different ways to regulate cellular processes that function over a wide dynamic range (FIG. 1). At the synaptic junction, for example, Ca²⁺ triggers exocytosis within microseconds, whereas at the other end of the scale Ca²⁺ has to operate over minutes to hours to drive events such as gene transcription and cell proliferation. One of the challenges is to understand how these widely different Ca²⁺-signalling systems can be set up to control so many divergent cellular processes.

At any moment in time, the level of intracellular Ca²⁺ is determined by a balance between the 'on' reactions that introduce Ca²⁺ into the cytoplasm and the 'off' reactions through which this signal is removed by the combined action of buffers, pumps and exchangers (FIG. 1). During the on reaction, a small proportion of the Ca²⁺ binds to the effectors that are responsible for stimulating numerous Ca²⁺-dependent processes (FIG. 1). These heterogeneous Ca²⁺-signalling systems are assembled from an extensive Ca²⁺-signalling toolkit¹ (BOX 1). Through alternative splicing, many of the toolkit components have different isoforms with subtly different properties, which further expands the versatility of Ca²⁺ signalling.

This review concentrates on the nature of the Ca²⁺signalling toolkit and how it is exploited to create many
different Ca²⁺-signalling systems. We have gathered

emerging evidence that the expression patterns of the different signalling components might be regulated by Ca²⁺ itself to set up a 'self-assessment system' to ensure that these signalling systems remain constant. Alterations in this Ca²⁺-dependent homeostatic mechanism might be the cause of many prominent diseases, as exemplified by end-stage heart failure.

Ca2+-signalling toolkit and signalling dynamics

Each cell type expresses a unique set of components from the Ca²⁺-signalling toolkit to create Ca²⁺-signalling systems with different spatial and temporal properties. Almost all Ca²⁺-signalling systems have one thing in common — they function by generating brief pulses of Ca²⁺. These Ca²⁺ transients are created by variations of the basic on/off reactions that are outlined in Fig. 1.

Signal Ca²⁺ is derived either from internal stores or from the external medium (FIG. 1). In the case of the latter, there are many different plasma-membrane channels (BOX 1) that control Ca²⁺ entry from the external medium in response to stimuli that include membrane depolarization, stretch, noxious stimuli, extracellular agonists, intracellular messengers and the depletion of intracellular stores. The release of Ca²⁺ from the internal store — usually the endoplasmic reticulum (ER) or its muscle equivalent, the sarcoplasmic reticulum (SR) — is controlled by Ca²⁺ itself, or by an expanding group of messengers³, such as inositol-1,4,5-trisphosphate (Ins(1,4,5)P₃), cyclic ADP ribose (cADPR), nicotinic

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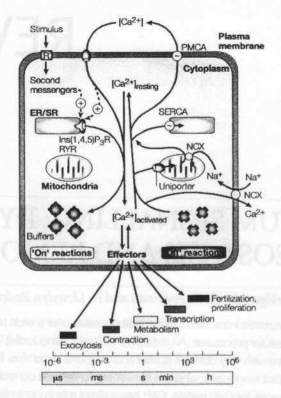


Figure 1 | Calcium-signalling dynamics and homeostasis During the 'on' reactions, stimuli induce both the entry of external Ca2+ and the formation of second messengers that release internal Ca2+ that is stored within the endoplasmic/ sarcoplasmic reticulum (ER/SR). Most of this Ca2+ (shown as red circles) is bound to buffers, whereas a small proportion binds to the effectors that activate various cellular processes that operate over a wide temporal spectrum. During the 'off' reactions, Ca2+ leaves the effectors and buffers and is removed from the cell by various exchangers and pumps. The Na⁺/Ca² exchanger (NCX) and the plasma-membrane Ca2+-ATPase (PMCA) extrude Ca2+ to the outside, whereas the sarco(endo)plasmic reticulum Ca2+-ATPase (SERCA) pumps Ca2+ back into the ER. Mitochondria also have an active function during the recovery process in that they sequester Ca2 rapidly through a uniporter, and this is then released more slowly back into the cytosol to be dealt with by the SERCA and the PMCA. Cell survival is dependent on Ca2+ homeostasis whereby the Ca2+ fluxes during the off reactions exactly match those during the on reactions. [Ca2+], Ca2+ concentration; Ins(1,4,5)P₃R, inositol-1,4,5-trisphosphate receptor; RYR, ryanodine receptor.

acid adenine dinucleotide phosphate (NAADP) and sphingosine-1-phosphate (S1P), that either stimulate or modulate the release channels on the internal stores.

Inositol-1,4,5-trisphosphate. Many stimuli function through Phospholipase C (PLC) to generate $Ins(1,4,5)P_3$ that functions to release Ca^{2+} from an internal store (FIG. 2). There are several PLC isoforms (BOX 1) that are activated by different mechanisms, such as G-protein-coupled receptors (PLC β), tyrosine-kinase-coupled receptors (PLC γ), an increase in Ca^{2+} concentration (PLC γ) or activation through Ras (PLC γ). The $Ins(1,4,5)P_3$ that is responsible for triggering the Ca^{2+}

oscillations that are required to activate mammalian eggs during fertilization is generated by a newly discovered PLC, PLC ζ , that is injected into the egg by the sperm⁵ (FIG. 2).

The dynamics of Ins(1,4,5)P, production can be very different depending on the receptor type being activated⁶. Bradykinin and neurokinin A receptors give large rapid Ca2+ transients, whereas lysophosphatidic acid (LPA), thrombin and histamine receptors give smaller responses that develop slowly but persist for much longer. Some of this variability might arise from the fact that receptors engage different transducing elements and PLC isoforms in a cell-type-specific manner7,8. For example, at the parallel fibre/Purkinje cell synapse, glutamate operates through a metabotropic glutamate receptor type I (mGluR1)→G_{or}→PLCβ4→ $Ins(1,4,5)P_1 \rightarrow Ins(1,4,5)P_1$ receptor 1 ($Ins(1,4,5)P_3R1$) →Ca2+ cascade, whereas hippocampal neurons respond to the same agonist using a mGluR5→G_{11α}→PLCβ1→ $Ins(1,4,5)P_1 \rightarrow Ins(1,4,5)P_1R \rightarrow Ca^{2+}$ sequence. The reason why neurons use these different signalling cascades is not known, but there is evidence from other cell types that mGluR1 and mGluR5 can result in radically different Ca2+ signals9. mGluR1 produces a single Ca2+ transient, whereas mGluR5 generates an oscillatory pattern.

Another example is found in pancreatic acinar cells in which muscarinic receptors generate small, highly localized Ca²⁺ transients, whereas the cholecystokinin (CCK) receptors produce much larger global Ca²⁺ transients. These different outputs might depend on the action of the regulators of G-protein signalling (RGS)¹⁰. In comparison to CCK receptors, muscarinic receptors are much more sensitive to the inhibitory action of RGS proteins (RGS2, RGS4 and RGS16), and this might limit the supply of Ins(1,4,5)P₃. Qualitatively different Ca²⁺ signals might also arise if receptors combine Ins(1,4,5)P₃ with other Ca²⁺-mobilizing messengers/modulators such as cADPR or NAADP¹¹⁻¹³, as discussed in the next section.

cADPR and NAADP. These two nucleotides mobilize intracellular Ca2+ through different mechanisms, even though they are generated through the same enzymatic pathway14 (FIG. 2). Mammalian cells express CD38, which is a multifunctional ADP ribosyl cyclase with both synthase and hydrolase activity. The synthase component of CD38 can use either NAD to produce cADPR or NADP to generate NAADP. CD38 has been located both on the cell surface and on intracellular membranes. On the cell surface, CD38 has been suggested to both produce cADPR and NAADP and to transport them into cells. Different activation mechanisms have been proposed for the cytosolic enzyme. External agonists might activate it, but a consistent mechanism for the transduction process is still lacking. An alternative possibility is that the formation of cADPR and NAADP is sensitive to cellular metabolism (FIG. 2). In other words, cADPR and NAADP might be metabolic messengers that can relay information about the state of cellular metabolism to the Ca2+-signalling pathways. Such an idea is supported by the fact that cADPR metabolism by the hydrolase is inhibited by either ATP15 or NADH16 (FIG. 2).

PHOSPHOLIPASE C (PLC). A phosphodiesterase that splits the bond between the phosphorus atom and the oxygen atom at C-1 of the glycerol moiety.

Box 1 | Calcium-signalling toolkit

The Ca²⁺-signalling system has a very large toolkit of signalling components that can be mixed and matched to create a diverse array of signalling units that can deliver Ca²⁺ signals with very different spatial and temporal properties. The following list is by no means inclusive but it summarizes some of the main toolkit components in mammalian cells:

Receptors

Receptors $(C_{1,A-C})$ and $(C_{1,A-C})$ are a receptor $(C_{1,A-C})$ branch $(C_{1,A-C})$ branch $(C_{1,A-C})$ and $(C_{1,A-C})$ are a receptor $(C_{1,A-C})$ and $(C_{1,A-C})$ and $(C_{1,A-C})$ are a receptor $(C_{1,A-C})$ and $(C_{1,A-C})$ are a receptor $(C_{1,A-C})$ and $(C_{1,A-C})$ are a receptor

Tyrosine-kinase-linked receptors: platelet-derived growth factor receptors (PDGFRα, PDGFRβ) | epidermal growth factor receptors (ERBB1–4)

Transducers

G proteins: $G_{q\alpha}$, $G_{11\alpha}$, $G_{14\alpha}$, $G_{16\alpha} \mid G_{\beta\gamma}$

Regulators of G-protein signalling (RGS): RGS1 | RGS2 | RGS4 | RGS16

Phospholipase C (PLC): PLCβ1-4 | PLCγ1, PLCγ2 | PLCδ1-4 | PLCε | PLCζ

ADP ribosyl cyclase: CD38

Channels

Voltage-operated channels (VOCs): Ca_v1.1, Ca_v1.2, Ca_v1.3, Ca_v1.4 (L-type) | Ca_v2.1 (P/Q-type) | Ca_v2.2 (N-type) | Ca_v2.3 (R-type) | Ca_v3.1, Ca_v3.2, Ca_v3.3 (T-type)

 $Receptor-operated\ channels\ (ROCs): \ NMDA\ receptors\ (NR1, NR2A, NR2B, NR2C, NR2D) \ |\ ATP\ receptor\ (P2X_{y}) \ |\ nACh\ receptor\ (P2X_{y}) \ |\ n$

 $Second-messenger-operated\ channels\ (SMOCs):\ cyclic\ nucleotide\ gated\ channels\ (CNGA\ 1-4,CNGB\ 1,CNGB\ 3)\ |\ arachidonate-regulated\ Ca^{2+}\ channel\ (I_{ARC})$

Transient receptor potential (TRP) ion-channel family: TRPC1-7 | TRPV1-6 | TRPM1-8 | TRPML (mucolipidin 1,2) | TRPNI (ANKTM1)

Inositol-1,4,5-trisphophate receptors (Ins(1,4,5)P,Rs): Ins(1,4,5)P,R1-3

Ryanodine receptors (RYRs): RYR1-3

Polycystins: PC-1 | PC-2

Channel regulators: triadin | junctin | sorcin | FKBP12 | FKBP12.6 | phospholamban | IRAG | IRBIT

Calcium buffers

Cytosolic buffers: calbindin D-28 | calretinin | parvalbumin

ER/SR buffers and chaperones: calnexin | calreticulin | calsequestrin | GRP 78 | GRP 94

Calcium effectors

Ca2+-binding proteins: calmodulin | troponin C | synaptotagmin | S100A1-12, S100B, S100C, S100P | annexin I-X | neuronal Ca2+ sensor family (NCS-1) | visinin-like proteins (VILIP-1, VILIP-2, VILIP-3) | hippocalcin | recoverin | Kv-channel-interacting proteins (KchIP1-3) | guanylate-cyclase-activating proteins (GCAP1-3)

Calcium-sensitive enzymes and processes

Ca2+-regulated enzymes: Ca2+/calmodulin-dependent protein kinases (CaMKI–IV)| myosin light chain kinase (MLCK)| phosphorylase kinase | Ins(1,4,5)P, 3-kinase | PYK2 | protein kinase C (PKC- α , PKC- β I, PKC- β II, PKC- γ) | cyclic AMP phosphodiesterase (PDE1A–C) | adenylyl cyclase (AC-1, AC-III, AC-VIII, AC-V, AC-VI) | nitric oxide synthase (endothelial NOS, eNOS; neuronal NOS, nNOS)| calcineurin | Ca2+-activated proteases (calpain I and II)

Transcription factors: nuclear factor of activated T cells (NFATc1-4) | cyclic AMP response element-binding protein (CREB) | downstream regulatory element modulator (DREAM) | CREB-binding protein (CBP)

 Ca^{2+} -sensitive ion channels: Ca^{2+} -activated potassium channels (SK, small conductance Ca^{2+} -sensitive channel; IK, intermediate conductance Ca^{2+} -sensitive channel; BK, large conductance Ca^{2+} -sensitive channel) | Human Cl^- channel, Ca^{2+} -activated (HCLCA1)

Calcium pumps and exchangers

Na⁺/Ca²⁺ exchangers (NCXs): NCX1-3

 $\label{eq:mitochondrial} \emph{Mitochondrial channels and exchangers: } \textit{permeability transition pore} \ | \ Na^+/Ca^{2^+}\ exchanger \ | \ Ca^{2^+}\ uniporter \ | \ H^+/Ca^{2^+}\ exchanger \ | \ Ca^{2^+}\ exchanger \ | \ Ca^{2$

Plasma membrane Ca2+-ATPases (PMCAs): PMCA1-4

Sarco(endo)plasmic reticulum Ca2+-ATPases (SERCAs): SERCA1-3

Golgi pumps: SPCA1, SPCA2

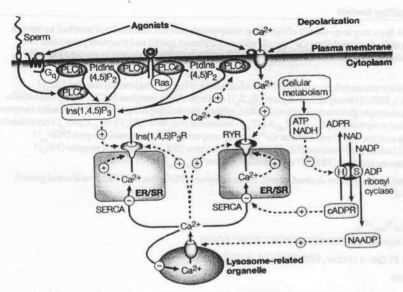


Figure 2 | Calcium-mobilizing messengers and modulators. Various second messengers or modulators regulate the release of Ca²⁺ from internal stores by the inositol-1,4,5-trisphosphate receptor (Ins(1,4,5)P₃R) or the ryanodine receptor (RYR). These release channels are sensitive to factors that function from the cytosol and from within the lumen of the endoplasmic/sarcoplasmic reticulum (ERVSR). The Ins(1,4,5)P₃R is regulated by inositol-1,4,5-trisphosphate (Ins(1,4,5)P₃) which is generated by various signalling pathways using different isoforms of phospholipase C (PLC; $\beta, \delta, \epsilon, \gamma$ and ζ). The nucleotides cyclic ADP ribose (cADPR) and nicotinic acid adenine dinucleotide phosphate (NAADP) are generated by ADP ribosyl cyclase, which has both synthase (S) and hydrolase (H) activity. This dual-function enzyme might be sensitive to the cellular metabolism, as ATP and NADH inhibit the hydrolase. Just how cADPR and NAADP function is still unclear, but they seem to have an indirect action, cADPR might function by stimulating the sarco(endo)plasmic reticulum Ca2+-ATPase (SERCA) pump, which increases the luminal level of Ca2+, which results in sensitization of the RYR. NAADP functions through a channel that is located on a lysosomal-like organelle to release Ca2+ that can either stimulate the Ins(1,4,5)P₃R or the RYR directly, or might function indirectly, like cADPR, to increase the lumenal level of Ca2+. Ptdlns(4,5)P2, phosphatidylinositol-4,5-bisphosphate.

RYANODINE RECEPTOR
(RYR). A Ca²⁺-release channel
that is located in the membrane
of the endoplasmic/sarcoplasmic
reticulum and is regulated by
several factors including Ca²⁺
itself, as well as the intracellular
messenger cyclic ADP ribose.

Ca**-INDUCED Ca** RELEASE (CICR). An autocatalytic mechanism by which cytoplasmic Ca** activates the release of Ca** from internal stores through channels such as inositol-1,4,5-trisphosphate receptors or ryanodine receptors.

VOLTAGE-OPERATED CHANNEL (VOC). A plasma-membrane ion channel that is activated by membrane depolarization.

NAADP functions by releasing Ca2+ from an internal store, which was recently identified in sea urchin eggs as a reserve granule store. This store is distinct from the stores that are regulated by Ins(1,4,5)P,Rs and RYANODINE RECEPTORS (RYRs), because the latter stores can be depleted without affecting the ability of NAADP to release Ca2+. This separate reserve granule store might be equivalent to a lysosome-related organelle in mammalian cells17 (FIG. 2). In contrast to the Ins(1,4,5)P, Rs and RYRs, the NAADP release mechanism is not sensitive to Ca2+ and therefore does not support the process of Ca11-INDUCED Ca11 RELEASE (CICR). NAADP seems to have a role in both initiating and coordinating various Ca2+signalling systems11,12. NAADP might sensitize the Ins(1,4,5)P, Rs and RYRs through two mechanisms (FIG. 2): it might function directly by providing trigger Ca2+, or indirectly by releasing a bolus of Ca2+, which is then taken up by the other stores in which it can sensitize the release channels by functioning from the lumen.

The function of cADPR resembles that of a modulator rather than a messenger. When cADPR is introduced into cells there usually is no immediate effect^{18,19}. In those cases in which it elicits a Ca²⁺ response, there usually is a

long latency^{20,21}, which indicates that it might be functioning indirectly by increasing the Ca²⁺ sensitivity of RYRs, as has been shown in neurons^{18,19,22} and in the heart^{21,23}. In these two excitable cells, voltage operated channels (VOCs) respond to membrane depolarization by admitting a small pulse of Ca²⁺, which then stimulates the RYRs to release further Ca²⁺ through CICR (FIG. 2). The degree to which the initial entry signal is amplified by CICR, which is referred to as the 'gain' of the signalling system, can be regulated by cADPR. This might have pathological consequences, as cardiac arrhythmias can develop if cADPR sets the gain too high²⁴.

Just how cADPR functions to modulate the sensitivity of the RYRs is still unclear. One view is that cADPR functions as a messenger to stimulate Ca²⁺ release by the RYRs¹⁴. However, cADPR does not seem to bind directly to the RYR, instead it seems to function through some intermediary — the FK506-binding protein 12.6 (FKBP12.6), which is a subunit that is associated with the RYR (BOX 2, part a), has been suggested as a possible candidate²⁵. An alternative view is that cADPR functions by activating the sarco(endo)Plasmic reticulum Ca²⁺-ATPASE (SERCA) pump²³. Such an indirect action is consistent with many of the properties of cADPR. By stimulating the SERCA pump, cADPR would enhance the load of Ca²⁺ within the lumen of the ER — a process that is known to increase the Ca²⁺ sensitivity of RYRs^{23,26} (FIG. 2).

Sphingolipid-derived messengers. Some agonistevoked Ca2+ signals might be controlled by a sphingolipid-activated Ca2+-release pathway that functions independently of Ins(1,4,5)P,Rs or RYRs27. In mast cells, the sphingolipid-derived messenger S1P functions together with Ins(1,4,5)P3 to generate the Ca2+ signals that underlie the synthesis and release of inflammatory mediators. The dual activation of these pathways leads to a Ca2+ signal with a rapid peak (S1P dependent) and a sustained plateau (Ins(1,4,5)P, dependent)28. Exactly how S1P stimulates Ca2+ release from intracellular stores is unclear. Until recently, the best candidate for the S1P receptor was a widely expressed protein known as SCaMPER (sphingolipid Ca2+-release-mediating protein of endoplasmic reticulum). However, it has no similarity to any known intracellular Ca2+ channel and is a small (~20-kDa) protein with only one transmembrane domain. A recent reinvestigation of SCaMPER showed that there was little correlation between its intracellular location and that of known intracellular Ca2+ stores. Furthermore, expression of SCaMPER was found not to confer sensitivity to sphingolipids or to affect Ca2+ homeostasis, but could lead to cell death29.

Ca²⁺-entry mechanisms. Ca²⁺ that enters the cell from the outside is a principal source of signal Ca²⁺ during the on reaction (FIG. 1). Entry of Ca²⁺ is driven by the presence of a large electrochemical gradient across the plasma membrane. Cells use this external source of signal Ca²⁺ by activating various entry channels with widely different properties. We know the most about the VOCs, which are found in excitable cells

Box 2 | Multimolecular complexes of calcium-signalling components

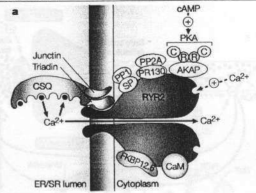
Like many other signalling systems, the components of the different Ca²⁺-signalling systems are often grouped together into large complexes.

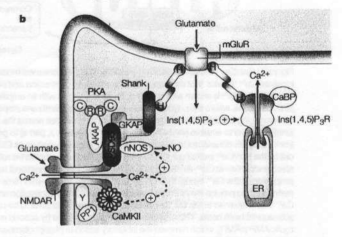
The RYR2 Ca2+-release complex in cardiac cells Ryanodine receptor 2 (RYR2) is composed of four subunits that form the channel, which is associated with various proteins that function to modulate its opening (see figure, part a). The endoplasmic/sarcoplasmic reticulum (ER/SR) luminal Ca2+binding protein calsequestrin (CSQ) modulates the sensitivity of RYR2 (FIG. 2). The interaction between CSQ and RYR2 is facilitated by the transmembrane proteins triadin and junctin. The reversible phosphorylation of RYR2 by cyclic AMP (cAMP) is controlled by protein kinase A (PKA), which is composed of regulatory (R) and catalytic (C) subunits that are attached through an A kinase anchoring protein (AKAP). Dephosphorylation depends on protein phosphatase 2A (PP2A), which is attached through the isoleucine-zipper-binding scaffolding protein PR130, and on protein phosphatase 1 (PP1), which is attached through spinophilin (SP). RYR2 is also modulated by calmodulin (CaM) and by FK506-binding protein 12.6 (FKBP12.6).

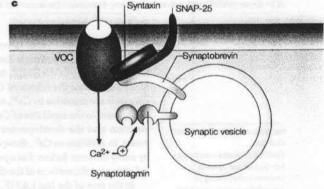
NMDA and Ins(1,4,5)P₃ receptor complexes in neurons Glutamate-induced Ca²⁺ entry is carried out by NMDA (N-methyloaspartate) receptors (NMDARs) that are linked to other signalling components (see figure, part b), some of which are Ca²⁺ sensitive, such as Ca²⁺/CaM-dependent kinase II (CaMKII) and neuronal nitric oxide synthase (nNOS). NMDARs are also associated with other proteins such as yotiao (Y), which binds PP1, and the scaffolding protein PSD95, which links into other signalling components such as the AKAP-PKA complex and guanylate kinase-associated protein (GKAP). Proteins such as shank and Homer (H) might link the metabotropic glutamate receptor (mGluR) to both the NMDAR and the inositol-1,4,5-trisphospate receptor (Ins(1,4,5)P₃R), which is also associated with a Ca²⁺-binding protein (CaBP).

Synaptic-vesicle complex

The exocytotic release of synaptic vesicles, which occurs within microseconds of membrane depolarization, is activated by an influx of Ca²⁺ through the Ca₂2 voltage-operated channels (see figure, part c), for example, the P/Q-, N- and R-type VOCs (BOX 1). These entry channels have a binding site that anchors them to components of the exocytotic machinery such as syntaxin or SNAP-25. In this way, the Ca²⁺ that enters through the channel has immediate access to the synaptotagmin Ca²⁺ sensor that is thought to initiate the fusion event that is carried out by syntaxin, SNAP-25 and synaptobrevin.







SARCO(ENDO)PLASMIC
RETICULUM Ca**-ATPASE
(SERCA). A pump located in
sarcoplasmic or endoplasmic
reticulum membranes that
couples ATP hydrolysis to the
transport of Ca** from cytosolic
to lumenal spaces.

RECEPTOR-OPERATED CHANNEL (ROC). A plasma-membrane ion channel that opens in response to the binding of an extracellular ligand. and generate the rapid Ca²⁺ fluxes that control fast cellular processes such as muscle contraction or exocytosis at synaptic endings (BOX 2, part c).

There are many other Ca²⁺-entry channels (BOX 1) that open in response to different external signals, such as the RECEPTOR-OPERATED CHANNELS (ROCs), for example the NMDA (N-methyl-D-aspartate) receptors (NMDARs) that respond to glutamate (BOX 2, part b). There also are SECOND-MESSENGER-OPERATED CHANNELS (SMOCs) that are controlled by internal messengers, such as the cyclic-nucleotide-gated channels that are found in sensory systems and the arachidonic-acid-sensitive channel³⁰. In addition to these more clearly defined channel-opening mechanisms, there are many other channel types that

are sensitive to a diverse array of stimuli, such as the STORE-OPERATED CHANNELS (SOCs), thermosensors and stretch-activated channels. Many of these channels belong to the large transient receptor protein (TRP) ion-channel family³¹⁻³⁴, which are encoded by up to 23 different genes. This family consists of three groups — the canonical TRPC family, the vanilloid TRPV family and the melastatin TRPM family (BOX 1). TRP channels tend to have low conductances and therefore can operate over much longer time scales without swamping the cell with too much Ca²⁺. Members of the TRP family are particularly important in controlling slow cellular processes such as smooth-muscle contractility and cell proliferation.

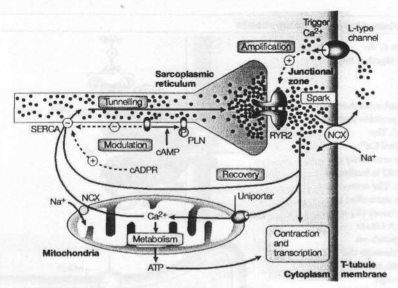


Figure 3 | Cardiac calcium-signalling module. This self-contained module generates the localized Ca²+ 'sparks' that are responsible for activating contraction and perhaps also gene transcription (Ca²+ is shown as red circles). Signalling begins with an amplification step in the junctional zone, where the L-type channel on the T-tubule membrane responds to depolarization by introducing a small pulse of trigger Ca²+, which then diffuses across the narrow gap of the junctional zone to activate ryanocline receptor 2 (RYR2) (BOX 2, part a) to generate a spark. Ca²+ from this spark diffuses out to activate contraction. Recovery occurs as Ca²+ is pumped out of the cell by the Na¹/Ca²+ exchanger (NCX) or is returned to the sarcoplasmic reticulum (SR) by sarco(endo)plasmic Ca²+-ATPase (SERCA) pumps on the non-junctional region of the SR. A proportion of this Ca²+ travels through the mitochondria, during which time it stimulates the metabolism to provide the ATP that is necessary to maintain contraction and transcription. The Ca²+ that is returned to the SR 'tunnels' back to the junctional zone to be used again for subsequent heart beats. This circulation of Ca²+ is modulated by second messengers such as cyclic AMP (cAMP), which removes the inhibitory action of phospholamban (PLN), or by cyclic ADP ribose (cADPR) that activates the pump to increase the amount of releasable Ca²+ in the SR.

SECOND-MESSENGER-OPERATED CHANNEL (SMOC). A plasma-membrane

ion channel that opens in response to the binding of intracellular second messengers such as diacylglycerol, cyclic nucleotides or arachidonic acid.

STORE-OPERATED CHANNEL (SOC). A plasma-membrane ion channel, of uncertain identity, that opens in response to the depletion of internal Ca²⁺ stores.

FIGHT-OR-FLIGHT RESPONSE
This response occurs in the
hypothalamus, which, when
stimulated by stress, initiates a
sequence of nerve-cell firing and
chemical release (including
adrenaline and noradrenaline)
that prepares our body for
running or fighting.

Ca²⁺ release from internal stores. The other principal source of Ca²⁺ for signalling is the internal stores that are located primarily in the ER/SR, in which Ins(1,4,5)P₃Rs or RYRs regulate the release of Ca²⁺ (FIG. 2). These two channels are sensitive to Ca²⁺, and this CICR process contributes to the rapid rise of Ca²⁺ levels during the on reaction and the development of regenerative Ca²⁺ waves. In addition to Ca²⁺, these channels are regulated by many different factors that operate on both the luminal and cytosolic surfaces of the channel (BOX 2, part a).

In the case of the Ins(1,4,5)P,Rs, the primary determinants are Ins(1,4,5)P, and Ca2+. The binding of Ins(1,4,5)P, increases the sensitivity of the receptor to Ca2+, which has a biphasic action (that is, it activates at low concentrations, but becomes inhibitory at the higher concentrations that occur after Ca2+ release). This Ca2+ regulation is mediated by the direct action of Ca2+ on the receptor, as well as indirectly through calmodulin (CaM), which can be activating or inhibitory35,36. Recently, a Ca2+-binding protein (CaBP) was shown to activate Ca2+ release in the absence of Ins(1,4,5)P, (REF 37). However, this new function for CaBP is contentious38. In addition to these cytosolic actions, Ca2+ can also sensitize the Ins(1,4,5)P, R by functioning from the lumen (FIG. 2). This luminal sensitivity might be conferred by the ER luminal lectin chaperones calreticulin and calnexin, which are Ca^{2+} -binding proteins that are known to interact with the $Ins(1,4,5)P_3R$. The Ca^{2+} -binding sites on calreticulin have affinities that are sufficiently low to enable them to regulate $Ins(1,4,5)P_3Rs$ through luminal Ca^{2+} levels.

The InsP,R can also be modulated by other signalling pathways, including phosphorylation by Ca2+/CaMdependent kinase II (CaMKII), cGMP-dependent protein kinase (PKG), protein kinase C (PKC) and cAMPdependent protein kinase (protein kinase A; PKA)2. For some of these kinases, the scaffolding proteins that mediate recruitment to their site of action on the Ins(1,4,5)P, R have been identified. For example, PKG is recruited by the Ins(1,4,5)P, R-associated cGMP kinase substrate (IRAG), which results in Ins(1,4,5)P,R phosphorylation and a decrease in receptor activity39. In B cells, the Ins(1,4,5)P,R is phosphorylated by the tyrosine kinase Lyn, which results in increased activity. This phosphorylation event is facilitated by the scaffolding protein Bank that links together Lyn, the Ins(1,4,5)P,R and the B-cell receptor 10. Similarly, in neuronal cells Ins(1,4,5)P, Rs are tethered to mGluRs by the Homer protein, thereby linking the source of Ins(1,4,5)P, production to its site of action (BOX 2, part b). The protein phosphatases 1 and 2a (PP1 and PP2a) have been found to co-purify with PKA and the Ins(1,4,5)P,R, which is reminiscent of their interaction with the RYR41. This protein complex of phosphatase, kinase and substrate allows the rapid regulation of Ins(1,4,5)P,R activity by phosphorylation/dephosphorylation.

RYRs are also controlled by several signalling pathways, as illustrated by the RYR2 that is found in cardiac cells. Like the Ins(1,4,5)P₃R, RYR2 responds to Ca²⁺ in a bell-shaped fashion (that is, RYR2 is inactive at low nM concentrations of Ca²⁺, active at low µM concentrations of Ca²⁺ and inactivated by high concentrations of Ca²⁺ that are in the mM range)⁴². In the cardiac myocyte, Ca²⁺ that enters through the L-type Ca²⁺ channel activates RYR2 to create the 'spark' that triggers contraction (FIG. 3). This role of RYR2 in excitation—contraction (E—C) coupling is a highly regulated process that involves many accessory factors that are bound to both its luminal and cytosolic domains (BOX 2, part a).

Opening of RYR2 is inhibited by CaM, which is present in Ca2+-bound and non-bound forms, which are known as CaCaM and apoCaM, respectively43. Several accessory proteins contribute to the control of heart function by the sympathetic nervous system, through the FIGHT-OR-FLIGHT RESPONSE⁴⁴. After β-adrenergic stimulation, RYR2 is phosphorylated by PKA, which is attached through an A kinase anchoring protein (AKAP)45 (BOX 2, part a). Phosphorylation of RYR2 by PKA results in the dissociation of the 12.6-kDa FKBP12.6, which normally stabilizes the RYR in a closed conformation45. Furthermore, FKBP12.6 is required for the coupled gating between neighbouring receptors that coordinates the activation and inactivation of physically linked receptors during E-C coupling44. PKA also phosphorylates sorcin, which is another regulator of RYR2 (REF. 46).

The RYR2 macromolecular complex also includes the phosphatases PP1 and PP2a, which interact with RYR2 through the leucine/isoleucine-zipper-binding scaffolding proteins spinophilin and PR130, respectively (BOX 2, part a)44. The presence of these phosphatases in the same protein complex as the kinase and substrate ensure that there is a tight regulation of the phosphorylation status of the receptor and, therefore, its activity. The membrane and luminal region of the RYR2 is present in a complex with three other proteins: junctin, triadin and calsequestrin (BOX 2, part a)47,48. Calsequestrin is the principal Ca2+-binding protein of muscle cells and is highly concentrated in the junctional region of the SR. In the lumen of the SR, calsequestrin does not bind directly to the RYR, but is anchored adjacent to the Ca2+-release site through junctin and triadin, which are both membrane-bound proteins (BOX 2, part a)48. Triadin and junctin interact with calsequestrin in a Ca2+-dependent manner48, and this interaction might account for the sensitivity of RYR2s to Ca2+ in the lumen23,26. Indeed, transgenic studies have shown that there is a significant role for calsequestrin, junctin and triadin in cardiac Ca2+ signalling and hypertrophy (see below).

Emerging Ca2+ channels. AUTOSOMAL-DOMINANT POLYCYSTIC KIDNEY DISEASE (ADPKD) has been linked to mutations in two membrane-spanning proteins, which are known as polycystin-1 (PC-1) and polycystin-2 (PC-2)49. PC-1 has a large extracellular domain and might function in transducing sensory information, such as shear stress during fluid flow⁴⁹. PC-2 has been shown to function as an intracellular Ca2+-release channel50 and to form a non-selective cation channel when it is inserted into the plasma membrane51. PC-2 has homology with VOCs and TRP channels, and when expressed by itself it shows spontaneous channel activity. Mutations in either protein somehow corrupts the PC-1-PC-2 complex, which leads to abnormal Ca2+ signalling and, consequently, altered rates of cell proliferation and function. These proteins seem to have a widespread expression, although their role outside the kidney is not well understood. Interestingly, PC-2 is localized to the ER in some cell types52, which indicates that it might be a ubiquitous CICR channel.

Ca²⁺-binding proteins. During the on reaction (FIG. 1), Ca²⁺ flows into the cell and interacts with different Ca²⁺-binding proteins, of which there are ~200 encoded by the human genome that function either as Ca²⁺ effectors or buffers².

The buffers, which become loaded with Ca²⁺ during the on reaction and unload during the off reaction, function to fine-tune the spatial and temporal properties of Ca²⁺ signals. They can alter both the amplitude and the recovery time of individual Ca²⁺ transients. These buffers have different properties and expression patterns. For example, calbindin D-28 (CB) and calretinin (CR) are fast buffers, whereas parvalbumin (PV) has much slower binding kinetics and a high affinity for Ca²⁺. These are mobile buffers that increase the diffusional range of Ca²⁺ (REF. 53). Of the Ca²⁺ that enters the cytosol, only a very small proportion ends up being free, because most of it is rapidly bound to the buffers and, to a lesser extent, the effectors (FIG. 1). The Ca¹⁺-BINDING RATIO (K_S) is used to

compare the buffering capacity of cells. Some cells, such as motoneurons and adrenal chromaffin cells, have K_s values of approximately 40, but this can increase to values as high as 2,000 in Purkinje neurons⁵⁴. The low buffering capacity of motoneurons enables them to generate rapid Ca²⁺ signals, but this does make them much more susceptible to excitotoxic stress, which might contribute to motoneuron disease⁵⁴.

The importance of buffers in Ca²⁺ signalling has emerged from studying transgenic animals in which individual buffers have been deleted. When the *PV* gene that encodes *PV* is knocked out, the relaxation of fast-twitch muscles is impaired, but they become fatigue resistant through a remarkable compensatory mechanism that involves an upregulation of the mitochondria, which increases their capacity to sequester Ca²⁺, such that they can partially replace the loss of *PV*⁵⁵. These *PV*^{-t-} animals also have defects in their short-term synaptic plasticity⁵⁶ owing to defects in Ca²⁺ signalling.

Several different effectors, such as troponin C, CaM, synaptotagmin, S100 proteins and the annexins (BOX 1), are responsible for activating different Ca²⁺-sensitive cellular processes. For those processes that respond rapidly, there is a close juxtaposition of the signalling and effector components as occurs for the pre-synaptic and post-synaptic events in neurons (BOX 2, parts b,c). For those processes that operate over longer time scales, such as cell proliferation, Ca²⁺ functions more globally (see below).

Pumps and exchangers. During the course of a typical Ca²⁺ transient, the on reactions are counteracted by the off reactions, during which time various pumps and exchangers remove Ca²⁺ from the cytoplasm (FIG. 1). The pumping mechanisms also have important homeostatic functions in that they maintain the resting level of Ca²⁺ at approximately 100 nM and ensure that the internal stores are kept loaded. Four different pumping mechanisms are responsible for the off reaction — the PLASMA-MEMBRANE Ca¹⁺-ATPASE (PMCA), the Na⁺/Ca⁺ EXCHANGER (NCX), SERCA and the MITOCHONDRIAL UNIPORTER (FIG. 1).

These pumping mechanisms have different thresholds for activity. PMCA and SERCA pumps have lower transport rates but high affinities, which means that they can respond to modest elevations in Ca²⁺ levels and set basal Ca²⁺ levels. The NCX and mitochondrial uniporter have much greater transport rates, and can limit Ca²⁺ transients over a wider dynamic range. For example, mitochondria accumulate Ca²⁺ even when presented with modest nM global Ca²⁺ changes, but the rate of mitochondrial Ca²⁺ uptake is optimal at μ M Ca²⁺ concentrations^{57,58}.

The diverse PMCA, SERCA and NCX molecular toolkit (BOX 1) enables cells to select the combination of off reactions that exactly meets their Ca²⁺-signalling requirements. For example, cells, such as stereocilia, skeletal and cardiac muscle, that generate rapid Ca²⁺ transients have PMCA isoforms (PMCA2a and PMCA3f) that pump at fast rates, whereas cells that produce slower Ca²⁺ transients to activate cell proliferation express PMCA4b that pumps much more slowly⁵⁹.

AUTOSOMAL-DOMINANT POLYCYSTIC KIDNEY DISEASE (ADPKD). A fatal disease that is characterized by the progressive development of fluid-filled cysts in the kidney, liver and pancreas

 Ca^{3} -BINDING RATIO $(K_{s_{s}})$. The ratio between the amount of Ca^{2s} that is bound compared to the Ca^{2s} that is free in the cytosol.

PLASMA-MEMBRANE

Ca*-ATPASE

(PMCA). A pump on the plasma
membrane that couples ATP
hydrolysis to the transport of

Ca*- from cytosolic to
extracellular spaces.

Na¹/Ca²¹ EXCHANGER
(NCX). A plasma-membrane
enzyme that exchanges three
mokes of Na⁴ for one mole of
Ca²⁴, either inward or outward,
depending on the ionic
gradients across the membrane.

MITOCHONDRIAL UNIPORTER A 'channel' that is located in the inner mitochondrial membrane that transports Ca²⁺ from the cytosol into the mitochondrial matrix. Recent studies have highlighted the contribution of a hitherto unrecognized family of Ca²⁺ pumps in regulating Ca²⁺ stores. These Ca²⁺ pumps, which are known as secretory-pathway Ca²⁺-ATPases (SPCAs), are related to SERCAs but have distinct functional properties and cellular roles⁶⁰. It seems that SPCAs might be responsible for Ca²⁺ sequestration into Golgi compartments. The expression of these pumps increases in lactotrophs before parturition, and mutations can lead to HAILEY-HAILEY DISEASE, which indicates that the maintenance of Golgi Ca²⁺ levels by SPCAs is crucial in regulating secretion and cellular contacts.

Spatial and temporal organization of Ca²⁺ signalling In addition to the extensive Ca²⁺-signalling toolkit, another factor that contributes to the versatility of Ca²⁺ signalling is its high degree of spatial and temporal diversity¹. Spatial properties are particularly relevant for rapid responses when components of the on reactions and their downstream effectors are closely associated. This spatial contiguity is less apparent for slower responses such as gene transcription and cell proliferation when Ca²⁺ signals are usually presented in the form of repetitive Ca²⁺ transients and waves.

Spatial aspects of Ca²⁺ signalling. Many Ca²⁺-signalling components are organized into macromolecular complexes (BOX 2) in which Ca²⁺ signalling functions within highly localized environments. These complexes can function as autonomous units, or modules, that can be multiplied, or mixed and matched, to create larger, more diverse signalling systems. For example, the cardiac Ca²⁺-release unit (FIG.3) can be recruited independently of its neighbours to produce graded contractions. Similarly, the individual spines on neurons operate as autonomous Ca²⁺-signalling units, which greatly increases the computational capacity of neurons. For example, individual spines can undergo input-specific, Ca²⁺-dependent synaptic modifications during the process of learning and memory.

Temporal aspects of Ca2+ signalling. Almost all Ca2+-sensitive processes are tuned to respond to Ca2+ transients that are generated by the on/off mechanisms that are summarized in FIG. 1. At the fast end of the scale, for example, synaptic transmission or cardiac contraction, the effector systems respond to pulses within the microsecond to millisecond range. Moving up the timescale, the Ca2+ transients tend to last for longer (seconds to minutes) and the resulting signal usually spreads out as a Ca2+ wave (see below) to reach targets that are distributed throughout the cell. During prolonged stimulation, these Ca2+ transients are repeated to set up regular Ca2+ oscillations that have been implicated in the control of many different processes, such as oocyte activation at fertilization61, growth-cone migration⁶², growth-cone turning⁶³, axonal growth of cortical neurons64, neuronal-cell migration65, development of neurotransmitter phenotypes66, formation of nodules in plant root hairs67, contributing to astrogliosis and epilepsy in neocortical slices68, development of muscle69, release of cytokines from renal epithelial cells70 and the disassembly of adhesive structures during cell migration71.

Cells respond to such oscillations using highly sophisticated mechanisms, including an ability to interpret changes in frequency. Such frequency-modulated Ca²⁺ signalling occurs in many cells (for example, hepatocytes, salivary glands, endothelial cells and smooth muscle cells), in which it can regulate specific responses such as exocytosis⁷², mitochondrial redox state⁷³ and differential gene transcription^{74–78}. The molecular machines that are responsible for decoding frequency-modulated Ca²⁺ signals include CaMKII⁷⁹ and PKC⁸⁰. The function of repetitive Ca²⁺ spiking in differential gene transcription is explored more fully in the section on cardiac hypertrophy.

Ca²+ waves. The Ca²+ signal that makes up Ca²+ oscillations often spreads through the cytoplasm as a regenerative wave¹, and depends on successive rounds of Ca²+ release and diffusion from clusters of Ins(1,4,5)P₃Rs or RYRs that are located on the ER/SR. In the case of Ins(1,4,5)P₃Rs, the channel clusters give rise to local signals that are known as Ca²+ 'puffs', whereas RYRs generate Ca²+ 'sparks'. Typically, these elementary Ca²+ signals produce a modest elevation of the cytosolic Ca²+ concentration (~50–600 nM), with a limited spatial spread (~2–6 μm), and reflect the transient opening of channels (duration of <1 s)^{81,82}. Ca²+ that is released by one channel cluster can diffuse to a neighbouring site and activate it. This SALTATORIC PROPAGATION mechanism allows the initial local signal to trigger global Ca²+ waves and oscillations¹.

In some cases, these waves are highly localized, for example in starburst amacrine cells in the retina, in which localized dendritic signals are used to compute the direction of motion⁸³. In addition to such intracellular waves, information can also spread from cell to cell through intercellular waves as has been described in endocrine cells⁸⁴, the vertebrate gastrula⁸⁵ and the intact perfused liver⁸⁶. In some cases, intercellular waves can cross from one cell type to another as occurs between endothelial and smooth muscle cells⁸⁷.

On and off reactions in cardiac cells. The highly organized cardiac Ca2+-signalling module illustrates several of the important dynamic aspects of Ca2+ signalling, such as amplification, homeostasis, tunnelling and modulation through crosstalk with other signalling pathways (FIG. 3). Activation begins when the cardiac action potential depolarizes the T-TUBULE to open the L-type VOC to introduce a small pulse of trigger Ca2+ a 'sparklet'88 — that diffuses across the junctional zone to stimulate RYR2s (BOX 2, part a) to generate a spark. As the sparklet activates a cluster of 4-6 RYR2s, the spark is much larger and results in a considerable amplification of the initial sparklet. The Ca2+ within the spark then diffuses away from the junctional zone to induce contraction by activating sarcomeres that are situated in the immediate vicinity. During the recovery phase, the off reactions that were described earlier (FIG. 1) begin to remove Ca2+ from the cytoplasm (FIG. 3).

An important feature of Ca²⁺ signalling is homeostasis — cells avoid a net loss or gain of Ca²⁺ by ensuring that the fluxes occurring during the on and off reactions

HAILEY-HAILEY DISEASE
A rare autosomal-dominant skin
disease that is characterized by
disturbed keratinocyte adhesion.

SALTATORIC PROPAGATION
A mechanism by which Ca²⁺
signals leap from one group of
Ca²⁺ channels to the next.

T-TUBULE (transverse tubule). A tubular invagination of the plasma membrane in a muscle fibre, the function of which is to pass the excitation signal from the muscle-cell surface to the sarcomeres, to ensure rapid and synchronous activation.

JUNCTIONAL ZONE
The narrow space that is located
between the T-tubule and
sarcuplasmic reticulum in
cardiac cells in which the process
of excitation—contraction
coupling occurs.

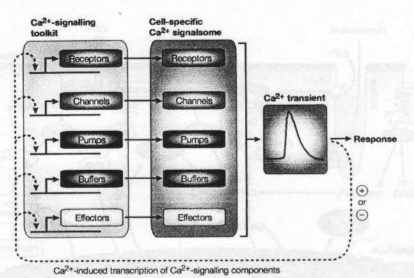


Figure 4 | A calcium-induced calcium-signalling remodelling hypothesis. Cell-specific Ca²⁺ signalsomes that are created by selecting components from the Ca²⁺-signalling toolkit (BOX.1) generate Ca²⁺ transients that are characteristic for each cell type. It is proposed that the Ca²⁺ transient has two functions. In addition to activating cellular responses, it also functions as part of a feedback mechanism to regulate the transcriptional events that are responsible for maintaining the signalsome. This Ca²⁺-dependent transcriptional regulation might have a central role in the compensatory mechanisms that enable cells to adapt to any modifications of their Ca²⁺-signalling systems. It seems that Ca²⁺ can adjust transcriptional activity both positively and negatively. Diseases such as manic depression, hypertension, diabetes and congestive heart failure might develop from a failure of this proposed phenotypic remodelling system.

are always balanced. Homeostasis is particularly important in cardiac cells, in which there is a large circulation of Ca2+ with every heart beat (FIG. 3). For homeostasis to occur, the same amount of Ca2+ that enters through the L-type channel during the on reaction must be removed by the NCX during the off reaction. Likewise, the same amount of Ca2+ that is released by the RYR2s is returned to the SR by the SERCA pump. As the SERCA pumps are distributed over the non-junctional regions of the SR, the newly pumped Ca2+ has to 'tunnel' its way through the SR lumen to reach the junctional region to become available for subsequent release cycles. This part of the recovery process is complicated by the fact that a proportion of the released Ca2+ travels through the mitochondria while en route to the SERCA pump, and this is particularly evident in neonatal cells89. While in transit, Ca2+ within the mitochondrial matrix activates the metabolism that ensures that the formation of ATP is maintained at the required rate to sustain contraction.

The force of contraction can be adjusted by varying the amount of Ca^{2+} that circulates during each on/off cycle. The positive inotropic response that is produced by β -adrenergic stimulation is mediated by cyclic AMP/PKA, which has three main actions on Ca^{2+} signalling. First, it stimulates the L-type VOCs to increase the amount of Ca^{2+} that enters during each action potential. Second, it phosphorylates phospholamban to reduce its inhibitory effect on the SERCA pump, which is then able to increase the luminal Ca^{2+} concentration so that more Ca^{2+} is released from the SR — as described earlier, an increase in the activity of the

SERCA pump is also enhanced by cADPR²³. Third, cAMP/PKA phosphorylates the RYRs, thereby enhancing their ability to release Ca²⁺ to form sparks⁴⁵.

The expression of the individual components of this cardiac Ca²⁺-signalling module, as for Ca²⁺-signalling systems in other cells, is constantly under review. The next section describes how Ca²⁺ has a direct role in remodelling its signalling pathways in both health and disease.

Remodelling Ca²⁺-signalling systems

The full exploitation of the Ca2+-signalling toolkit depends on differential gene transcription to assemble diverse Ca2+-signalling systems. However, such cell-specific Ca2+ SIGNALSOMES are not fixed in stone, in that they are constantly being remodelled to ensure that each cell type continues to deliver the Ca2+ signals that characterize its unique function. It seems that each cell has a signalling blueprint and 'knows' exactly what kind of signal it is expected to deliver. If the spatio-temporal properties of this output signal change because of a loss or defect of a key component, compensatory mechanisms come into play to restore the normal output signal. This remodelling process indicates that there is an element of quality assessment, in that the output of the signalling system is constantly monitored. We propose the hypothesis that Ca2+ itself has an important function in this internal assessment mechanism by remodelling its own signalling pathway (FIG. 4).

Ca2+-induced Ca2+-signalling remodelling. The central pillars of this Ca2+-induced Ca2+-signalling remodelling hypothesis are that Ca2+ is a potent activator of gene transcription2,90-93, and that some of the genes that are activated code for components of the Ca2+-signalling toolkit. The various Ca2+-dependent transcription factors (BOX 1) are activated by different mechanisms. In the case of downstream regulatory element modulator (DREAM), which functions as a repressor, Ca2+ that enters the nucleus functions by removing DREAM from its DNA-binding site91. Ca2+ can also function indirectly through protein kinases (CaMKII and CaMKIV) or protein phosphatases (for example, calcineurin) to alter the phosphorylation state of various transcription factors (FIG. 5). Alternatively, Ca2+ can promote gene transcription by recruiting either the Ras/mitogen-activated protein kinase (MAPK)- or cAMP-signalling pathways.

There is increasing evidence that Ca²⁺ can alter the expression level of Ca²⁺-signalling components such as pumps and channels⁹⁴⁻⁹⁸ (FIG. 4). The Ca²⁺-dependent development of cerebellar granular neurons was marked by an upregulation of Ins(1,4,5)P₃R and two PMCA isoforms, PMCA2 and PMCA3. On the other hand, PMCA4 and the type 2 NCX (NCX2) were rapidly downregulated^{94,96}. Some of these changes, such as the expression of Ins(1,4,5)P₃R, are mediated through the calcineurin/nuclear factor of activated T cells (NFAT) transcriptional cascade^{97,98}.

This ability of Ca²⁺ to regulate its signalling pathways might help to explain the ability of the system to compensate for changes in the activity of individual

INOTROPIC
Affecting the force of cardiac contractions.

SIGNALSOME
The collection of components
that constitute the different
signalling pathways found in
specific cell types.

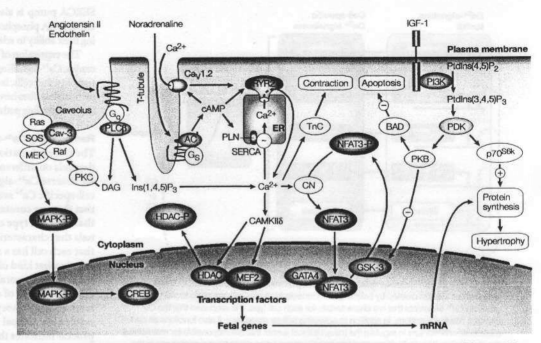


Figure 5 | **Signalling pathways that participate in the control of compensatory hypertrophy.** A key element of this remodelling process is the Ca²⁺ that is released by the cardiac signalling module (FIG.3). This Ca²⁺ signal is enhanced during hypertrophy by the adrenergic pathway that functions through adenylyl cyclase (AC) to increase cyclic AMP (cAMP), which enhances signalling by phosphorylating the entry channel (Ca₂1.2), ryanocline receptor 2 (RYR2) and phospholamban (PLN). Ca²⁺ functions through calcineurin (CN) to dephosphorylate the transcription factor NFAT3, thereby enabling it to enter the nucleus to incluce the transcription of genes, some of which are fetal genes. Hypertrophy can also be enhanced by angiotensin II and endothelin that function through G_a and phospholipase C-β (PLCβ) to generate inositol-1,4,5-trisphosphate (Ins(1,4,5)P₃), which could enhance Ca²⁺, and diacylglycerol (DAG), which seems to function by recruiting the mittogen-activated protein kinase (MAPK)-signalling pathway to stimulate cAMP response element-binding protein (CREB). The phosphatidylinositol 3-kinase (Pl3K)-signalling pathway, which is activated by insulin-like growth factor (IGF-1), contributes to hypertrophy by inhibiting the glycogen synthase kinase 3 (GSK-3) that inactivates NFAT3, by stimulating Bcl-associated death promoter (BAD) to inhibit apoptosis and by stimulating p70^{six} (70-kD ribosomal S6 kinase) to stimulate the expression of the newly transcribed genes. Cav-3, caveolin-3; GATA4, zinc-finger transcription factor; HDAC, histone deacetylase; MEF2, myocyte enhancer factor 2; MEK, MAPK kinase; NFAT, nuclear factor of activated T cells; PDK, phosphoinositide-dependent protein kinase; PKB, protein kinase B; PKC, protein kinase C; PtdIns(4,5)P₂, phosphaticlylinositol-4,5-bisphosphate; PtdIns(3,4,5)P₃, phosphaticlylinositol-3,4,5-trisphosphate; SOS, Son of sevenless; TnC, troponin C.

components⁹⁹. Compensatory changes in Ca²⁺ signalling have also been recorded in cells from mice that carry a null mutation in one copy of the SERCA2 gene, which resembles the mutation that is found in DARIER'S DISEASE. Despite a 40–60% decline in the Ca²⁺ content of the intracellular store, the mouse cardiac cells were still able to generate moderate Ca²⁺ spikes through compensatory changes in the expression levels of phospholamban and NCX¹⁰⁰. Exocrine glands adjusted to the same mutation by upregulating specific PMCA isoforms (PMCA3 and PMCA4) and by adjusting the levels of its Ca²⁺ sensors to increase the sensitivity of exocytosis ¹⁰¹. Each cell type adopted a different strategy to overcome the same mutation, thereby emphasizing the flexibility of Ca²⁺-signalling remodelling systems.

Cardiac hypertrophy. Several important disease states (hypertension, heart disease, diabetes, manic depression and Alzheimer's disease) might result from abnormal remodelling of the Ca²⁺ signalsome. A good example is CONGESTIVE HEART FAILURE, which begins when the heart adapts to stress by increasing in size and by showing an

altered phenotype owing to the expression of neonatal genes. This initial hypertrophy is a compensatory change, because the heart returns to its original phenotype and size if the abnormal inputs are reduced. However, if the stresses persist, this compensated hypertrophy shifts to a state of congestive heart failure, which is more difficult to reverse than compensatory hypertrophy. The phenotypic remodelling that occurs during cardiac hypertrophy and congestive heart failure is controlled by several signalling pathways in which Ca²⁺ has a prominent role (FIG. 5).

This central role of Ca²⁺ signalling in cardiac hypertrophy is apparent from studies on transgenic mice in which the overexpression of genes encoding components of this pathway (FIG. 5), such as the L-type Ca²⁺ channel¹⁰², calcineurin and NFAT¹⁰³, triadin 1 (REF 104), junctin¹⁰⁵, calsequestrin¹⁰⁶ and $G_{q\alpha}$ (REF 107), all result in hypertrophy. Transgenic mice that overexpress calsequestrin have a particularly marked phenotype that is characterized by severe cardiac hypertrophy and decreased depolarization-induced Ca²⁺ release from the SR. The overexpression of calsequestrin resulted in a compensatory decrease in the

DARIER'S DISEASE
An autosomal-dominant skin disorder.

CONGESTIVE HEART FAILURE
A syndrome that is characterized
by the failure of the heart to
maintain the circulation of the
blood adequately.

expression of its binding partners (junctin, triadin and RYR2) within the RYR2 complex (BOX 2, part a)106,108. Conversely, hypertrophy can be prevented by either inactivating components such as Gq (REF. 109) or by promoting the activity of glycogen synthase-3 (GSK-3), which inactivates the Ca2+-sensitive transcription factor NFAT¹¹⁰ (FIG. 5). Ablating phospholamban was able to rescue the hypertrophy that resulted from the overexpression of calsequestrin111, but not hypertrophy caused by overexpressing Goa (REF. 112). The effect of manipulating phospholamban seems to depend on the initial cause of hypertrophy, and this might explain why the effects of ablating this regulator in mice are very different to those seen in humans¹¹³. In humans, hypertrophy and heart disease developed in individuals carrying a mutation that results in the deletion of phospholamban, thereby emphasizing the significance of alterations in Ca2+ signalling as a cause of heart failure in humans113.

A major problem in trying to understand how Ca2+ controls cardiac hypertrophy is the fact that the heart is continuously subjected to large periodic Ca2+ signals (FIG. 3) that flood through the cytoplasm and nucleus every time the heart contracts. Why is it then that cardiac cells subjected to this constant barrage of Ca2+ avoid triggering a hypertrophic response? It has been suggested that the normal functioning heart might not be transcriptionally silent, but might be under constant Ca2+dependent control114. In other words, the transcriptional events that are responsible for regulating the expression of the cardiac signalsome are continuously assessed as proposed earlier for the Ca2+-induced Ca2+-signalling remodelling hypothesis (FIG. 4). If this is the case, then the increase in transcription during hypertrophy might result from subtle alterations in the spatio-temporal properties of the individual Ca2+ spikes. Indeed, a broadening of the Ca2+ transient or an increase in its amplitude has been recorded in cases in which hypertrophy is induced by modifying the levels of proteins that associate with RYR2 (BOX 2, part a), such as triadin 104 or FKBP12.6 (REF. 115), respectively. The hypertrophy that occurred in the FKBP12.6- mice failed to develop in the females, who might be protected against the hypertrophic effect of Ca2+ by female sex hormones such as oestrogen. The failure of phospholamban ablation to prevent hypertrophy112 might be explained by the fact that these transgenic animals had a large increase in the amplitude of the Ca2+ transients. In all of these cases, the rising phase that is responsible for triggering contraction remains the same, but the increase in amplitude or spike broadening enhances the extent of the Ca2+ pulse. These subtle changes in the individual spikes might be integrated over time to stimulate a distinct programme of gene transcription, and some of the genes transcribed will be fetal genes. It is this change in the dynamics of the Ca2+ transient that seems to carry the information that is responsible for inducing hypertrophy.

A different phenotypic remodelling process occurs during the onset of congestive heart failure when the amplitudes of the Ca2+ transients are much reduced owing to a severe downregulation of the Ca2+-signalling system. One of the most noticeable changes is a decline in the activity of the SERCA pump that is due, in part, to the enhanced inhibition through phospholamban116, which results in the severe depletion of the SR store that characterizes congestive heart disease117. One reason for this decline in Ca2+ signalling is a marked downregulation of β-adrenergic-receptor signalling 118. During compensatory hypertrophy, an increased release of catecholamines contributes to the hypertrophic response by activating the cAMP pathway that enhances Ca2+ signalling (FIG. 5). This enhancement of Ca2+ signalling is seriously compromised when the β-adrenergic-signalling system is downregulated during congestive heart failure.

The severity of congestive heart failure might arise when this downregulation of the β-adrenergic-signalling system coincides with a Ca2+-induced Ca2+-signalling remodelling process to switch off the expression of key components such as the SERCA pumps. It is the coincidence of these two regulatory events that might explain the severe disfunction of the Ca2+ uptake and storage capacity, which results in the weak transients that characterize the failing human heart117.

Conclusion

There are numerous Ca2+-signalling systems that are designed to regulate many different cellular processes. This versatility is achieved by the existence of an extensive Ca2+-signalling toolkit that is used to assemble these cell-specific signalsomes that can deliver Ca2+ signals with the spatial and temporal characteristics that are necessary for its many control functions. A major challenge for the future is to determine the differential transcription and expression mechanisms that are responsible for putting together these different signalling pathways. In addition, it will be important to establish the quality-assessment mechanisms that are responsible for maintaining the integrity of these signalling pathways. There is already some indication that Ca2+ itself might function in regulating the stability of its signalling pathways. This Ca2+-dependent qualityassessment mechanism warrants further attention, because there are indications that several pathologies, such as cardiac disease, might develop after the abnormal remodelling of Ca2+ signalling.

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Online links

DATABASES

ms in this article are linked onl LocusLink: http://www.ncbi.nlm.nih.gov/LocusLink/ AKAP | CaBP | calsequestrin | CaMKII | Ins(1,4,5)P,Rs | mGluRs | NMDARs | NCX | phospholamban | PLCB | PMCA | RYRs | SERCA | syntaxin

Swiss-Prot: http://www.expasv.ch/ FKBP12.6 | GKAP | PR130 | PSD95 | SNAP-25 | spinophilin |

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