Selective modulation of subtype III IP$_3$R by Akt regulates ER Ca$^{2+}$ release and apoptosis

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Ca$^{2+}$ efflux from intracellular stores, in particular endoplasmic reticulum (ER), has a fundamental role in modulating different cellular responses. Variations of Ca$^{2+}$ concentration [Ca$^{2+}$]$_c$ works as a trigger, for example, for the secretion of hormones and neurotransmitters, modulation of metabolism and mitotic division in numerous cell types.$^1$ Several extracellular stimuli induce ER depletion, with consequent cytosolic Ca$^{2+}$ increase, through activation of phospholipase C, which leads to the production of a second messenger, inositol 1,4,5-trisphosphate (IP$_3$). IP$_3$ in turn activates an internal membrane receptor, responsible for ER Ca$^{2+}$ release, the IP$_3$ receptor (IP3R). In birds and mammals, three different genes encode for three subtypes of IP$_3$R (IP$_3$R I, II and III), which share high similarity (70–80%) in their primary sequences and are highly conserved in different species,$^10$ which is a substrate for Roderick’s group.$^11$ Activated Akt exerts its pro-survival activity through phosphorylation of target proteins, inhibiting their pro-apoptotic function.$^5$ All three subtypes of IP$_3$R present, in their C-terminal tail, a robust phosphorylation motif RXRXX(S/T), highly conserved in different species,$^{10}$ which is a substrate for Akt kinase activity.$^{10,11}$ Both groups cited, using mutants of type I IP$_3$Rs, declared that IP$_3$R phosphorylation by Akt promotes survival, but only the work of Roderick’s group$^{11}$ connected it to an inhibition of ER Ca$^{2+}$ release. We demonstrated that Akt activation protects cells from apoptosis by strongly reducing Ca$^{2+}$ flux from the ER.$^{12}$ Here, we report a thorough analysis of Akt anti-apoptotic activity through the regulation of IP$_3$R Ca$^{2+}$ channels. Using cells that display different ratios of IP$_3$Rs, we show that Akt
functions specifically on IP₃R isoform III, independently of the presence or absence of isoform I. Finally, our data highlight how the Akt-dependent inhibition of Ca²⁺ transfer from the ER to mitochondria, through IP₃R III, preserves the mitochondrial integrity and protects from Ca²⁺-mediated apoptosis.

Results

We previously reported that Akt protects from Ca²⁺-dependent apoptotic stimuli through inhibition of ER Ca²⁺ release in HeLa cells. To investigate whether Akt might possess a potential functional role on a specific isoform of IP₃R, we used two different cell lines, COS7 and SH-SY 5Y, which lack IP₃R type I and type III, respectively. After confirmation of the effective suitability of our cellular systems (Figure 1a), we measured intracellular calcium homeostasis using aequorin probes, targeted to different subcellular compartments, that is, the cytoplasm and the organelles acting as sources (ER) or targets (mitochondria) of the Ca²⁺ signal. We compared mock-transfected (control) with myristoylated/palmitoylated Akt (m/p-Akt)-overexpressing cells, where m/p-Akt is a plasma membrane-targeted Akt chimera, considered a constitutively active form. In COS7 cells, Akt affects neither the total ER Ca²⁺ content (317.67 ± 15.75 μM control versus 288.33 ± 25.96 μM m/p-Akt) nor the kinetics of Ca²⁺ accumulation. After stimulation with ATP (which functions on Gq-coupled plasma membrane receptors and causes the production of IP₃, thus releasing Ca²⁺ from the ER through the IP₃Rs), Akt-overexpression cells display a very different behavior in the release phase (Figure 1b), with an 86.7 ± 1.9% decrease in the Vₘₙₐₓ of released Ca²⁺ (Vₘₐₓ: 14.32 ± 2.11 μM/s control versus 1.9 ± 0.27 μM/s m/p-Akt). Conversely, in SH-SY 5Y cells, the inhibition of ER Ca²⁺ release mediated by Akt overexpression is negligible (Figure 1c), with no significant differences in the ER plateau values (329.46 ± 11.09 μM control versus 308.25 ± 11.18 μM m/p-Akt), similarly to COS7 cells, and a tiny reduction in the [Ca²⁺]c flow through the IP₃-gated channels (Vₘₐₓ: 19.39 ± 0.97 μM/s control versus 17.49 ± 1.34 μM/s m/p-Akt). We used Carbachol (Chc) as agonist stimulus instead of ATP in SH-SY 5Y cells, because in this cell type, Chc generates higher ER Ca²⁺ release than ATP (data not shown).

The close physical association between ER and mitochondria explains how mitochondria detect the Ca²⁺ flux released by ER very efficiently. Using an aequorin form targeted to the mitochondrial matrix, we measured mitochondrial [Ca²⁺]c in COS7 and SH-SY 5Y cells, to have a further read-out system of Akt activity on IP₃R Ca²⁺-releasing properties. As expected, in COS7 cells, Akt overexpression markedly reduces Ca²⁺ transient in the mitochondrial compartment (11.89 ± 1.31 μM control versus 1.42 ± 0.06 μM m/p-Akt; Figure 1d); this is owing to the IP₃R inhibition described above. Figure 1e shows how, in SH-SY 5Y cells, Akt activity does not alter mitochondrial Ca²⁺ homeostasis in any marked manner, compared with COS7 (peak amplitude: 32.33 ± 4.23 μM control versus 27.34 ± 2.17 μM m/p-Akt), confirming the weak reticular Akt action on Ca²⁺ release in this cell type. To complete the scenario, we analyzed the cytosolic Ca²⁺-kinetics: in Akt-transfected COS7 cells, the [Ca²⁺]c increases evoked by stimulation with ATP are significantly smaller than in controls (peak amplitude: 0.8 ± 0.017 μM versus 1.72 ± 0.027 μM; Figure 1f), whereas in SH-SY 5Y this reduction is almost undetectable (peak amplitude: 1.45 ± 0.051 μM versus 1.83 ± 0.064 μM; Figure 1g).

Having shown the very different effect of Akt expression on the modulation of calcium homeostasis in COS7 and SH-SY 5Y cells, we sought to examine the rate of apoptosis after treatment with a Ca²⁺-mediated apoptotic stimulus. IP₃R is an important molecular component regulating apoptosis induced by a number of different stimuli, and IP₃R-mediated Ca²⁺ release can activate apoptotic pathways by inducing the release of a number of pro-apoptotic factors from mitochondria. As apoptosis inducer we used arachidonic acid (AA), a very specific stimulus that triggers Ca²⁺-dependent cell death. In neuronal and non-neuronal cell lines, AA leads to apoptosis via a mitochondrial-mediated pathway, opening the mitochondrial permeability transition pore, with consequent release of cytochrome c, followed by activation of caspase 3-dependent nuclear condensation and fragmentation. In COS7 cells, Akt overexpression protects from apoptosis induced by AA, with no marked reduction of the amount of cleaved poly-ADP-ribosomal polymerase (PARP) and caspase 3 (Figure 2a, lanes 1 and 4). Moreover, these data were confirmed by direct measurements of caspase activity after application of AA, showing a significant decrease of caspase 3 activity in m/p-Akt-overexpressing cells (Figure 2b). In SH-SY 5Y cells, the Akt anti-apoptotic effect seems to be lost: in fact, no differences were detected both in cleaved PARP levels and in the amount of cleaved caspase 3, comparing control and m/p-Akt-expressing cells after AA treatment (Figure 2c). Similar results are showed in Figure 2d, where AA induces an increase in caspase 3 activity which is mostly comparable between control and m/p-Akt-overexpressing SH-SY 5Y. This assay becomes necessary for a correct visualization of the real caspase 3 activity, because of the difficulty in the detection of cleaved form of the enzyme by immunoblotting in this cellular type.

To assess if the protective role of Akt in COS7 cells may be ascribed to a minor amount of reticular Ca²⁺ released, we analyzed the kinetics of cytosolic Ca²⁺ generated by AA application. Given that the increase in cytosolic [Ca²⁺]c ([Ca²⁺]c) evoked by apoptotic stimuli is in the nanomolar range, the [Ca²⁺]c was measured with the fluorescent indicator Fura-2/AM, as aequorin is not accurate enough to reveal small arises of [Ca²⁺]. We performed a cotransfection (in a 1:1 ratio) using green fluorescent protein with a mitochondrial marker (mtGFP) and m/p-Akt, to identify Akt-positive cells and, therefore, to compare changes in Fura-2/AM 340/380 ratio on the same coverslip. Thus, GFP-positive cells were distinguished from controls by the typical fluorescence emitted upon illumination with blue light. AA addition causes a cytosolic Ca²⁺ increment that is significantly reduced in Akt-expressing cells (Figure 3a, red trace), with a reduction of 47.8%. Parallel experiments were performed in SH-SY 5Y (Figure 3b), but in these cells m/p-Akt overexpression does not alter the amount of cytosolic Ca²⁺ evoked by AA application (blue trace). This result is in perfect agreement with the weak Akt activity at the reticular level (Figure 1c) and its lack of anti-apoptotic function (Figures 2c and d).
Mitochondrial fragmentation is one of early events that occur in apoptotic pathway, it is evolutionarily conserved, and it represents a clear hallmark of mitochondrial perturbation, which precedes effector caspase activation and cell death. We investigated if the Ca\(^{2+}\) transfer from ER to mitochondria promoted by AA might lead to fragmentation of the organelle and if Akt, modulating ER calcium release, might preserve the mitochondrial integrity, minimizing the apoptotic damage. First of all, we used HeLa cells not only because these cells express mostly type I and III IP\(_3\)R, with no detected

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**Figure 1** Akt-modulation of the Ca\(^{2+}\) signal in COS7, but not in SH-SY 5Y cells. (a) IP\(_3\)R levels in COS7, HeLa and SH-SY 5Y cells. HeLa cells were used as reference because they possess both type I and III IP\(_3\)Rs; the increment of p-Akt S473 levels reflects an effective Akt activation after m/p-Akt overexpression. (b) ER Ca\(^{2+}\) measurements in mock-transfected and m/p-Akt-overexpressing COS7 cells. To induce Ca\(^{2+}\) release from ER, cells were challenged with ATP, an agonist that, through interaction with G-protein-coupled receptors, evokes a rapid discharge from IP\(_3\)Rs. Akt activation markedly modulates ER Ca\(^{2+}\) kinetics (red trace), diminishing the amount of Ca\(^{2+}\) released, quantified after analysis of maximum rate of calcium release during agonist stimulation (see bars inside the dotted circle). (c) Experiments analogous to b were carried out in SH-SY 5Y cells. In these cells, the agonist used is Cch. (d) Mitochondrial Ca\(^{2+}\) homeostasis modulation after Akt activation in COS7 cells and (e) in SH-SY 5Y cells. (f) Cytosolic Ca\(^{2+}\) homeostasis in control and m/p-Akt-overexpressing COS7 and (g) SH-SY 5Y cells. Where indicated, cells were stimulated with 100 \(\mu\)M ATP or 500 \(\mu\)M Cch. The bars in d-g indicate the levels of Ca\(^{2+}\) uptake, expressed in percentage, control assumed as 100%. All traces are from single representative experiments and [Ca\(^{2+}\)] are presented as means of at least 20 experiments (± S.E.)
levels of subtype II, but mainly because it is a well-characterized cellular system in which it has been shown that Akt carries out its anti-apoptotic role. To ensure the overexpression of the protein of interest in all observed mitochondrial networks, we co-transfected cells with the indicated vector (empty vector for control cells) and mtGFP in a 3:1 ratio. Figure 4 in toto shows mitochondrial structures of different cells before and 20 min after treatment with AA; this duration of time reflects the maximum peak of cytosolic Ca\(^{2+}\) released by the ER. AA induces a robust mitochondrial fragmentation (Figure 4a), whereas only minor changes in morphology were detected in m/p-Akt-overexpressing HeLa cells (Figure 4b). Analogously, in COS7 cells, AA application leads to fragmentation of the organelle (Figure 4c) and Akt expression preserved the three-dimensional mitochondrial integrity (Figure 4d). On the contrary, on SH-SY 5Y cells, no effect of m/p-Akt was detected, with similar morphological modifications both in control and Akt-expressing cells (Figures 4e and f), once more revealing the Akt inability to exert its ‘usual’ anti-apoptotic activity in this cellular setting. Mathematical description of mitochondrial fragmentation is reported in Figure 4g.

We wanted to explore whether the huge discrepancy in Akt behavior between COS7 and SH-SY 5Y might be due to the IP\(_3\)R isoform III deficiency, which characterizes the latter cellular type. To demonstrate our hypothesis, we used a specific kind of SH-SY 5Y (herein referred as SH 2) that expresses IP\(_3\)R III (Figure 5a). First of all, we analyzed the ER kinetics, with particular attention to the release phase. After agonist induction, m/p-Akt-overexpressing SH 2 cells...
Akt), keeping the ER Ca\(^{2+}\) presence of isoform III of IP3R is the key point of Akt activity.

In this cellular type, Akt shows its greater activity, with a long-lasting \([\text{Ca}^{2+}]_{\text{c}}\) plateau. Interestingly, Akt protection from staurosporine-induced cell death in SH-SY 5Y seems due to the mitochondrial activity of the kinase. Nonetheless, the predominant (93%) endogenous IP3R isoform in COS7 cells has been reported to be type III, with no detectable levels of type I (Figure 1a).

The preferential role of Akt on subtype III may be related to the implications of this isoform in a variety of Ca\(^{2+}\)-dependent physiological processes. For example, it promotes Ca\(^{2+}\)-dependent insulin and somatostatin secretion in pancreatic \(\beta\) and \(\delta\) cells, and it is localized to the apical pole of different types of cells, a region responsible for Ca\(^{2+}\)-signals generation and thus termed ‘trigger zone’. These observations suggest that type III is more prone to initiate global release of intracellular Ca\(^{2+}\) in intact cells. This is further supported by the lack of Ca\(^{2+}\)-dependent negative feedback of IP3R III. Intriguingly, it has been recently reported that the phosphorylation of the receptor by Akt kinase is enhanced in the presence of Ca\(^{2+}\) and, considering the Akt inhibitory effect on ER calcium release, it may represent an intrinsic cellular mechanism to reinstate basal conditions, favoring ER refilling after store depletion due to agonist stimulation.

The in silico analysis of the mitochondrial network revealed no morphological changes were detected in Akt-expressing SH 2 cells, whereas AA induces mitochondrial fission in control cells, indicating how preservation of the mitochondrial integrity under apoptotic stress is required for correct Akt surviving activity.

**Discussion**

In this study, we have used different cell lines expressing different subtypes of IP3R ratio, to understand if Akt may have an IP3R isoform-specific action. Our work demonstrates that Akt preferentially exerts its anti-apoptotic role through inhibition of Ca\(^{2+}\) release by IP3R type III (Figure 6), and SH-SY 5Y cells deficient for this receptor isoform appear insensitive to the typical Akt surviving activity (Figures 2c and d), showing no differences in ER calcium release both after physiological (Figure 1c) and apoptotic stimulation (Figure 3b). As apoptosis inducer, we used AA: AA application causes a progressive release of Ca\(^{2+}\) from intracellular stores, thereby directly causing a \([\text{Ca}^{2+}]_{\text{c}}\) rise and activating capacitative Ca\(^{2+}\) influx, which in turn is responsible for maintaining a long-lasting \([\text{Ca}^{2+}]_{\text{c}}\) plateau.

Interestingly, Akt protection from staurosporine-induced cell death in SH-SY 5Y seems due to the mitochondrial activity of the kinase. Nonetheless, the predominant (93%) endogenous IP3R isoform in COS7 cells has been reported to be type III, with no detectable levels of type I (Figure 1a). In this cellular type, Akt shows its greater activity, with a marked reduction of Ca\(^{2+}\) released by ER after apoptotic stimulus (Figure 3a) and protection from apoptosis (Figures 2a and b). The inhibition of Ca\(^{2+}\) influx after agonist stimulation is more marked than previously observed in HeLa cells, an aspect which may be ascribed to the amount of IP3R III in COS7 cells. Moreover, in Sf9 insect cells Akt seems to phosphorylate type III IP3R more efficiently than type I.

The preferential role of Akt on subtype III may be related to the implications of this isoform in a variety of Ca\(^{2+}\)-dependent physiological processes. For example, it promotes Ca\(^{2+}\)-dependent insulin and somatostatin secretion in pancreatic \(\beta\) and \(\delta\) cells, and it is localized to the apical pole of different types of cells, a region responsible for Ca\(^{2+}\)-signals generation and thus termed ‘trigger zone’. These observations suggest that type III is more prone to initiate global release of intracellular Ca\(^{2+}\) in intact cells. This is further supported by the lack of Ca\(^{2+}\)-dependent negative feedback of IP3R III. Intriguingly, it has been recently reported that the phosphorylation of the receptor by Akt kinase is enhanced in the presence of Ca\(^{2+}\) and, considering the Akt inhibitory effect on ER calcium release, it may represent an intrinsic cellular mechanism to reinstate basal conditions, favoring ER refilling after store depletion due to agonist stimulation.

In a similar way to Akt, cAMP-dependent protein kinase (PKA), another component belonging to the AGC kinase group, is able to bind and phosphorylate all IP3Rs and type III with a certain efficiency. PKA is highly homologous to Akt, sharing ~45% sequence identity with the kinase domain, and rising to ~80% within the ATP site. Subtype III is phosphorylated by PKA at three different serine residues, and, similarly to Akt, a decrease in intracellular Ca\(^{2+}\) release mediated by...
PKA was observed. Conversely, an opposite role of PKA on 
$\text{Ca}^{2+}$ mobilization by IP$_3$R III has been reported, suggesting 
an alternative mechanism to justify the inhibition.

The most important aspect arising from our data refers 
to apoptosis. Akt activation does not protect from 
$\text{Ca}^{2+}$-mediated cell death in cells lacking for IP$_3$R III, a phenomenon

which cannot be considered cell-type dependent, because 
SH-SY 5Y-expressing isoform III showed the typical Akt 
surviving activity (Figure 5). This result can reasonably 
be ascribed to a modification in subtype III-$\text{Ca}^{2+}$ release 
properties, altered by Akt (Figure 5d). As Akt, another 
negative regulator of apoptosis, the Bcl-2 family member

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**Figure 4** Akt preserves the mitochondrial integrity in an IP$_3$R type III-dependent way. (a–f) The mitochondrial network of HeLa (a and b), COS7 (c and d) and SH-SY 5Y 
cells (e and f) was analyzed using confocal microscopy, 36 h post-transfection, before and after treatment with 80 $\mu$M AA for 20 min. Cells were co-transfected in a 3 : 1 ratio 
with an empty vector and mtGFP (left panels: control) or m/p-Akt and mtGFP (right panels: m/p-Akt). Greater magnification of the mitochondrial three-dimensional structure is 
presented in the insets. (g) Frequency distribution of mitochondrial fragmentation: C.F. was analyzed to represent the state of each single mitochondrial object (i.e., C.F. = 0.1 
‘elongated’ mitochondrion, considered not fragmented; C.F. = 1.1 ‘circular’ mitochondrion, considered fragmented). $\nu$, frequency (number of objects)
Bcl-xL seems to be able to interact with all three IP$_3$R isoforms, though isoform-specific effects have been described. The association between Bcl-xL, amount of ER Ca$^{2+}$ released and apoptosis is exclusively effective for type III, because although all three IP$_3$R isoforms provided apoptosis resistance when expressed with Bcl-xL, only the type III channel reduced ER [Ca$^{2+}$].

The role of IP$_3$R III in apoptosis is, currently, partially controversial, and it is described both as an anti- and a pro-apoptotic protein. We agree with the latter hypothesis,
Ca<sup>2+</sup>-dependent apoptotic stimulus evokes Ca<sup>2+</sup> release from the ER, especially through isoform III, with consequent mitochondrial Ca<sup>2+</sup> accumulation, fragmentation of the network and apoptosis. In situations with enhanced Akt activity, a typical condition of many cancers, subtype III is inhibited, therefore decreasing the amount of Ca<sup>2+</sup> accumulated by mitochondria. The mitochondrial network remains undamaged and apoptosis is blocked.

**Figure 6** Model depicting Akt anti-apoptotic activity through regulation of IP<sub>3</sub>R type III-Ca<sup>2+</sup> transfer to mitochondria. During basal condition, when Akt activation is physiological, a Ca<sup>2+</sup>-dependent apoptotic stimulus evokes Ca<sup>2+</sup> release from the ER, especially through isoform III, with consequent mitochondrial Ca<sup>2+</sup> accumulation, fragmentation of the network and apoptosis. In situations with enhanced Akt activity, a typical condition of many cancers, subtype III is inhibited, therefore decreasing the amount of Ca<sup>2+</sup> accumulated by mitochondria. The mitochondrial network remains undamaged and apoptosis is blocked.

and we are strongly convinced that it is a key factor in apoptosis, especially in Ca<sup>2+</sup>-dependent cell death: in fact, siRNA silencing of IP<sub>3</sub>R III antagonizes apoptosis induced by different Ca<sup>2+</sup>-mediated stimuli (P Pinton, unpublished data). Further support to this concept is provided by the evidence that, in CHO cells, subtype III co-localizes more closely with mitochondria, and thus preferentially transmits Ca<sup>2+</sup> signals to these organelles. This observation explains Akt ability to preserve the mitochondrial integrity (Figure 4), delaying Ca<sup>2+</sup> transfer from the ER to mitochondria by inhibition of type III channel gating. The Akt isoform-specific activity completes the molecular pathway describing a new role of promyelocytic leukemia (PML) protein as an apoptosis regulator at the ER–mitochondria interface. PML controls Ca<sup>2+</sup> transfer to mitochondria through the formation of a complex that, besides PML, includes Akt, IP<sub>3</sub>R III and PP2A.

In conclusion, our data collectively suggest the existence of a specific activity of Akt on IP<sub>3</sub>R type III, leading to channel inhibition, diminished Ca<sup>2+</sup> transfer to mitochondria and protection from apoptosis. We suppose that the function of the kinase at the ER–mitochondria level is the early stage of Akt itinerary in promoting cell survival, this concept is also supported by the very recent observation about the reticular localization of mTORC2, the complex responsible for Akt phosphorylation and activation. Thus, the modulation of type III-Ca<sup>2+</sup> release suggests an additional level of cell death regulation mediated by Akt.

**Materials and Methods**

**Cell culture and transfection.** HeLa cells were cultured in Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% fetal calf serum (FCS), l-glutamine and penicillin/streptomycin in 75-cm<sup>2</sup> Falcon flasks. Both kinds of SH-SY 5Y cells (with and without IP<sub>3</sub>R III) were cultured in Ham’s nutrient mixture F12 (HAM’S/F12), supplemented with 10% FCS, 2 mM l-glutamine, penicillin/streptomycin and MEM non-essential amino-acid solution (Sigma-Aldrich, St. Louis, MO, USA), in 75-cm<sup>2</sup> Falcon flasks.

For aequorin experiments, cells were seeded onto 13-mm glass coverslips and allowed to grow to 75% confluence; for Fura-2/AM measurements and mitochondrial morphology analysis, cells were seeded on 24-mm glass coverslip in the same conditions of growth.

For aequorin measurements in HeLa and COS7 cells, transfection with 4 μg of total DNA (3 μg of the indicated expression plasmids and 1 μg of aequorin) transfected with a standard calcium–phosphate procedure. In both kinds of SH-SY 5Y cells, we performed transfection with the same 3 : 1 DNA ratio, using Lipofectamine LTX (Invitrogen-Life Technologies, Grand Island, NY, USA). All measurements were performed 36 h after transfection.

**Aequorin measurements.** Probes used are chimeric aequorins targeted to the ER (erAEQmut), cytosol (cytAEQ) and mitochondria (mtAEQmut). ‘AEQ’ refers to wild-type aequorin, and ‘AEQmut’ refers to a low-affinity D119A mutant of aequorin. For the experiments with cytAEQ and mtAEQmut, cells were incubated with 5 μM coelenterazine for 1–2 h in DMEM supplemented with 1% FCS. A coverslip with transfected cells was placed in a perfused thermostated chamber located in close proximity to a low-noise photomultiplier with a built-in amplifier/discriminator. To reconstitute erAEQmut with high efficiency, the luminal [Ca<sup>2+</sup>] of the ER first had to be reduced. This was achieved by incubating cells for 1 h at 4 °C in Krebs-Ringer buffer (KRB) supplemented with 5 μM coelenterazine, 5 μM Ca<sup>2+</sup> ionophore ionomycin (Sigma-Aldrich) and 600 μM EGTA. After this incubation, cells were extensively washed with KRB supplemented with 2% bovine serum albumin and then transferred to the perfusion chamber. All aequorin measurements were carried out in KRB supplemented with either 1 mM Ca<sub>Cl<sub>2</sub></sub> (cytAEQ and mtAEQmut) or 100 μM EGTA (erAEQmut). Agonist was added to the same medium as specified in figure legends. The experiments were terminated by lysing cells with 100 μM digitonin in a hypotonic Ca<sup>2+</sup>-containing solution (10 mM CaCl<sub>2</sub> in H<sub>2</sub>O), thus discharging the remaining aequorin pool. The output of the discriminator was captured by a Thorn EMI photon-counting board and stored in an IBM-compatible computer for further analyses. The aequorin luminescence data were calibrated offline into [Ca<sup>2+</sup>] values using a computer algorithm based on the Ca<sup>2+</sup> response curve of wild-type and mutant aequorins.
Fura-2/AM measurements. Cytosolic-free [Ca\(^{2+}\)] was evaluated using fluorescent Ca\(^{2+}\) indicator Fura-2/AM (Molecular Probes-Life Technologies, Grand Island, NY, USA). Briefly, cells were incubated in medium supplemented with 2.5 μM Fura-2/AM for 30 min, washed with KR to remove extracellular probe, supplied with preheated KR (supplemented with 1 mM Ca\(^{2+}\)) and placed in a thermostated incubation chamber at 37 °C on the stage of an inverted fluorescence microscope (Zeiss Axiovert 200, Carl Zeiss, Oberkochen, Germany). Dynamic video imaging was performed using MetaFluor software (Universal Imaging Corporation Ltd, Marlow, Buckinghamshire, UK). Fluorescence was measured every 100 ms with the excitation wavelength alternating between 340 and 380 nm and the emission fluorescence being recorded at 510 nm. At the end of the experiment, a region free of cells was selected, and one averaged background frame was collected at each excitation wavelength for background correction.

Caspace 3 assay. Cells were centrifuged (1000 g for 5 min) and washed once in phosphate-buffered saline. The enzcheck caspase 3 assay kit # 2 by Molecular Probes was used for the determination of caspase 3 activity. Lysate (20 μg), resuspended in a final volume of 100 μl, was assayed using Wallac 1420 Victor3 multimark plate reader (Perkin Elmer, Waltham, MA, USA).

Imaging and analysis of mitochondrial morphology. HeLa, Cos7 and SH-SY5Y cells were seeded and transfected with mtGFP as described previously (see Results). Protein expression was allowed for 36 h and then cells were imaged with Nikon Swift Field Confocal (Nikon Instruments Inc., Melville, NY, USA) equipped with CFI Plan Apo VC60XH objective (n.a. 1.4) and an Andor Elements 3.2. Coverslips were placed in an incubated chamber with controlled temperature, CO\(_2\) and humidity and then z-stacks were acquired by 21 planes with 0.6 μm distance, to allow acquisition of the whole cell. After acquisition, images were restored with the Auoautogo 3D blind deconvolution module, installed on NIS-Elements (Nikon Instruments Inc.), using a theochtonous PSF.

After restoration, images were loaded in Imaris 4.0 (Bitplane AG, Zurich, Switzerland), then subtracted of background and used to generate a threshold Elements (Nikon Instruments Inc.), using a theoretical PSF.

Mathematical analysis of mitochondrial frequency distribution, based on calculation of circularity factor (C.F.), was performed on best-focused plane, using the FIJI software (http://fiji.sc/).

Immunoblot. Total cell lysates were prepared in RIPA buffer, likewise separated by SDS-PAGE and the standard immunoblotting procedure was used. Antibodies used were as follows: rabbit -Akt, rabbit -phospho Akt (S473), mouse -PARP and rabbit -Caspace 3 from Cell Signaling (Danvers, MA, USA); rabbit -actin from Sigma-Aldrich; goat -IP\(_{3}\) R I from Santa Cruz Biotechnology (Santa Cruz, CA, USA); and mouse -IP\(_{3}\) R III from BD Biosciences (Franklin Lakes, NJ, USA).

Conflict of Interest

The authors declare no conflict of interest.

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Author Contributions

SM and PP designed research; SM, MM, AB and MB performed all experiments; SM and PP wrote the paper; SM, AR and CG analyzed data.

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