Mitochondrial Participation in the Intracellular Ca²⁺ Network

Donner F. Babcock, James Herrington, Paul C. Goodwin,* Young Bae Park, and Bertil Hille

Department of Physiology & Biophysics, University of Washington, Seattle, Washington 98195-7290; and *Image Analysis Laboratory, Fred Hutchinson Cancer Research Center, Seattle, Washington 98104

Abstract. Calcium can activate mitochondrial metabolism, and the possibility that mitochondrial Ca²⁺ uptake and extrusion modulate free cytosolic $[Ca^{2+}]$ (Ca_c) now has renewed interest. We use whole-cell and perforated patch clamp methods together with rapid local perfusion to introduce probes and inhibitors to rat chromaffin cells, to evoke Ca²⁺ entry, and to monitor Ca²⁺-activated currents that report near-surface [Ca²⁺]. We show that rapid recovery from elevations of Ca_c requires both the mitochondrial Ca²⁺ uniporter and the mitochondrial energization that drives Ca²⁺ uptake through it. Applying imaging and single-cell photometric methods, we find that the probe rhod-2 selectively localizes to mitochondria and uses its responses to quantify mitochondrial free $[Ca^{2+}]$ (Ca_m). The indicated resting Ca_m of 100-200 nM is similar to the resting Ca_c reported by the probes indo-1 and Calcium

THE Ca²⁺ content of mitochondria within resting cells is believed to be low, yet isolated mitochondria can accumulate large quantities of Ca^{2+} when it is provided to them (Lehninger et al., 1967; Carafoli, 1979; Gunter and Pfeiffer, 1990). During the 1970s and 1980s, however, uptake of Ca2+ into mitochondria was often considered to require pathological levels of cytoplasmic Ca²⁺. Because elevated [Ca²⁺] increases the activity of key metabolic enzymes of mitochondria, reversible uptake of Ca²⁺ into mitochondria is proposed to coordinate energy production to cellular needs (McCormack et al., 1990; Gunter et al., 1994; Hajnóczky et al., 1995). Recently, the availability of new optical indicators for Ca²⁺ ions has enabled studies of mitochondrial Ca²⁺ uptake in living cells and tests of possible roles for it. Signals from populations of cells expressing mitochondrially targeted aequorin reported rapid and large transient increases of free intramitochondrial $[Ca^{2+}]$ (Ca_m)¹ and suggested that mitochondria sequester

Green, or its dextran conjugate in the cytoplasm. Simultaneous monitoring of Ca_m and Ca_c at high temporal resolution shows that, although Cam increases less than Ca_c, mitochondrial sequestration of Ca²⁺ is fast and has high capacity. We find that mitochondrial Ca²⁺ uptake limits the rise and underlies the rapid decay of Cac excursions produced by Ca2+ entry or by mobilization of reticular stores. We also find that subsequent export of Ca²⁺ from mitochondria, seen as declining Ca_m, prolongs complete Ca_c recovery and that suppressing export of Ca²⁺, by inhibition of the mitochondrial Na⁺/ Ca²⁺ exchanger, reversibly hastens final recovery of Ca_c. We conclude that mitochondria are active participants in cellular Ca²⁺ signaling, whose unique role is determined by their ability to rapidly accumulate and then release large quantities of Ca²⁺.

Ca²⁺ from locally high microdomains of cytosolic free $[Ca^{2+}]$ (Ca_c) associated with Ca²⁺ entry or with release from intracellular stores (Rizzuto et al., 1992, 1993, 1994; Rutter et al., 1993; Lawrie et al., 1996). In complementary studies, inhibitors that should block mitochondrial Ca²⁺ uptake severely compromised clearance of cytoplasmic Ca²⁺ after an imposed elevation, indicating that accumulation by mitochondria also is important for cellular Ca²⁺ homeostasis (Thayer and Miller, 1990; Friel and Tsien, 1994; Werth and Thayer, 1994; White and Reynolds, 1995; Herrington et al., 1996; Park et al., 1996). Comparing the rates of Cac decay after selective blockade of known mechanisms for cellular Ca2+ sequestration and extrusion showed that mitochondrial uptake (defined pharmacologically) accounts for as much as 70% of cytosolic Ca²⁺ removed during the initial rapid phase of recovery from large imposed Ca²⁺ loads in chromaffin cells (Herrington et al., 1996).

Here we introduce two fluorescent Ca^{2+} probes, one into the cytosol and the other into mitochondria of single rat chromaffin cells. Simultaneous optical monitoring demonstrates that even modest elevations of Ca_c produce rapid transient redistributions of Ca^{2+} from the cytosol into mitochondria. The mitochondrial Ca^{2+} uptake shapes and dampens the elevations of Ca_c during stimulation, and subsequent Ca^{2+} export from mitochondria delays com-

Please address all correspondence to B. Hille, Department of Physiology & Biophysics, G424 Health Sciences Building, University of Washington, Box 357290, Seattle, WA 98195-7290. Tel.: (206) 543-8639. Fax: (206) 685-0619. E-Mail: hille@u.washington.edu

^{1.} Abbreviations used in this paper: Ca_c , free cytosolic $[Ca^{2+}]$; Ca_m , free intramitochondrial $[Ca^{2+}]$; CGP-37157, 7-chloro-3,5-dihydro-5-phenyl-1H-4,1-benzothiazepine-2-on; BHQ, 2,5 di(*t*-butyl)-1,4-hydroxyquinone; CCCP, carbonyl cyanide *m*-chlorophenylhydrazone.

plete recovery of Ca_c and may facilitate refilling of intracellular stores. Cellular Ca^{2+} signaling may best be considered as an interactive, multicompartmental network in which mitochondria are active participants.

Materials and Methods

All experiments were performed at a room temperature of 24–27°C. Results are reported as mean \pm SEM.

Chemicals

CGP-37157 (7-chloro-3,5-dihydro-5-phenyl-1H-4,1-benzothiazepine-2-on) was the generous gift of Dr. Alain DePover, CIBA-GEIGY (Basel). Dyes were from Molecular Probes (Eugene, OR); ionomycin and BHQ (2,5 di(*t*-butyl)-1,4-hydroxyquinone) from Calbiochem (La Jolla, CA); CCCP (carbonyl cyanide *m*-chlorophenylhydrazone) and all other reagents from Sigma Chem. Co. (St. Louis, MO).

Cells and Media

Adrenal chromaffin cells were prepared as described previously (Herrington et al., 1996; Park et al., 1996) from \sim 300 g male rats. Briefly, adrenal medullae were dissected and digested 50–70 min at 37°C with (mg/ml in modified Hanks' solution): 1.2 collagenase D; 0.4 trypsii; 0.1 DNase type I. Dispersed cells were plated on \sim 5-mm square coverglass slips coated with poly-t-lysine and laminin, then maintained 1–4 d as primary cultures in a fortified DMEM medium.

Patch Clamp Procedures

Voltage clamp recordings used an Axopatch 1C amplifier (Axon Instruments, Foster City, CA) and the BASIC-FASTLAB analysis system (Indec Systems, Capitola, CA). The standard voltage protocol was a 0.05-1 s step to +10 mV from a holding potential of -80 mV. Tail currents in SK-type Ca²⁺-activated K-channels were recorded at a holding potential of -60 mV (Park et al., 1996).

For the whole-cell configuration of the patch clamp technique (Hamill et al., 1981), we usually used a standard pipette solution (mM): 75 Cs₂SO₄; 15 CsCl; 6.5 NaCl; 2.5 Na pyruvate; 2.5 malic acid; 1 NaH₂PO₄; 1 MgSO₄; 50 Hepes; 5 MgATP; 0.3 Tris GTP; 0.5 Tris cAMP; 0.1 leupeptin; pH to 7.3 with CsOH. In some cases, 78 K₂SO₄ and 10 KCl replaced 75 Cs₂SO₄ and 15 CsCl. Perforated-patch recording (Horn and Marty, 1988) also used methods detailed previously (Park et al., 1996), and a simplified pipette solution: 111 Cs₂SO₄; 6 CsCl; 10 NaCl; 5 MgSO₄; 20 Hepes; pH to 7.4 with CsOH (where noted, Cs⁺ salts were replaced with K⁺). Briefly, the tips of patch pipettes were filled by dipping for 1–3 s in simplified pipette solution, then backfilled with the same solution supplemented with 480 µg amphotericin/ml.

The standard external solution used during seal formation contained: 150 NaCl, 2.5 KCl, 2 mM CaCl₂, 1 MgCl₂, 10 Hepes, and 20 mM glucose, pH 7.4. For perforated-patch recordings external solutions also contained 2.5 mM Na pyruvate and 2.5 mM malic acid to optimize mitochondrial function (Villalba et al., 1994). For depolarization-evoked Ca^{2+} entry, the external CaCl₂ was increased to 10 mM.

Activation of voltage-dependent Ca²⁺ channels by depolarization provides precisely timed, but imperfectly predictable control of Ca²⁺ entry, as judged by the peak Ca_c produced. The degree of buffering by introduced dyes and chelators, and the extent of time-dependent channel 'rundown' (more severe in whole-cell- than in perforated-patch-recordings), and previous stimulus history all contribute to this variability. For these reasons, the durations of depolarizing stimuli often were adjusted upwards during the course of an experiment in an attempt to produce similar Ca_c challenges. In the interpretation of these experiments, differences in peak Ca_c reflect mainly imperfect compensation for this rundown.

Dye Loading and Photometry

Rhod-2-AM (0.2 mM) and Calcium GreenTM-AM (10 mM) were dispensed from DMSO stocks, dispersed in 10% Pluronic 147, and diluted to 1 and 12 μ M in 0.25 ml standard external solution. A single coverslip of cells was immediately added, incubated 35–50 min at 22–25°C in the dark, then transferred to an examination chamber containing 2–4 ml fresh buffer. Indo-1 (100 μ M), or either rhod-2 or Calcium Green dextran (3,000 M_r)

at 10 μ M together with 90 μ M BAPTA, were introduced by inclusion in the whole-cell pipette solution. All photometric measurements were performed on an inverted microscope using a 1.3 NA 40× oil objective (Nikon), an attenuated (1.0 NDF) 100 W Hg source, and paired photon-counting detectors (Hamamatsu). An electronic shutter (Uniblitz) restricted excitation to the sampling interval (50–100 ms; duty cycles of 0.5% in normal and 50% in fast mode). Pinhole diaphragms limited the excitation and emission fields to ~25 μ m diameter regions. Indo-1 was excited at 365 \pm 10 nm and detected at 405 \pm 35 and 500 \pm 40 nm. A filter cube with dual band excitation (490 \pm 15 and 560 \pm 15 nm) and dual band emission (530 \pm 15 and 650 \pm 75 nm) (XF-52; Omega Optical) allowed simultaneous excitation of Calcium Green and rhod-2. Detection at 525 \pm 10 and 580 \pm 30 nm was accomplished with additional interference filters separated by a 540nm dichroic mirror.

Calibration of Indo-1, Rhod-2, and Calcium Green Signals

The ratiometric probe indo-1 was calibrated as detailed previously (Herrington et al., 1996; Park et al., 1996). Briefly, the ratio R (F_{405}/F_{500}) of background-corrected signals was applied to the standard calibration equation (Grynkiewicz et al., 1985):

$$[Ca^{2+}] = K^* (R - R_{min}) / (R_{max} - R)$$

where R_{min} (0.47), R_{max} (7.15), and K* (2540 nM) were determined empirically from cells dialyzed with pipette solutions containing 50 mM EGTA, 15 mM CaCl₂, or 20 mM EGTA and 15 mM CaCl₂ (calculated free [Ca²⁺] of 251 nM), respectively.

Fluorescence signals from the single-wavelength indicators rhod-2 and Calcium Green were corrected first for autofluorescence observed with nonloaded cells (<5% of total signal), then for the spillover of fluorescence from Calcium Green (22–48%) or its dextran conjugate (10–15%) into the long-wavelength (rhod-2) detector channel, as determined from cells loaded only with these probes. The extent of spillover increased with age of the dual pass excitation filter set, presumably as a consequence of deterioration. Spillover of fluorescence from rhod-2 into the short wavelength (Calcium Green) channel was negligible. Compensation also was applied for time-dependent loss of signal (primarily photobleaching) as determined from exponential fits of resting-state fluorescence before and after recovery from stimulus (mean rate constants: rhod-2, 4.89 ± 0.10 × 10^{-4} s⁻¹; Calcium Green, 5.80 ± 0.17×10^{-4} s⁻¹). The corrected signals (F) were converted to [Ca²⁺] using the simplified calibration equation:

$$[\operatorname{Ca}^{2^+}] = K_{d} (F - F_{free}) / (F_{CaSat} - F)$$

where the K_d determined in vitro for rhod-2 (500 nM) or Calcium Green (240 nM; Eberhard and Erne, 1991) was assumed to apply, and F_{free} and F_{CaSat} were calculated from postexperimental responses of each cell as follows. Cells were sequentially perfused with a Mg²⁺-free saline containing 10 mM CaCl₂ and 10 μ M ionomycin, then with the same solution in which 2 mM MnCl₂ replaced CaCl₂.

These treatments elicited fluorescence in an average ratio (F_{MnSat}/F_{CaSat}) of 0.75 ± 0.05 (n = 11) and 0.78 ± 0.04 (n = 9) for Calcium Green and its dextran conjugate, indistinguishable from F_{MnSat}/F_{CaSat} ratios determined in vitro in a 150 mM KCl, 1 mM MgCl₂, pH 7.4 medium, containing 1 μ M dye and 10 mM CaCl₂ or MnCl₂. Compared with dye in the same medium containing 10 mM EGTA, Ca²⁺-free:Ca²⁺-saturated fluorescence (F_{free}/F_{CaSat}) was 0.091. Assuming this ratio also applies in vivo, cellular F_{free} was calculated from the observed F_{CaSat} elicited by ionophore.

lated from the observed F_{CaSat} elicited by ionophore. For the K⁺ salt of rhod-2, $F_{free}F_{CaSat}$ determined in vitro was 0.068. When the rhod-2 salt was applied to the cytosol by whole cell dialysis, the F_{MnSat}/F_{CaSat} ratio (0.29 \pm 0.01; n = 3) agreed well with that determined fluorimetrically for rhod-2 in vitro (0.24). F_{MnSat}/F_{CaSat} also measured in vitro was unchanged from pH 7-8.2 and altered little or not at all by variation of ionic strength (0.1-0.2 M KCl), viscosity (0-2 M sucrose), or dielectric constant (0-10% ethanol) that might be encountered intramitochondrially. However, for rhod-2 introduced into mitochondria by AM-loading, F_{MnSat}/F_{CaSat} was much larger either before (0.50 ± 0.13; n = 14) or after $(0.68 \pm 0.11; n = 4)$ whole cell dialysis, presumably indicating that the much higher affinity (>100-fold) of rhod-2 for Mn^{2+} than for Ca^{2+} and the greater efficacy of the ionophore in Mn²⁺ transport (Erdahl et al., 1996) allowed full saturation of the compartmentalized probe with Mn²⁺ but not with Ca^{2+} . Therefore, cellular F_{free} and F_{CaSat} were both calculated from the F_{MnSat} observed for rhod-2AM loaded, ionophore-treated cells, using the F_{MnSat}/F_{CaSat} and F_{free}/F_{CaSat} ratios observed in vitro for rhod-2. These determined ratios (and the K_d for Ca²⁺ used here) agree with the specifications now provided by the manufacturer, but differ from those reported in the original description of rhod-2 (Minta et al., 1989), presumably due to removal of a Ca^{2+} -insensitive contaminant present in the initial preparations of this probe.

Imaging

Images were collected on a Deltavision SA3.1 wide-field deconvolution microscope system (Applied Precision, Inc., Issaquah, WA) using a Nikon PlanApo 60× 1.4 NA objective on an Olympus IMT-2 inverted microscope with a Photometrics PXL-2 cooled CCD camera containing a Kodak KAF1400 chip. Images were collected with an intermediate magnification of 1.5×. Binning was used to increase signal-to-noise. For every 2-s time point, four optical sections were collected at 0.2 μ m spacing each with 0.3-s exposure. Final voxel resolution was 0.157 μ m (planar) × 0.200 μ m (axial). Images of each z-axis series were corrected for photobleaching and variations in lamp intensity; only intensity corrections were applied between time points. The deconvolution reconstruction algorithms of Agard and Sedat (Agard et al., 1989; Hiraoka et al., 1990, 1991; Chen et al., 1995) were applied to each z-series using an optical transform function determined empirically from a 0.1- μ m bead.

Results

We begin with the Ca²⁺-sensitive dye rhod-2 as a probe of Ca_m. Extrusion of protons during electron transport creates a large inside-negative potential difference $(\Delta \psi_m)$ across the inner mitochondrial membrane that is subsequently harnessed by the ATP synthase in the production of ATP. This negative $\Delta \psi_m$ has been exploited in the past to label energized mitochondria with membrane-permeant cations like Rhodamine 123 (Johnson et al., 1981). Similarly, the permeant cation rhod-2 acetoxymethyl ester (rhod-2AM) should also accumulate preferentially in mitochondria where it should become trapped when subsequent esterolysis generates zwitterionic rhod-2 (Minta et al., 1989). Unlike the membrane permeant Rhodamine 123, the trapped rhod-2 should not be sensitive to subsequent changes in mitochondrial membrane potential. The following experiments confirm these predictions for rat chromaffin cells.

Rhod-2 Localizes to Mitochondria and Responds to Ca^{2+} *Entry*

To verify compartmentalization of dye, cells loaded with rhod-2AM were visualized by fluorescence deconvolution imaging. In Fig. 1, rhod-2 fluorescence was found primarily in punctate and filamentous extranuclear regions, consistent with a mitochondrial localization, and in 1-3 small $(\sim 1 \ \mu m)$ intranuclear bodies, tentatively identified as nucleoli. With long exposure images, the characteristic vermiform mitochondrial morphology (Loew et al., 1995) was more apparent (Fig. 1 E). Pharmacological validation of the mitochondrial localization of rhod-2 is found in the following sections below. When the neutral ester of Calcium Green was coloaded in the same cell together with cationic rhod-2AM, the Calcium Green fluorescence showed a more pancellular distribution (Fig. 1 F), consistent with a localization in the cytosol and nucleus as well as some additional association with mitochondria.

Calcium entry induced by KCl depolarization increased the rhod-2 fluorescence within the presumptive mitochondria (Fig. 1 *B*). The fluorescence returned to baseline levels within a minute after the depolarization (Fig. 1, *C* and *D*). No fluorescence increase occurred during application of KCl in the nominal absence of external Ca^{2+} (not

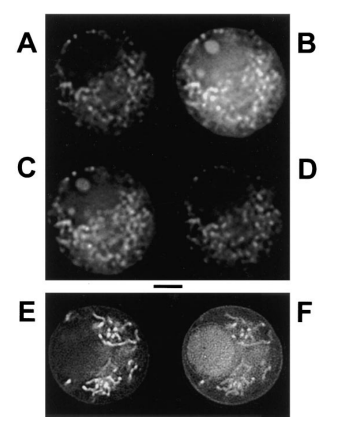


Figure 1. Rhod-2 images of chromaffin cell responses to depolarization and coimages of rhod-2 with Calcium Green. (A–D) Fluorescence deconvolution images of a single cell after loading with AM ester of rhod-2. Images were collected before, during, ~4 s after, and ~1 min after a 10-s, fast local perfusion with a depolarizing (70 mM KCl) medium. (E–F) After coloading with AM esters of rhod-2 and Calcium Green, images were collected from a central plane of a single cell not subjected to stimulus, using excitation and emission filters for rhod-2 in E and then for Calcium Green in F. Scale bar represents 4 µm.

shown). Hence, rhod-2 reports a transient rise of mitochondrial calcium following depolarization-induced Ca^{2+} entry. This is the phenomenon we wish to study.

Rapid Clearance of Cytosolic Ca^{2+} Requires Energized Mitochondria and the Ca^{2+} Uniporter

Uptake of Ca2+ by isolated mitochondria involves electrophoretic transport through a Ca²⁺ uniporter driven by the mitochondrial membrane potential $\Delta \psi_m$ (Gunter and Pfeiffer, 1990; Gunter et al., 1994; but also see Sparagna et al., 1995). We now show that Ca^{2+} uptake by mitochondria within the cell also depends on the negative $\Delta \psi_m$ and the uniporter. We perturb mitochondrial function with four agents, oligomycin, cyanide, carbonyl cyanide m-chlorophenylhydrazone (CCCP), and ruthenium red as illustrated in Fig. 2. Oligomycin blocks mitochondrial ATP production by direct inhibition of the ATP synthase but should not decrease $\Delta \psi_m$ by itself. Cyanide blocks electron transport but should allow maintenance of $\Delta \psi_m$ by an alternate route, the reversal of the ATP synthase fueled by cytoplasmic ATP. However, oligomycin and cyanide in combination block both routes for maintaining $\Delta \psi_m$ and

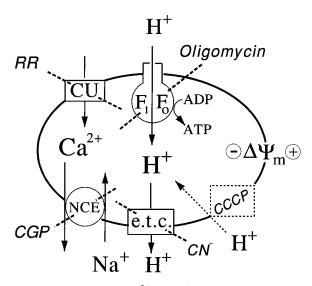


Figure 2. Pathways of Ca²⁺ and H⁺ transport in the inner mitochondrial membrane and diagnostic inhibitors. Abbreviations: CU, Ca²⁺ uniporter inhibited by ruthenium red, RR; F₁F₀, ATP synthase inhibited by oligomycin; NCE, Na⁺-Ca²⁺ exchanger inhibited by CGP-37157, CGP; e.t.c., electron transport chain inhibited by cyanide; CCCP, a protonophore that collapses the mitochondrial membrane potential, $\Delta \psi_m$.

should depolarize mitochondria. Likewise, the ionophore CCCP collapses $\Delta \psi_m$ by making membranes permeable to protons. Finally, ruthenium red blocks the Ca²⁺ uniporter directly.

The first of these experiments (Fig. 3) showed that CCCP and ruthenium red block rapid Ca²⁺ clearance. Ca²⁺ entry into the cell was induced by a step depolarization of the plasma membrane, and removal of cytosolic Ca^{2+} was monitored with the Ca^{2+} probe indo-1 introduced directly into the cytoplasm from a whole-cell pipette. In Fig. 3 A, spatially averaged Ca_c rose to $>4 \mu$ M during a brief depolarizing voltage step (first arrow) and then decayed within a few seconds to submicromolar levels. As reported previously (Herrington et al., 1996), addition of CCCP produced a small transient Ca_c increase of variable duration and size apparently dependent on mitochondrial Ca²⁺ content at the time of addition. CCCP also greatly slowed the recovery of Ca_c after a second depolarization. When this experiment was repeated with other cells internally dialyzed with ruthenium red to block the uniporter, the initial decay was already slowed even without the addition of CCCP (Fig. 3 B). Subsequent treatment with CCCP had no apparent further effect on recovery from Ca^{2+} entry (n = 4). These experiments show that mitochondria cannot clear Ca²⁺ rapidly from the cytoplasm when the $\Delta \psi_m$ is collapsed or the Ca²⁺ uniporter is blocked.

The second set of experiments (Fig. 3, *C* and *D*) tested the ability of oligomycin and cyanide to block Ca^{2+} clearance. Ca^{2+} entry was again induced by a brief depolarization, but here clearance of submembrane cytosolic Ca^{2+} was monitored electrophysiologically by the decay (tail) of a Ca^{2+} -activated K⁺ current that is prominent in rat chromaffin cells (SK current: Neely and Lingle, 1992*a*; Park, 1994). This method requires neither cell dialysis nor intro-

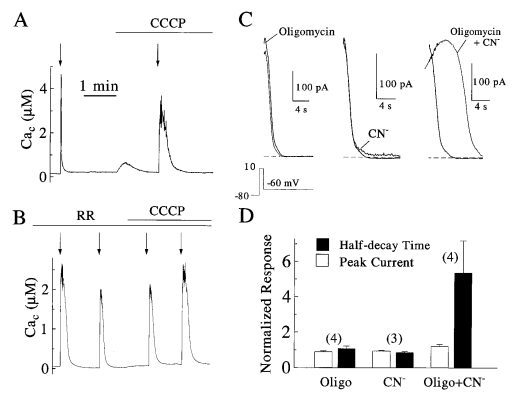
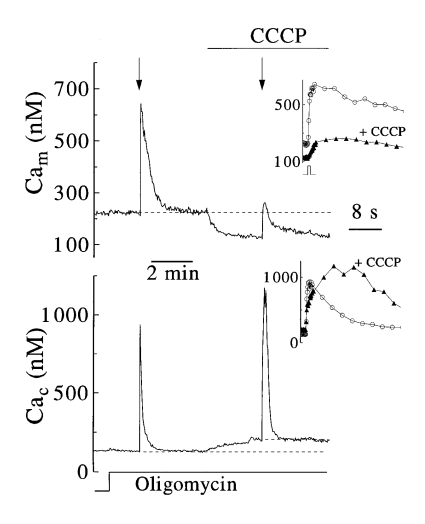


Figure 3. Prolongation of Ca²⁺ clearance by blockade of mitochondrial energization or of the mitochondrial (A-B)uniporter. Time course of cytoplasmic [Ca²⁺] monitored with indo-1 (100 μM) introduced into the cytosol by inclusion in the pipette solution alone (A) or, in a different cell, together with 2 µM RR (B). Beginning 3-5 min after the wholecell patch-clamp configuration was established, dual emission fluorescence signals were recorded. One or two 500-600-ms depolarizing pulses were applied (arrows) while each cell was perfused with control medium alone. Another one or two pulses were applied after 2 µM CCCP was included in the perfusion medium. (C-D) Effect of mitochondrial inhibitors on clearance of submembrane Ca_c as measured by the decay (tail) of a

 Ca^{2+} -activated K⁺ current. Tail currents were recorded in perforated-patch configuration in response to 1-s depolarizations to 0 mV applied every 2 min (Park, 1996). After 9.5 min, the perfusion medium was supplemented with 3 μ M oligomycin or 5 mM NaCN and at 11.5 min, with both. (*D*) Peak amplitude and half-decay times of currents observed after treatment were normalized to those obtained in the preceding control stimulus applied to the same cell. The numbers of cells examined are shown in parentheses.



duction of dyes into the cytoplasm and thus provides a useful, non-invasive indication of $[Ca^{2+}]$ at the plasma membrane. As in previous work (Park et al., 1996), the K⁺ current that was activated by Ca^{2+} entry during a 1-s depolarization then declined with a half-time of 2–7 s as the $[Ca^{2+}]$ fell and the SK channels closed. The decay was not slowed by oligomycin or cyanide alone but was prolonged greatly by the combination of these agents. Blockade of both available routes for mitochondrial energization increased the half-time of current decay 5.4 ± 1.8-fold (n = 4), without appreciably affecting the peak amplitude. These experiments without cell dialysis show again that rapid clearance of cytoplasmic Ca^{2+} depends on the existence of a negative $\Delta \psi_m$ and not, in the short term, on the ongoing oxidative production of ATP.

Monitoring of the CCCP-sensitive Mitochondrial Compartment

Using compartmentalized dyes, we can now verify the assumption that CCCP slows Ca_c clearance by preventing uptake of Ca^{2+} into mitochondria. We have seen that coloading of chromaffin cells by coincubation with AM esters compartmentalizes rhod-2 into mitochondria and disperses Calcium Green throughout the cell. Signals from the two dyes were followed at high time resolution using photometry and a filter cube designed for simultaneous dual-wavelength excitation and dual-wavelength detection.

Figure 4. Reciprocal actions of CCCP on mitochondrial and cytosolic Ca²⁺. Fluorescence signals from rhod-2 and Calcium Green, coloaded as AM esters in a preliminary incubation, were recorded simultaneously from a single cell, beginning 3-5 min after the perforated patch clamp configuration was established. The perfusion medium then was supplemented with 10 mM Ca^{2+} and 3 μM oligomycin and then with 2 µM CCCP as indicated. At the arrows 500-ms depolarizations were applied. A postexperimental calibration procedure (see Materials and Methods) provided parameters for conversion of rhod-2 and Calcium Green fluorescence to the indicated mitochondrial and cytosolic free Ca2+ concentrations (Ca_m and Ca_c, respectively). The results shown are representative of four similar experiments. (Insets) The initial portions of Ca_m and Ca_c responses are aligned in time to the depolarizing stimulus.

After sequential corrections for autofluorescence, signal cross-talk, and photobleaching, Ca_m and global free [Ca²⁺] were calculated from rhod-2 and Calcium Green fluorescence by application of calibration parameters determined for each cell examined. Voltage-clamp depolarizations controlled Ca²⁺ entry, and inhibitors were applied and removed rapidly by fast, local perfusion. We applied CCCP in the presence of oligomycin to prevent any accelerated consumption of cellular ATP by reverse-mode operation of the ATP synthase in proton-permeant mitochondria (Budd and Nichols, 1996; Park et al., 1996). As expected, oligomycin alone did not affect Ca_m or Ca_c of resting cells (Fig. 4).

The rapid rise and decline of $[Ca^{2+}]$ calculated from the Calcium Green signal during and following Ca^{2+} entry (Fig. 4, *lower traces*) resemble the Ca_c responses reported from indo-1 applied directly in the cytosol (Fig. 3 A). Other common features include a delayed final return to the resting level (Figs. 4, 6, and 8) and a small $[Ca^{2+}]$ increase when CCCP was applied late in the recovery from brief Ca^{2+} entry (Fig. 4). Calcium Green and indo-1 signals also both report that CCCP slows the initial clearance of Ca_c yet hastens the final recovery to resting levels. These similarities in the signals from the two probes confirm the usefulness of AM-loaded Calcium Green to measure Ca_c . We note, however, that in the experiments of Fig. 3, A and B and in previous studies using pipette-loaded indo-1 (Herrington et al., 1996), resting Ca_c was

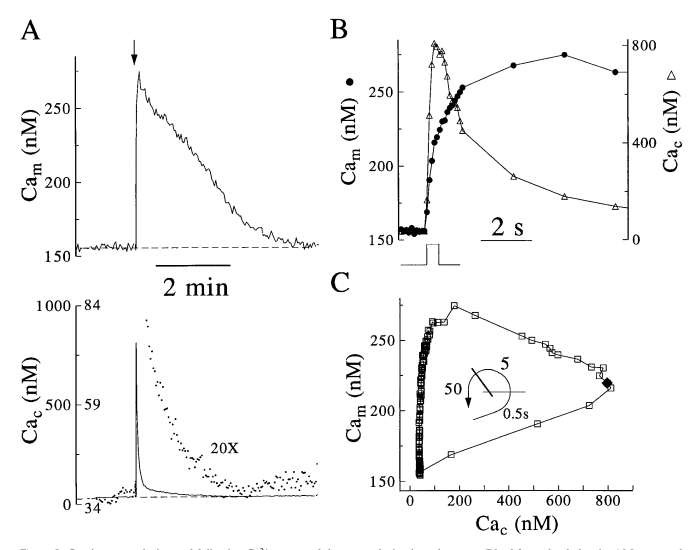


Figure 5. Ca_m increases during and following Ca^{2+} entry and decreases during late clearance. Rhod-2 was loaded as its AM ester and Calcium Green Dextran by whole cell dialysis. Calcium Green fluorescence observed in the on-cell patch clamp configuration was subtracted before calibration of signals simultaneously recorded from a single cell 5 min after the whole-cell configuration was established. (*A*) Ca_m and Ca_c responses to a 500-ms depolarization applied at the arrow. The final portion of Ca_c recovery is shown (*dots*) on a 20× expanded scale. (*B*) The initial portions of Ca_m (*open circle*) and Ca_c (*open triangle*) traces from *A* are aligned in time to the depolarizing stimulus. (*C*) The relationship of Ca_m to Ca_c during this response to stimulus and subsequent recovery; a closed diamond marks the Ca_m and Ca_c values at the termination of stimulus. The curved arrow indicates the approximate nonlinear time course; the lines divide elapsed time into segments of ~0.5, 5, and 50 s duration.

lower (by 100–200 nM) and peak Ca_c evoked by Ca^{2+} entry usually was higher than the values reported here by Calcium Green loaded as its AM ester and examined in perforated patch protocols. It is not yet clear whether these differences reflect alterations in the extent of Ca^{2+} entry, in cytosolic buffering or clearance mechanisms (all of which may be affected by the dye loading and patch clamp procedures employed), or by differences in dye localization and inaccuracies in calibration.

In comparison to the quick changes of Ca_c during and after brief plasma membrane depolarizations, rhod-2 reported a more limited but far more sustained elevation of Ca_m (Fig. 4, *insets*). Subsequent addition of CCCP caused a simultaneous fall in Ca_m and rise in Ca_c (Fig. 4; n = 4), presumably reflecting release of Ca^{2+} from depolarized mitochondria. CCCP also strongly attenuated the rise of Ca_m in response to elevation of Ca_c . The small residual signal may result from incomplete inhibition by CCCP, from incomplete compensation for crossover of the Calcium Green signal, or from rhod-2 reporting from a nonmitochondrial compartment.

Redistribution of Ca^{2+} during and after Ca^{2+} Entry

We further refined the compartmentation of the two dyes by using the patch pipette in whole-cell rather than perforated patch mode to allow introduction of the Calcium Green as a dextran conjugate in the pipette solution. This strategy should restrict Calcium Green to the cytosol and also remove any cytosolic rhod-2 and remaining ester by dialysis. Little or no loss of rhod-2 fluorescence occurred during entry of Calcium Green (not shown), confirming the extensive compartmentation of this dye indicated by imaging

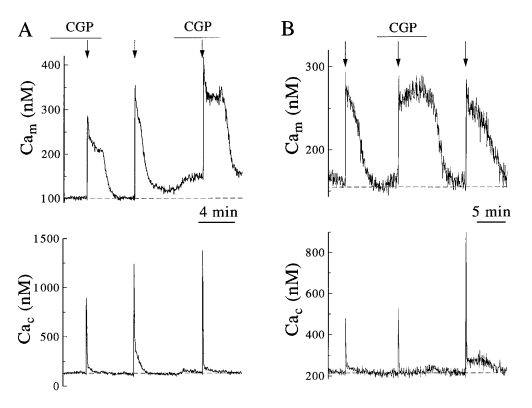


Figure 6. Reversible elimination of the sustained phase of Ca_c recovery by blockade of mitochondrial Ca2+ extrusion. Cells were coloaded with rhod-2 and Calcium Green esters, examined in the perforated patch clamp configuration, and the data analyzed as in Fig. 4. Responses are representative of five similar experiments. (A) Arrows mark the application of sequential depolarizations of 1.5, 2.0, and 2.5 s duration, applied to a single cell in the presence or absence of perfusion with 10 µM CGP-37157 as indicated. (B) Another cell received 0.5, 1.0, and 1.5 s stimuli before, during, and after application of inhibitor.

in Fig. 1. Subsequent responses of the two dye signals to Ca^{2+} entry evoked by depolarizations in the presence and absence (Fig. 5) of CCCP were qualitatively similar to those in Fig. 4. Because removal of CCCP restored Ca_m responses to Ca^{2+} entry (not shown), mitochondrial depolarization apparently does not result in artifactual release and diffusional loss of rhod-2.

Under these more stringent conditions of dye compartmentation, simultaneous recording of Ca_c and Ca_m at high temporal resolution suggests that three phases of Ca^{2+} traffic are initiated by Ca^{2+} entry. Fig. 5 *A* shows typical (*n* = 4) time courses of rhod-2 and Calcium Green dextran responses to a 500-ms depolarization, and Fig. 5 *B* shows overlaid initial segments of these records (sampled every 100 ms) on an expanded time scale. The three phases are best seen by plotting each Ca_m value against Ca_c (Fig. 5 *C*). During the 500-ms depolarization-induced Ca^{2+} entry, Ca_m rose without delay and in direct proportion to the rise in Ca_c . In the following 5 s, Ca_m continued to rise as Ca^{2+} was cleared from the cytosol, and Ca_c fell proportionately. In the final >50 s, Ca_m fell slowly during the final return of Ca_c to its resting value.

Our previous work suggested that mitochondria sequester the majority of the Ca^{2+} that enters the cell in response to depolarization (Herrington et al., 1996; Park et al., 1996). As was expected, we now see directly that Ca_m increases as Ca_c decreases. It now also is clear that half of mitochondrial uptake is accomplished while entry is still in progress and that uptake thus must attenuate the peak Ca_c reached.

Ca^{2+} Is Extruded from Mitochondria by Na^+ - Ca^{2+} Exchange

In the simplest interpretation, the fall in Ca_m that occurs when Ca_c returns to near resting values would represent export of mitochondrial Ca^{2+} . Alternatively, Ca_m might fall as a consequence of conversion of Ca^{2+} to an inaccessible form within the mitochondrial matrix (Lehninger, 1970; Friel and Tsien, 1994). The following experiments distinguish between these possibilities.

The benzothiazepine CGP-37157 selectively and potently inhibits the Na⁺-Ca²⁺ exchanger that exports Ca²⁺ from isolated mitochondria (Cox and Matlib, 1993). Fig. 6, A and B show that this agent has no effect on the rise of Ca_m evoked by Ca²⁺ entry into chromaffin cells but reversibly slows the Ca_m fall (n = 5), providing good evidence that extrusion of Ca^{2+} by the mitochondrial Na⁺-Ca²⁺ exchanger mediates the normal fall in Ca_m. While Ca_m is elevated and Ca²⁺ is being extruded from mitochondria, Ca_c often remains slightly above its resting value (Figs. 3-5). This small elevation has been attributed to a steady flux of Ca²⁺ into the cytoplasm from mitochondria (Freil and Tsien, 1994; Werth and Thayer, 1994; Herrington et al., 1996). The reversible reduction of this persistent Ca_c elevation by the exchange inhibitor CGP-37157 (Fig. 6) confirms this hypothesis. Unexpectedly, the fall in Ca_m that followed removal of CGP-37157 was accompanied by no elevation (Fig. 6 A) or only minor elevation (Fig. 6 B) of Ca_c . It would be useful to repeat these experiments in cells where the other mechanisms of Ca²⁺ clearance are inhibited.

Ca_m Responds to Release of Ca^{2+} from Intracellular Stores

Having established that mitochondria take up Ca^{2+} that enters through voltage-gated Ca^{2+} channels of the plasma membrane, we now consider if they also take up Ca^{2+} released from intracellular stores. Chromaffin cells have IP₃sensitive Ca^{2+} stores that can be mobilized by agonists acting at G protein–coupled receptors (O'Sullivan et al., 1989; Malgaroli et al., 1990; Neely and Lingle, 1992*b*). We

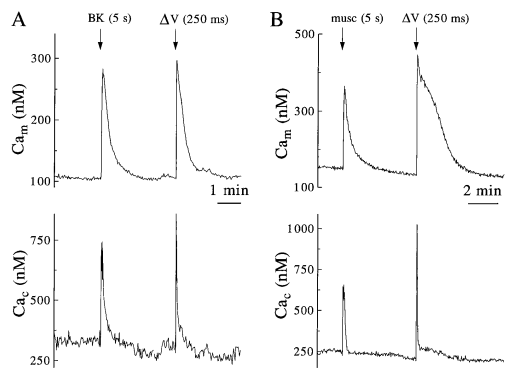


Figure 7. Similar mitochondrial responses to calcium entry and to calcium mobilization by agonists. Cells were coloaded with rhod-2 and Calcium Green esters, examined in perforated patch, and the data analyzed, all as in Fig. 4. Each record is representative of 3-5 similar experiments. (A) At the indicated times a single cell was perfused with 100 nM bradykinin (BK) for 5 s. After recovery, a 250-ms depolarization (ΔV) was applied. (B) Cam and Cac responses of another cell to a 5-s perfusion with 50 μ M muscarine (musc) and subsequent depolarization.

tested the selectivity of mitochondrial Ca^{2+} uptake for this source of Ca^{2+} by comparing responses of Ca_m and Ca_c to depolarizing steps and to agonists. For these experiments we used the perforated-patch configuration for recording since intracellular stores tend to run down quickly in wholecell recording so responses to agonists vanish (Herrington, J., Y.B. Park, D.F. Babcock, and B. Hille, unpublished observations).

Cells were first briefly perfused with bradykinin (n = 5) or muscarine (n = 3) and 3–4 min later given a brief depolarizing step (Fig. 7). The Ca²⁺-mobilizing agonists and voltage steps raised Ca_c to similar, but not identical, levels (see Materials and Methods), and the amplitudes of the corresponding Ca_m signals also were similar. Hence, mitochondria appear to sequester cytoplasmic Ca²⁺ ions, whether entering from outside or released from intracellular stores. The initial recovery of Ca_c after stimulation by agonist was consistently slower than after depolarization-induced Ca²⁺ entry, very likely because IP₃-dependent mobilization continues for some seconds after the agonist is washed away.

Chromaffin cells also contain a Ca^{2+} store that can be mobilized by caffeine or by Ca^{2+} entry during perforatedpatch recording (Park et al., 1996). This Ca^{2+} -induced Ca^{2+} release (CICR) is especially apparent in cells with impaired mitochondrial Ca^{2+} uptake (as in Figs. 3 *C* and 4) and can be recognized in experiments like that in Fig. 8 *A* where depolarizations of increasing duration were applied to a cell in perforated-patch mode. The plots of averaged Ca_m vs Ca_c from four such experiments (Fig. 8 *B*) show that Ca_c continued to rise for several hundred milliseconds after the Ca^{2+} entry evoked by depolarization had ceased (*large filled symbols*). These plots can be compared with the similar plot for whole-cell recording (Fig. 5 *C*) where Ca_c began to fall as soon as the membrane depolarization ended. In the perforated patch experiments (Fig. 8 *B*), Ca_m also rose more steeply during the period of CICR, as if mitochondria were taking up the additional Ca^{2+} released from stores. While spatially averaged Ca_c was still highly elevated, Ca_m then began to fall, as if reuptake into the depleted stores was creating a local sink for Ca^{2+} fed from mitochondria.

To substantiate the interpretation that mitochondria sequester Ca²⁺ released from CICR stores, we coapplied caffeine and the reticular Ca²⁺-ATPase inhibitor BHQ to deplete CICR stores and prevent their refilling (Fig. 9). When caffeine and BHQ were added, Cam and Cac increased transiently (Fig. 9 A; n = 4), showing that when caffeine induces release of Ca²⁺ from stores, mitochondria will take it up. For the preparation of cells used in these experiments, channel rundown (see Materials and Methods) was more severe than for those of Fig. 8. A control stimulus of 500 ms was needed to produce a Ca_c challenge similar to that found with a 250-ms pulse in the previous experiments (compare the trajectories marked with circles in Figs. 8 B and 9 B). Lengthening of the depolarization to 1 and 1.5 s then was required for subsequent stimuli applied in the presence of inhibitors.

Consider the trajectories of Ca_c and Ca_m that follow the depolarizing pulses (Fig. 9 *B*). Under the control conditions, there was a full second during which Ca_c was relatively constant while Ca_m continued to rise. In the presence of caffeine and BHQ, subsequent depolarizations still gave large averaged peak Ca_c responses but the rise of Ca_m after the end of the depolarization was greatly reduced (Fig. 9 *B*), confirming that much of the late rise of Ca_m represents uptake of Ca^{2+} released by CICR.

The initial rate of rise of Ca_m during Ca²⁺ entry is faster in these perforated-patch experiments than it was with the whole-cell recordings of Fig. 5. Using perforated-patch it was 674 ± 150 nM s⁻¹ (n = 9), and in whole-cell mode it was 79 ± 30 nM s⁻¹ (n = 6). We suspect that the faster rise

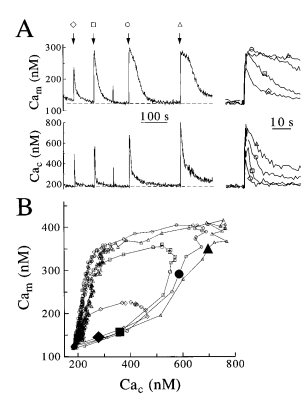


Figure 8. Calcium-induced release of Ca^{2+} and its mitochondrial sequestration. Cells were coloaded with rhod-2 and Calcium Green and examined in perforated patch as in Fig. 4. (*A*) Calibrated (*left*) and aligned (*right*) Ca_m and Ca_c records are from a single cell subjected to sequential depolarizations of 50 (*diamond*), 100 (*box*), 250 (*circle*), and 500 (*triangle*) ms depolarizations, applied at the arrows. Results are representative of four similar experiments. (*B*) Averaged responses from four cells treated exactly as above. The enlarged, closed symbols mark the observed or interpolated Ca_m and Ca_c values at the termination of stimulus.

reflects a combination of CICR and greater proximity of mitochondria to locally high microdomains of Ca_c around Ca^{2+} channels (Neher and Augustine, 1992; Robinson et al., 1995) in the undialyzed cells. Dialysis may disrupt cytoskeletal elements that establish a mitochondrial localization near the plasma membrane and other organelles. In the first second after entry ceased, when such microdomains should be dispersed, Ca_m continued to rise at ~40 nM s⁻¹ in the experiment of Fig. 5 *B*.

Discussion

The original description of the synthesis and characterization of rhod-2 noted that its AM-ester might be selectively accumulated, hydrolyzed, and trapped in mitochondria (Minta et al., 1989), and recent confocal imaging has shown compartmentation of this probe (Burnier et al., 1994; Tsien and Bacskai, 1995). It has also been shown qualitatively that Ca^{2+} entry (Tsien and Bacskai, 1995) and Ca^{2+} mobilization (Hajnóczky et al., 1995, Rutter et al., 1996) increase the signal from compartmentalized rhod-2. Our imaging of rhod-2 by fluorescence deconvolution methods and our quantitative photometric observations during application of pharmacological agents further validate and

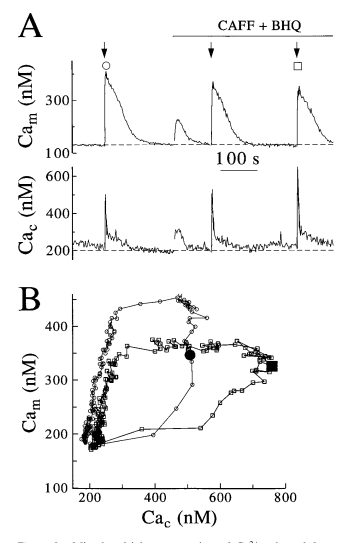


Figure 9. Mitochondrial sequestration of Ca^{2+} released from caffeine-sensitive stores. Cells were coloaded with rhod-2 and Calcium Green and examined in perforated patch as in Fig. 4. (*A*) Ca_m and Ca_c responses from another cell subjected to sequential depolarizations of 0.5 s applied before (*circle*), and of 1.0 or 1.5 s (*box*) applied after perfusion with 10 mM caffeine and 10 μ M BHQ. (*B*) Averaged responses from three cells treated exactly as above. The enlarged, closed symbols mark the observed Ca_m and Ca_c values at the termination of stimulus. The intermediate response to the first depolarization applied after treatment is not shown, to avoid excessive overlap of records.

extend the use of rhod-2 as a probe of mitochondrial $[Ca^{2+}]$ in living cells.

Previous work using blockers of mitochondrial function implicated mitochondria in the recovery of neurons and chromaffin cells from imposed Ca²⁺ entry (Thayer and Miller, 1990; Friel and Tsien, 1994; Werth and Thayer, 1994; White and Reynolds, 1995; Park et al., 1996; Herrington et al., 1996). For chromaffin cells we had found that blockers of mitochondrial Ca²⁺ uptake but not blockers of other cellular Ca²⁺ transporters greatly slow the initial clearance of Ca_c yet shorten the total time to return to the resting state (Park et al., 1996; Herrington et al., 1996). Now our simultaneous monitoring of Ca_m and Ca_c clearly establishes that Ca²⁺ uptake through the mitochondrial uniporter underlies rapid removal of cytosolic Ca²⁺. It shows further that subsequent Ca²⁺ export from mitochondria requires the mitochondrial Na⁺-Ca²⁺ exchanger and accounts for the slowness of the final decrease in Ca. With the less-invasive perforated-patch method we find that mitochondria accumulate Ca²⁺ mobilized from either IP₃- or caffeine-sensitive internal stores, and speculate that stores may be refilled with Ca²⁺ exported from mitochondria. A picture emerges of a highly interactive network of cellular Ca²⁺ signaling that involves mitochondria and at least two mobilizable pools. The rapid rise of Ca_m to even modest physiological Ca_c loads presumably also means that mitochondrial energy metabolism adjusts itself to cellular activities from moment to moment (McCormack et al., 1990; Gunter et al., 1994; Hajnóczky et al., 1995; Rutter et al., 1996).

What Is the Free Calcium Level in Mitochondria?

In the next sections we show that the Ca²⁺-handling properties of mitochondria within cells are similar to those found in detailed studies of isolated mitochondria. Some previous work used permeant precursors to load the fluorescent Ca²⁺ probes indo-1 and fura-2 into isolated mitochondria (Lukács et al., 1987; Gunter et al., 1988; Reers et al., 1989; McCormack et al., 1989; Leisey et al., 1993). Mitochondria within cells have also been loaded nonselectively with these dyes and studied after quench of cytosolic dye with Mn^{2+} (Mivata et al., 1991) or by imaging of mitochondria-rich regions (Sheu and Jou, 1994) to estimate a compartmentalized signal. For both isolated and cellular mitochondria, these studies with ratiometric dyes obtained low resting values of Ca_m in the range of 80-200 nM, much as for nonratiometric rhod-2 in our work. Thus it is agreed that there is hardly any gradient of free $[Ca^{2+}]$ across the resting mitochondrial membrane, despite a large, insidenegative $\Delta \psi_{\rm m}$. Presumably the uptake via the uniporter is in balance with extrusion via exchange and other mechanisms, and perhaps the uniporter is largely shut down at rest (Gunter and Pfeiffer, 1990; Gunter et al., 1994).

When cytoplasmic or extramitochondrial $[Ca^{2+}]$ is raised to $1-2 \mu M$, all studies with dyes report elevations in Ca_m to 250–1,000 nM, again as we have found. The general agreement supports the calibration protocols we have used for the nonratiometric rhod-2 dye. On the other hand, work with cells expressing mitochondrially targeted aequorin reports that Ca_m increases (for only a few seconds) to $>5 \ \mu M$ upon Ca²⁺ entry into excitable cells or mobilization of Ca²⁺ stores in nonexcitable cells (Rizzuto et al., 1992, 1993, 1994; Rutter et al., 1993). This higher level could be reconciled with the dye results if heterogeneity among dye-containing compartments allowed some to rise briefly to very high Ca²⁺ levels, which would saturate dyes and give high acquorin emission, while others had only moderate and longer lasting $[Ca^{2+}]$ elevations, keeping the spatially averaged dye signal below saturation. Alternatively, there may be subtle difficulties in calibrating aequorin signals. For single CHO cells responding to a Ca^{2+} mobilizing agonist, Rutter et al. (1996) found that rhod-2 fluorescence transiently increased 2-3-fold. Assuming a resting Ca_m of 100 nM and using the spectral properties and affinities of rhod-2 that we observe in vitro, we estimate that this corresponds to a peak Ca_m of 400–800 nM, much like the responses found in the present study, but much smaller than the >10 μ M peak Ca_m estimated from the responses of the same CHO cells transfected with mitochondrial aequorin.

What is the State of Mitochondrial Calcium?

Mitochondria can accumulate enormous quantities of Ca^{2+} . Classical experiments showed that isolated mitochondria exposed to modest Ca^{2+} concentrations, socalled "limited loading," accumulate up to 100 nmol Ca^{2+} (mg protein)⁻¹ in half a min without damage (Lehninger et al., 1967; Gunter and Pfeiffer, 1990). Exposure to millimolar Ca^{2+} ("massive loading") can lead to accumulations up to 2,600 nmol Ca^{2+} (mg protein)⁻¹, accompanied by irreversible damage, morphological changes, and precipitation of hydroxyapatite crystals. These values would be equivalent to Ca^{2+} contents of 36 and >900 mM within the mitochondria, respectively!

We now calculate how much the mitochondrial Ca²⁺ content should rise in our experiments if, for example, 70% (Herrington et al., 1996) of the Ca^{2+} in a single 1.5- μ M cytoplasmic challenge were taken up into the mitochondria. The mitochondria would actually remove much more than the 1.5- μ M free Ca²⁺ because almost all the cytoplasmic Ca²⁺ is reversibly bound to proteins and other Ca²⁺ buffers. In rat chromaffin cells (see Figs. 2 and 3 of Park et al., 1996) and in other excitable cells (Neher, 1995), typical values for cytoplasmic Ca2+-binding ratios are near 100. Thus, mitochondria would have to remove $0.7 \times 100 \times 1.5$ μ M = 105 μ M total Ca²⁺. Since the mitochondrial volume is only 6% of the combined cytosolic and nuclear volumes in rat chromaffin cells (Tomlinson et al., 1987), this uptake should raise the mitochondrial Ca²⁺ content by 1.75 mM. These numbers emphasize how remarkable it is that compartmentalized dyes report Cam rises of only 250-450 nM during such a challenge.

Why is the rise of Ca_m several thousand times less than expected? Like the cytoplasm, the mitochondrial matrix must contain Ca^{2+} buffers. These could include membrane phospholipids, Ca^{2+} -binding proteins, and phosphate compounds. The inner leaflet of the inner mitochondrial membrane contributes 6–20 mM of phospholipids (Carafoli, 1979; Schwerzmann et al., 1986), and the matrix contains tens of millimolar of phosphate compounds. The kind and concentration of Ca^{2+} -binding proteins is not well known.

We would suggest that the Ca²⁺ taken up in our experiments binds quickly and reversibly to sites in the mitochondrion and does not precipitate as insoluble compounds. If this were not true, the full Ca²⁺ load would not be able to re-enter the cytoplasm within seconds after application of CCCP (e.g., Figs. 1 *D* and 5 *B* of Herrington et al., 1996). We conclude therefore that the mitochondrial matrix buffers Ca²⁺ reversibly with an effective Ca²⁺ binding ratio of ~4,000 (see also Coll et al., 1982; Lukács and Kapus, 1987). This is 40 times the Ca²⁺-binding ratio of cytoplasm.

How Fast Can Mitochondria Transport Ca²⁺?

Rates of Ca²⁺ transport for isolated mitochondria have been extensively studied. Taking typical reported values,

the Ca²⁺ uniporter is half maximally activated around 10– 20 μ M Ca²⁺ and has a maximum velocity at room temperature ~10–30 nmol (mg protein)⁻¹ s⁻¹ in energized cardiac and liver mitochondria (Carafoli, 1979; Gunter and Pfeiffer, 1990). The maximum velocity of the mitochondrial Na⁺-Ca²⁺ exchanger is 30–400 times slower. Although the maximum velocity may be significantly affected by the bathing phosphate, Mg²⁺, pH, and other conditions, we will use it to compare with the uptake rates we measure in cells.

What would be the effect of a uniporter operating near its maximum velocity, at 20 nmol Ca²⁺ (mg protein)⁻¹ s⁻¹? Since 1 nmol (mg protein)⁻¹ corresponds to 385 µmol/liter of mitochondria (Schwerzmann et al., 1986), we would expect that a liter of mitochondria could take up 7,700 µmol Ca²⁺ s⁻¹. Once again we apply the mitochondrial volume (6%) and the cytoplasmic Ca²⁺-binding ratio (100) and predict that Ca_c (the free Ca²⁺) should fall at 4.6 µM s⁻¹ when the uniporter is working at full velocity.

We have previously measured the rates of Ca_c decay in chromaffin cells attributable to mitochondrial uptake after depolarization-induced Ca^{2+} entry (Herrington et al., 1996). At 1.5 μ M Ca_c the mean rate was 0.8 μ M s⁻¹. In Park et al. (1996), we gave evidence for appreciable uptake *during* a 200-ms depolarization, and in the present paper, we find that Ca_m rises more rapidly during the Ca²⁺ entry than afterward; indeed more than half of the rise in Cam occurs during a 500-ms depolarizing pulse (Figs. 5, 8, and 9). Thus, we presume that early uptake proceeds at rates equivalent to a decay of $Ca_c > 0.8 \mu M s^{-1}$. These velocities for Ca²⁺ import by mitochondria in living cells seem quite compatible with the 4.6 μ M s⁻¹ maximum velocity predicted from uptake by isolated mitochondria; however, they are higher than would be expected at 1.5 μ M Ca²⁺ if we take into account the reported Ca²⁺ affinity and cooperativity of uptake. A model in which a subset of mitochondria lies so close to the plasma membrane that they transiently see elevated Ca_c values well above those reported by the cytoplasmic dye (Lawrie et al., 1996) could explain these results. Perhaps also Ca²⁺ transport is more effective for mitochondria bathed in cytoplasm than for isolated mitochondria.

*Exploring the Cellular Ca*²⁺ *Network*

We propose that mitochondria have a unique role in the cytoplasmic Ca²⁺ network. The plasma membrane and reticular stores deliver Ca^{2+} rapidly to the cytoplasm, by opening ion channels, and then remove it slowly by primary and secondary active transport. Conversely, and consistent with their origins as a cellular symbiont within eukaryotic cytoplasm, mitochondria rapidly remove Ca²⁺ by a uniporter that acts like a channel, then return it more slowly by secondary active transport. Hence, the plasma membrane and internal stores create pulses of Ca_c with microdomains of high concentration suitable for initiating exocytosis or rapid muscle contraction. Mitochondria, on the other hand, act in an apparently constitutive manner, capturing Ca²⁺ pulses and then returning the Ca²⁺ slowly to the cytoplasm where it can prolong activation of highaffinity, Ca²⁺-dependent processes and perhaps feed refilling of stores. In some cells this transient sequestration into mitochondria is important in generation of Ca²⁺ waves and oscillations (Jouaville et al., 1995; Hehl et al., 1996). In addition, the transiently elevated Ca_m probably stimulates mitochondrial ATP production in anticipation of cellular needs. Such a major role for mitochondria in Ca²⁺ clearance holds for chromaffin cells and neurons but not for skeletal and cardiac muscles with a highly elaborated sarcoplasmic reticulum. We need now to examine the importance of mitochondria in other cell types and to determine if mitochondrial Ca²⁺ uptake is a regulated process.

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