## **Electron Transfer between Cytochrome c and p66Shc Generates Reactive Oxygen Species that Trigger Mitochondrial Apoptosis**

Marco Giorgio, <sup>1,2,8,\*</sup> Enrica Migliaccio, <sup>1,2,8</sup><br>Francesca Orsini, <sup>1,2</sup> Demis Paolucci, <sup>3</sup> Francesco Paolucci,<sup>3</sup> and Pier Giuseppe Pelicci<sup>1,2,7,8,\*</sup> 2004; Pelicci, 2004). 1Experimental Oncology Department European Institute of Oncology The release of proapoptotic factors from mito-

**oxidative damage and have been implicated in the regu-** per se sufficient to mediate complete cyt c release re-<br>lation of specific cellular functions, including apoptosis, mains, however, controversial. lation of specific cellular functions, including apoptosis. a mains, however, controversial.<br>Mitochondrial ROS increase markedly after proapoptotic. Beactive oxygen species (ROS: superoxide, hydro-**Mitochondrial ROS increase markedly after proapoptotic** Reactive oxygen species (ROS; superoxide, hydro-<br>
signals, though the biological significance and the un-<br>
gen peroxide, and hydroxyl radicals) are potent intrasignals, though the biological significance and the un**derlying molecular mechanisms remain undetermined.** cellular oxidants, which have been proposed as critical **P66<sup>Shc</sup> is a genetic determinant of life span in mammals,** regulators of apoptosis (Danial and Korsmeyer, 2004).<br>which regulates ROS metabolism and apoptosis. We re-<br>Indeed, ROS induce the opening of the PTP through which regulates ROS metabolism and apoptosis. We re-<br>nort here that p66<sup>Shc</sup> is a redox enzyme that generates oxidation-dependent mechanisms and are potent in**port here that p66<sup>Shc</sup> is a redox enzyme that generates** a oxidation-dependent mechanisms and are potent in-<br>mitochondrial ROS (hydrogen peroxide) as signaling and ducers of apoptosis, both in cultured cells and in vivo mitochondrial ROS (hydrogen peroxide) as signaling molecules for apoptosis. For this function, p66<sup>Shc</sup> uti-<br>lizes reducing equivalents of the mitochondrial electron vond their documented proapoptotic activity when adlizes reducing equivalents of the mitochondrial electron yond their documented proapoptotic activity when ad-<br>transfer chain through the oxidation of cytochrome c. ministered exogenously, endogenously produced ROS **transfer chain through the oxidation of cytochrome c.** ministered exogenously, endogenously produced ROS **Redox-defective mutants of p66<sup>Shc</sup> are unable to in-** are implicated in the execution of the apoptotic pro-<br>duce mitochondrial ROS generation and swelling in gram itself, regardless of the type of triggering signal. **duce mitochondrial ROS generation and swelling in** gram itself, regardless of the type of triggering signal. **vitro or to mediate mitochondrial apoptosis in vivo.** In fact, a marked increase of intracellular ROS is con-**These data demonstrate the existence of alternative** sistently associated with apoptosis, and treatments redox reactions of the mitochondrial electron transfer with antioxidants<br>chain, which evolved to generate proapoptotic ROS and Noble, 1994). chain, which evolved to generate proapoptotic ROS **in response to specific stress signals.** The molecular mechanisms underlying increased

**\* Enrica Migliaccio,1,2,8** degenerative diseases, and aging. Mitochondria play a **Francesca Orsini,**<sup>1,2</sup> Demis Paolucci,<sup>3</sup><br>
Maurizio Moroni,<sup>4</sup> Cristina Contursi,<sup>4</sup><br>
20 **10** apoptotic factors, such as cytochrome c (cyt c), aif, apoptotic factors, such as cytochrome c (cyt c), aif, Giovanni Pelliccia,<sup>2</sup> Lucilla Luzi,<sup>2</sup> Saverio Minucci,<sup>1,4</sup> smac/diablo, and endoG, which are released into the **Massimo Marcaccio,<sup>3</sup> Paolo Pinton,<sup>6</sup>** cytosol during apoptosis, where they activate a series **Rosario Rizzuto,<sup>6</sup> Paolo Bernardi,<sup>5</sup> 6** *Paolo Bernardi,*<sup>5</sup> *Conserved by Paymatic activities that lead to the specific degra*dation of proteins and DNA (Danial and Korsmeyer,

Milan chondria is due to the disruption of the organelle integ- <sup>2</sup> FIRC Institute of Molecular Oncology rity, which is achieved through multiple and concomi-Milan tant events. In the case of cyt c, for example, its release <sup>3</sup>G Ciamician Chemistry Department requires permeabilization of the outer membrane, re-University of Bologna modeling of the mitochondrial *cristae*, and dissociation Bologna **both constituent** of cyt c from cardiolipin, a constituent of the inner <sup>4</sup> Congenia srl **EXECONGER 10** membrane (Green and Kroemer, 2004). These events Milan are triggered by the opening of a high-conductance<br>
<sup>5</sup> Biomedical Sciences Department **Algerity conducts** channel, the permeability transition pore (PTP) (Berchannel, the permeability transition pore (PTP) (Ber-University of Padova **nardi et al., 2001).** Opening of the PTP provokes an Padova **increase of inner membrane permeability to ions and** increase of inner membrane permeability to ions and <sup>6</sup> Experimental and Diagnostic Medicine Department solutes (the so-called permeability transition, PT), fol-University of Ferrara and the mitochondrial ma-Ferrara **the intervelling of the organelle, and physical rupture of its** frisch rupture of its <sup>7</sup> Department of Medicine and Surgery **7** Network outer membrane, with the consequent release of proteins University of Milan **of the intermembrane space, including cyt c (Bernardi** cyt c (Bernardi Milan et al., 2001). Another mechanism leading to alterations Italy of the mitochondrial membrane integrity is the translocation of proapoptotic proteins such as Bax and Bid from the cytosol to the outer mitochondrial membrane, **Summary** where they form channels and/or regulate the function of preexisting channels (Scorrano and Korsmeyer, **Reactive oxygen species (ROS) are potent inducers of** 2003). Whether these events then trigger the PT or are

ROS generation during apoptosis are unclear. It has **Introduction** been proposed that ROS accumulation is secondary to the progress of the apoptotic process and is caused by Apoptosis is essential for the maintenance of tissue ho-<br>the interruption of the mitochondrial electron transfer meostasis, and its deregulation is implicated in cancer, chain (ETC), a consequence of the PT and of cyt c release (Cai and Jones, 1998). Alternatively, ROS might \*Correspondence: marco.giorgio@ifom-ieo-campus.it; piergiuseppe. function to trigger mitochondrial apoptosis, serving as<br>pelicci@ifom-ieo-campus.it<br>a signaling molecules to initiate the PT. Notably, intracel-<br>a signaling m signaling molecules to initiate the PT. Notably, intracellular ROS arise prior to cyt c release during the actiinvolving p53 or TNF $\alpha$  or those triggered by hypergly- ptosis. cemia, ischemia/reperfusion, or viral infection (Trinei et al., 2002; Sakon et al., 2003; Brownlee, 2001; Becker, **Results** 2004).

ROS are generated accidentally, as by products of P66<sup>Shc</sup> Induces Permeability Transition aerobic metabolism, and mitochondria are the major **of Isolated Mitochondria** source of intracellular ROS (Wallace, 1999). During res-<br>To investigate whether the effect of p66Shc on the mitopiration, electrons are extracted from substrates and chondrial PTP is direct, we analyzed its ability to induce are then transferred to molecular oxygen through a swelling of purified mitochondria, measured as dechain of enzymatic complexes (I–IV). In the final step of crease of light absorbance of the organelle suspension. the ETC, cyt c oxidase (COX; complex IV) ensures the Mitochondria were purified from wild-type (wt) mouse reduction of molecular oxygen to water, without forma-<br>
liver and pretreated with a low concentration of  $Ca^{2+}$ tion of oxygen radicals. However, partial reduction of  $(7 \text{ u.M})$ , a condition that does not induce PT but sensioxygen with generation of ROS can occur if molecular tizes PTP opening to other signals (Bernardi et al., oxygen interacts with the ETC upstream of complex IV.  $\frac{20011}{20011}$  Addition of recombinant n66<sup>Shc</sup> (5–20 uM) oxygen interacts with the ETC upstream of complex IV. 2001). Addition of recombinant p66<sup>Shc</sup> (5–20  $\mu$ M) did<br>Experimental data indicate that ROS are indeed contin-<br>not induce swelling as compared to a large load of Experimental data indicate that ROS are indeed contin-<br>uously produced during mitochondrial respiration and<br> $C_1a^{2+}$  (0.2 mM) a well-characterized inducer of the PT uously produced during mitochondrial respiration and  $Ca^{2+}$  (0.2 mM), a well-characterized inducer of the PT that up to 2% of the total oxygen consumption is con-<br>(Figure 1A). Since p66<sup>Shc</sup> is not imported by mitothat up to 2% of the total oxygen consumption is con-<br>verted to ROS (Chance et al., 1979). How ROS are requ-<br>chondria in vitro Neptura et al. 1979 How ROS are requ-<br>chondria in vitro Neptura et al. 2004), this finding indi verted to ROS (Chance et al., 1979). How ROS are regu-<br>lated during apoptosis is unclear.<br>cates that p66<sup>Shc</sup> is unable to alter mitochondrial per-

P66<sup>Shc</sup> deletion in mice (p66<sup>Shc–/-</sup>) decreases the in-<br>cidence of aging-associated diseases, such as athero-<br>sclerosis, and prolongs life span (Migliaccio et al., 1999;<br>when the protein is allowed to enter the mitochon sclerosis, and prolongs life span (Migliaccio et al., 1999;<br>
Napoli et al., 2003; Francia et al., 2004). The underlying<br>
molecular mechanisms are, however, unknown. P66<sup>Shc</sup><br>
is a splice variant of p52<sup>Shc</sup>/p46<sup>Shc</sup>, two (Pelicci et al., 1992). P66<sup>Shc</sup> has the same modular structure of p52<sup>Shc</sup>/p46<sup>Shc</sup> (SH2-CH1-PTB) and con-<br>tains a unique N-terminal region (CH2); however, it is addition of recombinant p66<sup>Shc</sup> induced marked and<br>not in

linking (Migliaccio et al., 1999; Orsini et al., 2004; Pacini **P66Shc** Localizes within the Mitochondrial et al., 2004). Similarly, the p66<sup>Shc−/−</sup> mice are resistant **P66<sup>Shc</sup> Localizes within the Mitochondrial** et al., 2004). Similarly, the p66<sup>Shc−/−</sup> mice are resistant **P66<sup>Shc</sup>** to apoptosis induced by paraquat, hypercholesterolemia, and ischemia (Migliaccio et al., 1999; Napoli et P66<sup>Shc</sup> is enriched in the submitochondrial fraction

tochondrial depolarization and the release of cyt c tions from MEFs (Figure 1E) were treated with digitonin<br>observed after treatment of cells with various proapo- (which allows extraction of soluble proteins from the observed after treatment of cells with various proapo- (which allows extraction of soluble proteins from the ptotic signals, suggesting that it regulates the mitochondrial pathway of apoptosis. Since inhibition of PTP extracts membrane bound proteins). After centrifugaopening blocks the proapoptotic function of  $p66^{Shc}$ , the tion, the resulting pellets and supernatants were ana-PTP itself might be the target of p66<sup>Shc</sup> activity (Trinei lyzed by Western blotting (WB) by using anti-p66<sup>Shc</sup>, et al., 2002; Orsini et al., 2004). We report here that anti-cyt c, anti-mtHsp70, or anti-porin antibodies. Digip66<sup>Shc</sup> is a redox enzyme that generates mitochondrial tonin or digitonin/high-salt treatments did not modify

vation of several apoptotic pathways, such as those hydrogen peroxide for the induction of PT and apo-

liver and pretreated with a low concentration of  $Ca<sup>2+</sup>$ cates that p66<sup>Shc</sup> is unable to alter mitochondrial per-

not involved in Ras regulation but rather functions in<br>the intracellular pathway(s) that regulates ROS metabo-<br>lism and apoptosis (Migliaccio et al., 1997; Migliaccio<br>et al., 1999; Trinei et al., 2002).<br>Intracellular ROS l intracellular encores are decreased in powin<br>
cells, as revealed by the reduced accumulation of POS-sen-<br>
stive probes and the reduced accumulation of endoge-<br>
nous markers of oxidative stress (8-oxo-guanosine)<br>
(Trinei et

al., 2003; Zaccagnini et al., 2004).<br>Expression of p66<sup>Shc</sup> is indispensable for the mi-<br>Expression of p66<sup>Shc</sup> is indispensable for the mi-<br>ther investigate its localization, mitochondria prepara-Expression of  $p66^{Shc}$  is indispensable for the mi-<br>chondrial depolarization and the release of cyt c tions from MEFs (Figure 1E) were treated with digitonin



Figure 1. P66<sup>Shc</sup> Localizes within the Mitochondrial Intermembrane Space and Induces Swelling of Isolated Mitochondria

Absorbance changes of purified liver mitochondria, either intact (A) or treated with 25  $\mu$ M digitonin (B-D), pretreated with 7  $\mu$ M CaCl<sub>2</sub> and supplemented with the following: (A) 20  $\mu$ M collagenase as inactive control (control [ctrl]; black trace), 20  $\mu$ M p66Shc (red), or 200  $\mu$ M CaCl<sub>2</sub> (brown); (B) 20  $\mu$ M collagenase in 0.25 M sucrose (ctrl, black) or different concentrations of  $p66^\text{Shc}$  (5, 10, and 20  $\mu$ M, as indicated; red); (C) different concentrations of  $H_2O_2$  (blue) or control buffer (gray); (D) 20  $\mu$ M p66<sup>Shc</sup> (added to all kinetics) and 1  $\mu$ M CsA (violet), 20  $\mu$ M NEM (green) or control buffer (red). Each experiment was performed six times using different mitochondrial preparations and two distinct preparations of the p66<sup>Shc</sup>. (E) (Left panels) WB analysis of MEF subcellular fractions using anti-p66Shc, anti-mtHp70, anti-calnexin, and anti-cytoplasmic Hsp70 antibodies. T, total extract; Mt, mitochondria; ER, endoplasmic reticulum; Ct, cytosol. (Right panels) ER and Mt fractions were treated with Proteinase K (PK) and analyzed by WB with anti-calnexin or anti-p66<sup>Shc</sup> antibodies. (F and G) Mitochondrial fractions from MEFs were treated with the indicated concentrations of NaCl and/or digitonin. After centrifugation, pellets (P) and supernatants (S) were analyzed by WB.

brane) and of mtHsp70 (matrix), indicating that these **by Energized Mitochondria** experimental conditions did not provoke gross struc-<br>
P66<sup>Shc</sup> expression is indispensable for the upregulation tural alterations of purified mitochondria (Figures 1F of intracellular ROS during p53- or ischemia-induced and 1G). As expected, digitonin/high salt but not digi- apoptosis (Trinei et al., 2002; Pacini et al., 2004; Zactonin alone induced cyt c release in the supernatant cagnini et al., 2004). To investigate whether p66<sup>Shc</sup>-reg-(Griparic et al., 2004). P66<sup>Shc</sup>, instead, was partially and ulated ROS derive from mitochondria, we analyzed in-<br>equally released by either digitonin or digitonin/high tact liver mitochondria from wt or p66<sup>Shc-/-</sup> mice equally released by either digitonin or digitonin/high salt, suggesting that it exists within the inner mitochon-<br>
drial space in the form of both soluble and nonsoluble chemical that induces ROS- and p66<sup>Shc</sup>-dependent protein (mitochondrial p66<sup>Shc</sup> is about 50 ng/mg of total apoptosis. We had previously shown that in vivo treatproteins; see Figure S1 in the Supplemental Data avail-<br>able with this article online). Similar results were ob-<br>but not p66<sup>Shc-/-</sup> mice and that apoptosis in wt animals tained by morphometric analysis of immunoelectron is prevented by treatment with the antioxidant N-acetyl microscopy sections of wt MEFs, which revealed asso- cysteine. Likewise, CCl4 induces p66Shc- and ROSmembrane (55.7%), intermembrane space (35.4%), and chondria were collected 20 hr after treatment, and ROS matrix (8.9%) (data not shown). were measured by spectrofluorimetry, using a mole-

# the localization of porin (a marker of the outer mem- **P66Shc Stimulates Generation of ROS**

chemical that induces ROS- and p66<sup>Shc</sup>-dependent but not p66<sup>Shc−/−</sup> mice and that apoptosis in wt animals dependent apoptosis of MEFs (Figure S2). Liver mito-





(A and B) Fluorescence changes (ex. 498, em. 527; I, intensity of fluorescence) and corresponding bar graph (rate of fluorescence intensity/ min) (C) of mitochondria from wt (black lines in [A] and [B] and empty bars in [C]) or p66<sup>Shc−/-</sup> (red lines and red bars) mouse livers, treated with placebo ([A], CTRL) or CCl<sub>4</sub> ([B], +CCl<sub>4</sub>), after sequential additions of H<sub>2</sub>DCFDA and ant A (black and red lines) or control ethanol (gray line, wt; green line, p66<sup>Shc-/-</sup>) with (plain) or without (dashed) succinate. (A) and (B) report one set of experiments representative of four; (C) reports the average of the four experiments ± SEM (#, wt ctrl versus wt +CCl<sub>4</sub>, p < 0,01; #, p66<sup>Shc-/-</sup> + CCl<sub>4</sub> versus wt + CCl<sub>4</sub>, p < 0.01). Respiration was controlled for each mitochondrial preparation. (D) Fluorescence changes (ex. 498, em. 527) of purified liver mitoplasts after sequential additions of H<sub>2</sub>DCFDA, succinate and ant A (blue), p66<sup>Shc</sup> (red), or 20 µM collagenase (ctrl, black). (D) reports one set of experiments. The average of six experiments is reported in the graph of the right panel ± SEM, where results are expressed as fold increase in fluorescence rate induced by p66<sup>Shc</sup>, ant A, or control buffer (\*, p66<sup>Shc</sup> versus ctrl, p < 0.01). (E) Fluorescence changes (ex. 498, em. 527) of purified liver mitoplasts after sequential additions of H<sub>2</sub>DCFDA and succinate (green), p66<sup>Shc</sup> (red), or ant A (blue). (F) Respiration analysis of mitoplasts after additions of p66<sup>Shc</sup> (red), ant A (blue), or control buffer (ctrl, black). ADP (100  $\mu$ m) was added to all samples to maximize respiration. (G and H) Fluorescence changes in intact wt (black) or p66Shc<sup>-/-</sup> (red) MEFs incubated with H<sub>2</sub>DCFDA (G) or HE (ex. 410, em. 490) (H) and treated with ant A (plain) or ethanol as control (dashed).

cular probe that becomes fluorescent upon oxidation pensable for the increased generation of mitochondrial by ROS (2',7'-dichlorofluorescin diacetate; H<sub>2</sub>DCFDA) ROS after challenge with proapoptotic stimuli and that (LeBel et al., 1992). The rate of mitochondrial H<sub>2</sub>DFCDA p66<sup>Shc</sup> requires a functional ETC for this activity. (LeBel et al., 1992). The rate of mitochondrial H<sub>2</sub>DFCDA p66<sup>Shc</sup> requires a functional ETC for this activity.<br>
oxidation increased significantly after CCl<sub>4</sub> treatment in To analyze the mechanisms underlying the regulat oxidation increased significantly after  $CCl<sub>4</sub>$  treatment in the wt mice, while it remained unchanged in the p66<sup>Shc−/−</sup> mice (Figures 2A–2C). Notably, H<sub>2</sub>DCFDA oxi-<br>dation was consistently detected only in the presence scavenging activities in mitochondrial preparations from dation was consistently detected only in the presence of respiratory substrates, such as succinate (Figures wt and p66<sup>Shc−/−</sup> mouse livers were comparable (Figure 2A–2C) or glutamate/malate (data not shown). Together, S3). Furthermore, we found no differences in any of the<br>these findings indicate that p66<sup>Shc</sup> expression is indis-<br>known ROS scavenging systems (catalase, SODs, and

effect of p66<sup>Shc</sup> on mitochondrial ROS generation, we known ROS scavenging systems (catalase, SODs, and GSH-peroxidase) in samples from wt or p66<sup>Shc−/−</sup> fibro- oxygen reduction using reducing equivalents from the blasts and tissues (Figure S3), suggesting that the ef-<br>fect of p66<sup>Shc</sup> on mitochondrial ROS is due to increased tion by p66<sup>Shc</sup> in the range of 10 nM/min/mg of mito-

tion might be direct or, rather, be the consequence of induced  $H_2$ DFCDA fluorescence in mitoplasts or mito-<br>the organelle swelling induced by  $p66^{Shc}$ . To distinguish chondria suspensions; data not shown). the organelle swelling induced by p66<sup>Shc</sup>. To distinguish between these two possibilities, we investigated the effect of recombinant p66Shc on ROS generation by puri- **P66Shc Is a Redox Protein that Mediates** fied mitoplasts. Mitoplasts are unable to swell after PTP **Electron Transfer** opening, yet they respire (due to a fraction of cyt c that We therefore investigated whether p66<sup>Shc</sup> is capable remains associated to the inner membrane; Figure 2F). itself to execute an electron transfer (ET) reaction. To Strikingly, addition of recombinant p66<sup>Shc</sup> to mitoplasts this end, we measured the ability of recombinant increased H<sub>2</sub>DCFDA oxidation (Figure 2D). These data  $p66^{Shc}$  to mediate ET when adsorbed onto a probe indicate that the effect of  $p66^{Shc}$  on mitochondrial ROS electrode by cyclic voltammetry (CV) (Armstrong, 2002). production is not a consequence of the organelle swell- Protein adsorption was performed through the carboxing and suggest that p66<sup>Shc</sup> directly stimulates mito-<br>
ylic group of 11-mercaptoundecanoic acid self-assemchondrial ROS generation. bled as a monolayer (SAM) onto a gold electrode. In the

 $(H<sub>2</sub>O<sub>2</sub>)$  by dismutation of superoxide anion  $(O<sub>2</sub>)$ , which is produced by reduction of oxygen by semiquinone subtraction from the oxidation and reduction scans (Balaban et al., 2005). A well-characterized mechanism resulted to zero (Figure 3A upper panel; red line). Strik-<br>Leading to mitochondrial ROS generation is inhibition of ingly, when the SAM electrode was coated with recom leading to mitochondrial ROS generation is inhibition of ETC downstream to quinone. Antimycin A (ant A), for binant  $p66^{Shc}$ , the two CV scans were modified by the example, blocks the ETC at the level of complex III and superimposition of additional oxidation and reduction is a potent inducer of O<sub>2</sub><sup>-</sup> and H<sub>2</sub>O<sub>2</sub> production (Bala- events, in the regions of 100 and –170 mV, respectively. ban et al., 2005). Therefore, we investigated the effects Baseline subtraction resulted in two sharp peaks (Fig-<br>of p66<sup>Shc</sup> on mitochondrial respiration and H<sub>2</sub>O<sub>2</sub>/O<sub>2</sub> ure 3A). The intensity of the two peaks was alway of p66<sup>Shc</sup> on mitochondrial respiration and H<sub>2</sub>O<sub>2</sub>/O<sub>2</sub><sup>-</sup> ure 3A). The intensity of the two peaks was always metabolism. proportional to the scan rate (data not shown). Fur-

nonrespiring mitochondria (data not shown) or mi- contributed to the faradaic ET reaction was about 6 toplasts (Figures 2D and 2E). Unlike ant A, however, pmol/cm<sup>2</sup>, consistent with a homogenous and thin p66<sup>Shc</sup> did not decrease oxygen consumption of mi-<br>
layer of an electrode-coated protein of about 66 kDa. toplasts (Figure 2F) or mitochondria (data not shown), These data indicate that electrode-coated p66<sup>Shc</sup> unindicating that it does not block respiration. To obtain dergoes an ET process. in vivo confirmation, oxygen consumption was measured in purified liver and heart mitochondria and in **P66Shc Oxidizes Cyt c In Vitro** intact primary fibroblasts from wt and p66<sup>Shc−/−</sup> mice The average of the oxidation and reduction peaks is a under either basal or uncoupled conditions and using direct measure of the redox potential of an ET protein, different substrates. In no case did we observe sig- which, in the case of p66<sup>Shc</sup>, is about −35 mV. Among nificant differences in respiration between wt and the redox systems that are present within the mito-

H<sub>2</sub>DCFDA or hydroetidine (HE), two fluorescent probes [2002]). Therefore, we evaluated whether cyt c and that are oxidized by  $H_2O_2$  or  $O_2$ ; respectively (LeBel et al., 1992; Zhao et al., 2003). Their specificity in our experimental system was confirmed by the findings not shown) on the p66<sup>Shc</sup>-coated electrode provoked a that basal and  $p66^{Shc}$ -induced H<sub>2</sub>DCFDA oxidation by dramatic change of its CV response (Figure 3B). First, mitoplasts was inhibited by catalase (an  $H_2O_2$  scaven-<br>ger) but not by superoxide dismutase (SOD) (which longer detected, while a novel, single ET event was reger) but not by superoxide dismutase (SOD) (which generates  $H_2O_2$  from  $O_2$ <sup>-</sup>), while HE oxidation was inhibited by SOD but not by catalase (Figure S4). Analysis tinct from that of either p66<sup>Shc</sup> or cyt c alone (Figure of basal and ant A-induced oxidation of H<sub>2</sub>DCFDA or 3A). Second, the calculated electrode capacitance of HE in wt and p66<sup>Shc-/-</sup> MEFs (Figures 2G and 2H) and p66<sup>Shc</sup> was reduced of about 30% in the presence of HE in wt and p66<sup>Shc−/−</sup> MEFs (Figures 2G and 2H) and purified liver mitochondria (data not shown) revealed cyt c, suggesting that cyt c formed an additional film decreased levels of H<sub>2</sub>O<sub>2</sub> but not O<sub>2</sub><sup>--</sup> in the p66<sup>Shc-/-</sup> onto the p66<sup>Shc</sup>-coated SAM electrode. These findings samples. It appears, therefore, that p66<sup>Shc</sup> stimulates indicate that p66<sup>Shc</sup> and cyt c interacted samples. It appears, therefore, that p66<sup>Shc</sup> stimulates  $H_2O_2$  generation without affecting mitochondrial respi-<br>were integrated in the same ET event. Finally, the reration or production of  $O_2^-$ . Since, however, it requires a functional ETC, these data suggest that p66<sup>Shc</sup> favors closer in the presence of cyt c (from 270 to 35 mV),

tion by p66<sup>Shc</sup> in the range of 10 nM/min/mg of mito-ROS production rather than diminished scavenging. chondrial protein, both in vivo (Figures 2A and 2B) The effect of p66<sup>Shc</sup> on mitochondrial ROS produc- and in vitro (Figure 2D) (using standard curves of H<sub>2</sub>O<sub>2</sub>-

absence of adsorbed proteins, the CV curve reflected **P66Shc Produces Mitochondrial Hydrogen Peroxide** the unperturbed and symmetrical capacitive response without Inhibiting Respiration **business and reduction** of the SAM electrode to alternate oxidation and reduc-Mitochondrial respiration generates hydrogen peroxide tion scans, which is typical of long-chain alkylthiols SAMs (Figure 3A, left panel). Accordingly, baseline Like ant A, p66<sup>Shc</sup> did not induce ROS production in thermore, the calculated concentration of p66<sup>Shc</sup> that

p66<sup>Shc−/−</sup> mitochondria (data not shown). chondria, the redox value of cyt c (17 mV) is one of the To characterize ROS production by  $p66^{Shc}$ , we used closest to that of  $p66^{Shc}$  (Figure 3A and Chen et al. p66<sup>Shc</sup> can exchange electrons. Absorption of cyt c (but not of glucose oxidase, used as negative control; data corded with a redox potential of 10 mV (Figure 3B), disduction and oxidation peaks of p66Shc became much



Figure 3. Redox Properties of p66<sup>Shc</sup> and of the p66<sup>Shc</sup>-Cyt c Couple

(A) CV experiments using uncoated (upper panel, Ctrl), p66<sup>Shc</sup> (middle panel)-, or cyt c (lower panel)-coated SAM electrodes.

(B) CV experiment using a p66<sup>Shc</sup>-coated SAM electrode in the presence of cyt c.

(C) Absorption spectrum of reduced cyt c in the presence of 20  $\mu$ M p66<sup>Shc</sup> (red) or control buffer (black).

(D) Absorbance changes at 550 nm of reduced cyt c in the presence of the indicated concentrations of recombinant p66Shc (red), yeast-

purified COX (green), control buffer (black), or 20 μM collagenase (gray line).<br>(E) Fluorescence changes of a 20 μM H<sub>2</sub>DCFDA solution during the sequential addition of 20 μM p66<sup>Shc</sup> (red) or control buffer (black), 10 μ reduced cyt c, and 50  $\mu$ M CuSO<sub>4</sub>.

indicating that the ET kinetics of p66<sup>Shc</sup> is accelerated right panel). Furthermore, when the initial potential was by cyt c. Therefore, it appears that p66<sup>Shc</sup> executes a set at +100 mV (which cannot reduce the p66<sup>Shc</sup>/cyt c redox reaction with cyt c under the CV experimental redox center), neither catalytic oxygen reduction nor conditions.  $H_2O_2$  oxidation was detected (data not shown). To-

dox status of cyt c by spectrophotometry. Addition of ET between p66shc and cyt c leads to oxygen reduction  $p66<sup>Shc</sup>$  modified the absorbance spectrum of reduced and formation of H<sub>2</sub>O<sub>2</sub>. cyt c, provoking a slight shift of the  $\gamma$  peak and a reduction of the amplitude of the α peak (Figure 3C). These **P66Shc Generates Hydrogen Peroxide** modifications are consistent with a strong interaction **from Reduced Cyt c In Vivo** between p66<sup>Shc</sup> and the prosthetic group of cyt c, and We then investigated whether, in vivo, the ETC site of the partial oxidation of cyt c. Analysis of the kinetics of p66<sup>Shc</sup>-induced ROS production is compatible with the absorbance at 550 nm ( $\alpha$  peak) revealed that the effect described in vitro activity of p66<sup>Shc</sup> on reduced cyt c. of p66<sup>Shc</sup> on cyt c oxidation status is dose dependent To this end, mitochondria were treated with ascorbate/ and comparable to that of purified yeast COX (Figure N,N'-tetramethyl-p-phenyldiamine (TMPD) in the ab-3D). Together, these findings indicate that p66<sup>Shc</sup> is a sence of energetic substrates. Under these experimen-

H<sub>2</sub>DCFDA when reacting with cyt c. Addition of p66<sup>Shc</sup> detectable ROS (Figure 3G; black trace). Addition of reto reduced cyt c did not provoke H<sub>2</sub>DFCDA oxidation combinant p66<sup>Shc</sup>, instead, induced H<sub>2</sub>DFCDA oxida-(data not shown). A significant fluorescence boost from tion, indicating that p66<sup>Shc</sup> is able to generate ROS by oxidized H2DCFDA was, instead, recorded in the pres- acting downstream to reduced cyt c. Consistently, ence of copper (but not zinc or iron) ions (Figure 3E; p66<sup>Shc</sup> did not increase ROS generation induced by red trace), indicating that, in vitro, the copper atom cat- inhibitors of complexes I (rotenone) or III (ant A) (Figalyzes generation of free radicals from the redox center ure S4). of the p66<sup>Shc</sup>-cyt c complex. Notably, under anoxic conditions, H2DCFDA oxidation was significantly de- **Mapping of the Redox Center of p66Shc** creased, suggesting a role for oxygen intermediates in  $\qquad$  To map the region of p66 $S<sup>the</sup>$  interacting with cyt c, we this process (Figure 3E; dashed trace). Since H<sub>2</sub>DCFDA compared, by ELISA, the cyt c binding properties of oxidation is specifically sensitive to H<sub>2</sub>O<sub>2</sub>, these data p66<sup>Shc</sup> with those of p52<sup>Shc</sup> or p46<sup>Shc</sup>, which ar oxidation is specifically sensitive to  $H_2O_2$ , these data suggest that this is the molecular species that forms involved in the regulation of ROS and apoptosis (Migduring the reaction. **liaccio et al., 1999**). Increasing amounts of coated cyt

the electrochemical reaction of p66<sup>Shc</sup> with cyt c, we cof Shc proteins (0.08–4  $\mu$ g) and bound proteins remeasured current generation at 650 mV (which corres-<br>vealed with anti-Shc antibodies. The p66<sup>Shc</sup>-cyt c assoponds to the oxidation potential of H<sub>2</sub>O<sub>2</sub>) in an electrode-<br>
ciation was dose dependent at each concentration of confined portions (Figure 4B). A residual binding activity confined p66Shc-cyt c system (Callegari et al., 2004). P66<sup>Shc</sup> and cyt c were entrapped within a conducting (about 10%) was detected for p52<sup>Shc</sup>, while p46<sup>Shc</sup> did polymeric film and immobilized onto a glassy carbon not bind at all, suggesting that the N-terminal region of electrode. The coated electrode was then transferred p66<sup>Shc</sup> (CH2-PTB) is involved in cyt c binding. into an air- or argon-saturated phosphate buffer and a To narrow down the cyt c binding region of  $p66^{\text{Shc}}$ . −100 mV potential applied for 150 s (this potential in- we expressed various portions of the CH2-PTB region duces the continuous reduction of the p66<sup>Shc</sup>-cyt c as GST fusion proteins (Figure 4C) and measured their complex, has no direct effect on molecular oxygen, and ability to recover cyt c from cellular lysates through allows accumulation of oxygen radicals, if formed, in pull-down experiments. Results showed that a 52 the gel; accumulation step, Figure 3F). Analysis of cur- amino acid region N terminal to the PTB domain is critirent generation at −100 mV revealed early oxygen cal for the binding to cellular cyt c (denominated CB, reduction current generated by the  $p66^{Shc}/cyt$  c for cyt c binding; Figures 4A and 4C). This region is electrode, only in the presence of air-saturated buffer highly conserved among the known p66Shc vertebrate (Figure 3F; left panel). After switching of the electrode orthologs (Figure 4D) and contains glutamic (E125, potential to 650 mV (to measure accumulated  $H_2O_2$ ; de-<br>tection step, Figure 3F), current was recorded specific-<br>which, in the context of COX IV and yeast cyt c peroxi-

We then investigated the effect of p66<sup>Shc</sup> on the re- gether, these results provide direct demonstration that

redox protein that oxidizes cyt c in vitro. The state of tal conditions, mitochondrial respiration is supported by reduction of cyt c (by TMPD), while complexes I–III **Electron Transfer between p66**<sup>Shc</sup> and Cyt c **remain inactive. As expected, since the ETC sites of** Generates Hydrogen Peroxide In Vitro **ROS** production are excluded in this system, ascor-We then investigated whether  $p66^{\text{Shc}}$  oxidizes bate/TMPD-treated mitochondria did not generate

To obtain a direct proof that  $H_2O_2$  is generated during c (0.25–4  $\mu$ g) were incubated with increasing amounts

which, in the context of COX IV and yeast cyt c peroxially in the presence of air-saturated buffer (Figure 3F; dase, are essential for the interaction and ET reaction

<sup>(</sup>F) Current measurement at −100 (accumulation step) or +650 (detection step) mV of the glassy carbon electrode coated with a gel containing p66<sup>Shc</sup> and cyt c. Recordings were performed in aerated (air, blue and black traces) or argon-flushed (argon, green and red traces) phosphate buffer, as indicated.

<sup>(</sup>G) Fluorescence changes of purified liver mitoplasts after sequential additions of H<sub>2</sub>DCFDA, 0.5 mM ascorbate, 0.2 mM TMPD, and p66<sup>Shc</sup> (red) or control buffer (black).





Figure 4. Mapping of the p66<sup>Shc</sup> Cyt c-Interacting Region

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(A) Modular organization of Shc proteins (p66, p52, p46) and p66Shc mutants (*qq* and *f*). The positions of the *qq* and *f* mutations are indicated within the CB region.

(B) ELISA binding assay of recombinant p66Shc, p52Shc, p46Shc, p66Shc*qq*, and p66Shc*f* proteins added at different concentrations to dish wells preadsorbed with different concentrations of cyt c, as indicated.

(C) In vitro pull-down experiment (right panel) using MEF lysates and the schematized (left panel) portions of p66Shc expressed in bacteria as GST fusion proteins. Recovered proteins were analyzed by Western blotting using anti-cyt c antibodies.

(D) Amino acid sequence of the p66<sup>Shc</sup> N-terminal region. Amino acid sequence alignment of the CH2-PTB N-terminal regions (aa 1–155) of the available p66<sup>Shc</sup> vertebrate horthologs. Identities are highlighted in gray. The most conserved region (CB) is boxed in red. The relevant glutamic (E132 and E133) and triptophan (W134) residues are indicated in the sequence (by asterisks and circle, respectively) and in the p66<sup>Shc</sup> modular organization scheme below.



Figure 5. Impaired Redox Properties of p66Shc Mutants Defective for Cyt c Binding (A) Voltammograms of cyt c (green), p66Shc*qq*

(orange), and p66Shc*qq* + cyt c (black). (B) Voltammograms of cyt c (green), p66Shc*f* (pink), and p66Shc*f* + cyt c (black).

(C) Fluorescence changes of purified liver mitoplasts during the sequential additions of H<sub>2</sub>DCFDA; succinate; and buffer (ctrl, black), p66Shc (red), p66Shc*qq* (orange), or p66Shc*f* (violet) proteins.

(D) Ratio of the H<sub>2</sub>DCFDA mean fluorescence, obtained by FACS analysis of p66<sup>Sh</sup> MEFs after infection with control, p66Shc, p66Shc*qq*, or p66Shc*f* retroviruses; results are .<br>the average ± SEM of three experiments.

with cyt c (Zhen et al., 1999). We engineered similar stained cells (Figure 5D). Therefore, mutations that immutations within p66<sup>Shc</sup> (p66<sup>Shc</sup>E132Q-E133Q muta-<br>tion, p66<sup>Shc</sup>qq; p66<sup>Shc</sup>W134F mutation, p66<sup>Shc</sup>f) and of p66<sup>Shc</sup> to stimulate ROS generation in vivo. tion, p66<sup>Shc</sup>qq; p66<sup>Shc</sup>W134F mutation, p66<sup>Shc</sup>f) and evaluated their effects on the ability of the protein to bind cyt c, to transfer electrons, and to stimulate mito- **Mutation of the Redox Center of p66Shc Impairs** chondrial ROS generation. Notably, both *qq* and *f* muta- **its Ability to Mediate Mitochondrial Apoptosis** tions abrogated the ability of recombinant  $p66^{Shc}$  to We then investigated whether the function of  $p66^{Shc}$  to bind cyt c, as revealed by ELISA and pull-down experi-<br>regulate mitochondrial apoptosis depends on its elecments (Figures 4B and 4C). trochemical properties, using the p66<sup>Shc</sup>qq mutant. To

showed symmetric peaks with redox potentials of -151 other functions of p66<sup>Shc</sup>, we first evaluated the ability mV and -140 mV, respectively (Figures 5A and 5B). of the two neighboring domains (PTB and CH2) to bind These values are markedly lower than that of wt  $p66<sup>Shc</sup>$  activated receptors and undergo phosphorylation after (−35 mV), suggesting that the E132-E133 and W134 res- oxidative stress, respectively, and the property of idues influence the p66Shc redox properties. In the pres- p66Shc*qq* to localize within mitochondria. In the context ence of cyt c, the CV responses of both mutant proteins of the *gq* mutation, the p66<sup>Shc</sup> PTB domain maintained resulted in two asymmetric peaks (Figures 5A and 5B), its ability to bind activated EGF receptors (Figure 6A). which could be easily deconvoluted into two compo-<br>Likewise, p66<sup>Shc</sup>qq was efficiently phosphorylated on nents associated to the ET processes of cyt c (17 mV; serine following expression into p66<sup>Shc−/−</sup> MEFs and Figures 5A and 5B, green traces) and the p66<sup>Shc</sup>gg treatment with H<sub>2</sub>O<sub>2</sub> (Figure 6B). Like the wt protein, (−150 mV; Figure 5A, orange trace) or p66Shc*f* (−141 mV; p66Shc*qq* localizes within mitochondria (Figure 6C). Figure 5B, pink trace) mutants. The detection of sepa- Mitochondrial p66<sup>Shc</sup> is PK resistant (Figure 6D); it is rated ET processes is consistent with a decreased in-<br>released by digitonin treatment (Figure 6E) and forms a teraction between the two p66<sup>Shc</sup> mutants and cyt c, complex with mtHsp70, from which it dissociates after as also revealed by ELISA and pull-down experiments proapoptotic stimuli (Figure 6F). (Figures 4B–4D). Therefore, both properties of execut-<br>We then measured the ability of the p66<sup>Shc</sup>qq mutant ing ET and interacting with cyt c are impaired in the *qq* to regulate mitochondrial PT in vitro and apoptosis in vivo. or *f* mutants of p66<sup>Shc</sup>, suggesting that the E132-E133 P66<sup>Shc</sup>qq was unable to induce swelling of digitonized and W134 residues of p66<sup>Shc</sup> concur in the formation mitochondria (Figure 7A), nor did it restore the apo-

in vivo. Addition of recombinant p66<sup>Shc</sup>qq and p66<sup>Shc</sup>f bility (Figure 7D). Notably, CsA inhibited the ability of to digitonin-treated mitochondria did not increase reexpressed p66<sup>Shc</sup> to restore mitochondrial apoptosis H<sub>2</sub>DCFDA oxidation (Figure 5C). Likewise, expression following H<sub>2</sub>O<sub>2</sub> (Figure 7E) or staurosporine (data not of p66<sup>Shc</sup>qq and and p66<sup>Shc</sup>f into p66<sup>Shc−/−</sup> MEFs did shown), supporting the theory that a functional PT is not increase the cellular fluorescence of  $H_2$ DCFDA- needed for the proapoptotic activity of p66 $\text{S}_{\text{hc}}$ . It ap-

The CV curves of the p66<sup>Shc</sup>qq and p66<sup>Shc</sup>f mutants ensure that the  $qq$  mutation had not interfered with

of its redox center and cyt c binding surface. ptotic response of p66<sup>Shc−/−</sup> MEFs upon treatment with We then compared the ability of p66<sup>Shc</sup>, p66<sup>Shc</sup>qq,  $H_2O_2$  or staurosporine, as measured by cyt c release and p66<sup>Shc</sup>f to regulate production of ROS in vitro and (Figure 7B), caspase activation (Figure 7C), and cell (Figure 7B), caspase activation (Figure 7C), and cell via-



Figure 6. Receptor Binding, Phosphorylation, Localization, and Hsp70 Association of the p66Shc Redox Mutants

(A) In vitro pull-down experiment using lysates of MEFs and treated or not with EGF, and GST, GST-PTBp66Shc, or GST-PTBp66<sup>Shc</sup>qq. Recovered proteins were analyzed by WB using anti-EGFR antibodies.

(B) P66Shc−/− MEFs were reconstituted with p66Shc or p66Shc*qq* by retroviral-mediated gene transfer and treated or not with  $H_2O_2$ . Anti-Shc immunoprecipitates were analyzed by WB using antibodies against phosphorylated p66<sup>Shc</sup> (upper panel) or total pool of p66Shc (lower panel).

(C) WB analysis of subcellular fractions from p66Shc−/− MEFs reexpressing p66Shc or p66Shc*qq*, using the indicated antibodies.

(D) Mitochondria (Mt) and ER-enriched fractions (ER) were treated with PK and analyzed by WB with indicated antibodies.

(E) MEFs mitochondrial fractions (as in [C]) were treated with the indicated concentrations of digitonin. After centrifugation, pellets (P) and supernatants (S) were analyzed by WB.

(F) MEFs mitochondrial fractions (as in [C]) were immunopreciptated with anti-p66<sup>S</sup> antibodies. Specific immunoprecipitates were analyzed by WB using anti-mtHsp70 (upper panel) or anti-p66<sup>Shc</sup> (lower panel) antibodies.

ter abrogates the functions of p66<sup>Shc</sup> to induce mito-<br>number of modifications of p66<sup>Shc</sup> to ende-

for the proapoptotic function of p66<sup>Shc</sup> and support a p66<sup>Shc</sup> appears to be a specialized function whereby ratory chain can generate ROS not only accidentally

fraction of the mitochondrial electron flow, which is ated from cyt c by p66<sup>Shc</sup> for the production of ROS and E.M. and F.O., unpublished data). The p66<sup>Shc</sup>of two functional states of p66<sup>Shc</sup>: inactive, under basal results with the inactivation of p66<sup>Shc</sup> and that pro-

pears, therefore, that mutation of the p66<sup>Shc</sup> redox cen- conditions, and active, after proapoptotic signals. A chondrial PT in vitro and to mediate cellular apoptosis scribed, which occur within minutes after treatment after stress. with various proapoptotic signals and might lead to activation of p66<sup>Shc</sup>: the cytosolic pool of p66<sup>Shc</sup> becomes **Discussion** serine phosphorylated (Migliaccio et al., 1999), the mitochondrial pool of p66<sup>Shc</sup> is released from a high The present findings provide a mechanistic explanation molecular weight complex (Orsini et al., 2004), and both for the proapoptotic function of p66<sup>Shc</sup> and support a mitochondrial and cytosolic p66<sup>Shc</sup> pools are increa model whereby p66<sup>Shc</sup> generates  $H_2O_2$  within mito-<br>chondria, which, in turn, induces opening of the PTP phosphorylation, genetic evidence indicates that this chondria, which, in turn, induces opening of the PTP phosphorylation, genetic evidence indicates that this and cellular apoptosis (Figure 8). ROS production by modification is required for the ability of p66<sup>Shc</sup> to mediand cellular apoptosis (Figure 8). ROS production by modification is required for the ability of p66<sup>Shc</sup> to medi-<br>n66<sup>Shc</sup> appears to be a specialized function whereby ate apoptosis (Migliaccio et al., 1999). The underly electrons are subtracted from the mitochondrial ETC to mechanism, however, is unknown. The mitochondrial catalyze the partial reduction of molecular oxygen. This pool of p66<sup>Shc</sup> is not serine phosporylated (E.M. and is the first demonstration that the mitochondrial respi-<br>
ratory chain can generate ROS not only accidentally<br>
p66<sup>Shc</sup> from cytosol to mitochondria does not occur afbut also through a specific enzymatic system. In this ter proapoptotic signals (Orsini et al., 2004), suggesting context, p66 $s$ hc can be regarded as an atypical signal that serine phosphorylation might serve other, nonmitotransducer that converts proapoptotic into redox signals. chondrial activities of p66<sup>Shc</sup> that are also needed to In vitro and in vivo experiments demonstrated that exert its proapoptotic function. Mitochondrial p66<sup>Shc</sup>  $p66^{Shc}$  oxidizes cyt c and generates  $H_2O_2$ . Therefore, a exists within a high-molecular-weight complex, which fraction of the mitochondrial electron flow, which is includes members of the TIM-TOM import complex mostly used by COX to reduce oxygen to water, is devi- (TIM44, TIM20, TIM23, and mtHsp70; Orsini et al. [2004] (Figure 8). P66<sup>Shc</sup> expression, however, does not influ- mtHsp70 complex is destabilized following treatment ence the mitochondrial transmembrane potential under with proapoptotic signals, leading to the release of steady-state conditions, while it is indispensable for its monomeric p66<sup>Shc</sup>. Since recombinant p66<sup>Shc</sup> poscollapse following proapoptotic signals (Orsini et al., sesses constitutive redox activity, it is tempting to 2004). These findings are compatible with the existence speculate that association with the TIM-TOM complex



Figure 7. The p66<sup>Shc</sup>qq Mutant Does Not Induce PT, Nor Does It Activate Apoptosis

(A) Absorbance changes of digitonized liver mitochondria pretreated with Ca<sup>2+</sup> and added of p66Shc (red trace) or p66Shc*qq* (orange) proteins.

(B) WT and p66Shc−/− MEFs infected with control, p66Shc, or p66Shc*qq* retroviruses were treated with 2  $\mu$ M staurosporine and, after 6 or 12 hr, cytosolic fractions analyzed by WB (left panels). Cytosolic cyt c was analyzed by ELISA in the same cells after 6 hR treatment with 2  $\mu$ M staurosporine or 800  $\mu$ M H<sub>2</sub>O<sub>2</sub> (right bar graph).

(C and D) The same cells as in (B) were treated with 2  $\mu$ M staurosporine or 800  $\mu$ M H2O2 and analyzed for caspase-3 activation (by flow cytometry using antibodies against cleaved caspase-3; [C]) or trypan blue (D).

(E) The same cells as in (B) were treated with  $H<sub>2</sub>O<sub>2</sub>$  and 5  $\mu$ m CsA and analyzed by trypan blue. Results represent the mean of triplicate cultures ± SEM.

apoptotic signals activate p66<sup>Shc</sup> by releasing it from concentrations of reduced cyt c, thus firing the ET reacthis inhibitory complex. This inhibitory complex. This in this inhibitory complex.

A further mechanism of regulation of the proapo-<br>ptotic function of mitochondrial p66<sup>Shc</sup> might be linked aged DNA are significantly reduced in p66<sup>Shc-/-</sup> primary to its intrinsic redox properties. Since the calculated cultures (fibroblasts, endothelial cells, lymphocytes;  $\mathsf{K}_\mathsf{eq}$  of the [p66<sup>Shc</sup>] + [cyt c]  $\leftrightarrow$  [p66<sup>Shc</sup>]<sup>-</sup> + [cyt c]<sup>+</sup> redox Nemoto and Finkel [2002]; Trinei et al. [2002]; Zaccagreaction is 0.1, cyt c oxidation by p66<sup>Shc</sup> is unfavored nini et al. [2004]; Pacini et al. [2004]). Likewise, markers when reduced cyt c is present at a concentration com-<br>
of oxidative stress (8-oxo-guanosine, isoprostane, nitparable to p66<sup>Shc</sup>. The reaction might, instead, occur if rotyrosine) are decreased in tissues from p66<sup>Shc−/−</sup> mice an excess of reduced cyt c is present, a condition that (Trinei et al., 2002; Napoli et al., 2003; Francia et al., is achieved when COX activity is decreased (e.g., dur- $2004$ ), suggesting that p66<sup>Shc</sup> might regulate ROS proing hypoxia) or by physiological inhibition by nitric ox- duction also in the absence of acute (proapoptotic) ide (NO) (Sarti et al., 2003). Notably, reduced cyt c might stress signals and that a fraction of intracellular ROS also accumulate during tBid/Bax-induced apoptosis, (and of oxidative stress) depends, under basal condisince both molecules induce remodeling of the inner tions, on the expression of p66<sup>Shc</sup>. Since p66<sup>Shc</sup> is actimembrane and increase of free intermembrane cyt c vated by virtually every type of stress, the basal pro-(Scorrano et al., 2002), which is then fully reduced by duction of ROS by  $p66<sup>Shc</sup>$  may reflect its moderate the outer membrane rotenone-insensitive NADH-cyt  $b_5$  activation by chronic stress. The redox balance has reductase (Bernardi and Azzone, 1981). Therefore, dif-<br>
profound effects on metabolism and transcription, thus ferent proapoptotic mechanisms may lead to increased providing the molecular basis for a function of p66<sup>Shc</sup>



aged DNA are significantly reduced in p66<sup>Shc−/−</sup> primary

Figure 8. Model of p66Shc Redox Activity during Mitochondrial Apoptosis

Proapoptotic signals induce release of p66<sup>Shc</sup> from a putative inhibitory complex. Active p66Shc then oxidizes reduced cyt c (red) and catalyzes the reduction of  $O_2$  to  $H_2O_2$ . PTP opening by  $H_2O_2$  then leads to swelling and apoptosis. NADH-cyt  $b<sub>5</sub>$  reductase is indicated as additional putative source of reduced cyt c.

in the cellular adaptation to chronic stress. Regulation **Acknowledgments** of redox equilibrium and apoptosis by p66<sup>Shc</sup> might<br>equally contribute to its effect on life span.<br>cal support, C. Tacchetti and C. Puri for EM analysis, and V. Raker

solic cyt c was measured after 6 hr by ELISA and after 6/12 hr by WB of cytosolic fractions; anti-caspase-3 positive cells after 6 hr, Received: November 10, 2004<br>by FACS; and cell survival after 20 hr, by trypan blue. Intracellular Revised: March 22, 2005 ROS were measured incubating cells for 30 min with 10  $\mu$ M<br>  $H_2$ DCFDA (Molecular Probes) in complete culture medium; cells<br>
were then recovered and suspended in PBS for FACS analysis. Mi-<br>
toplast (0.5 mg/ml) or 0.4 mg/ toplast (0.5 mg/ml) or 0.4 mg/ml mitochondrial suspension and 30 **References** !M DH2CFDA were sequentially added to a spectrofluorimetric cuvette. Fluorescence (excitation wavelength [ex.] 498; emission wavelength [em.] 525) was registered using a Perkin Elmer Ls55 Armstrong, F.A. (2002). Insights from protein film voltammetry into<br>wavelength [em.] 525) was registered using a Perkin Elmer Ls55 Armstrong, F.A. (2002). Insi spectrofluorimeter at 25°C. Anti-p66<sup>Shc</sup> antibodies have been de-<br>complex biological electron-transies antibodies against connected and the Soc. Dalton Trans. 5, 661. scribed (Orsini et al., 2004). Antibodies against caspase-3, mtHsp70, cytHsp70, calnexin, porin, phosphorylated p66Shc Ser36 Balaban, R.S., Nemoto, S., and Finkel, T. (2005). Cell *120*, 483–495. residue, phosphotyrosine, and cyt c were obtained from commer-<br>
cial sources. The p66<sup>Shc</sup>qq and f mutants were generated using the<br>
QuikChange Site-Directed Mutagenesis Kit (Stratagene) and<br>
cloned into the PINCO vector (

Cell lysates were subjected to differential centrifugations to sepa- Bernardi, P., Petronilli, V., Di Lisa, F., and Forte, M. (2001). A mitorate nuclei (700 rgf), mitochondria (8000 rgf), and the endoplasmic chondrial perspective on cell death. Trends Biochem. Sci. *26*, reticulum (15,000 rgf). Mice were intraperitoneally injected with 2 112-117. mg CCl<sub>4</sub>/g of body weight CCl<sub>4</sub> 20 hr before liver collection. Each<br>
mitochondrial preparation was assayed for respiration by using a<br>
Clark type oxygraph and swelling, as described (Petronilli et al.,<br>
1994), with minor min on ice). Callegari, A., Cosnier, S., Marcaccio, M., Paolucci, D., Paolucci, M.,

electrode (reference), platinum spiral wire (counter electrode), and Mat. Chem. *14*, 807–810. gold disk (working electrode). The SAM was formed by soaking the Chance, B., Sies, H., and Boveris, A. (1979). Hydroperoxide metab-<br>gold electrode for 5 hr in 100  $\mu$ M 11-mercaptoundecanoic acid. olism in mammalian organ Purified horse cyt c (Sigma), p66Shc, p66Shc*qq*, or p66Shc*f* solutions The direction of the electrode surface and left to evaporate<br>slowly. Measurements were preformed in phosphate buffer and<br>LiClO4 at constant ionic strength (100 mM). Before each experi-<br>ment, the electrolyte solution was pu (lower than 1 ppm O2), and the voltammetric curves were recorded Danial, N.N., and Korsmeyer, S.J. (2004). Cell death: critical control maintaining an argon blanket over the solution. CV experiments points. Cell 116, 205-2 maintaining an argon blanket over the solution. CV experiments. were carried out at 5–50 mV/s with an Autolab Model PGSTAT 30 at Diwan, J.J., Yune, H.H., Bawa, R., Haley, T., and Mannella, C.A.<br>25°C. Protein midpoint potentials (E1/2) were calculated as average (1988) Enhanced untake o

0.25  $\mu$ g purified horse cyt c (Sigma) by using a 4  $\mu$ g/ml coating Green, D.R., and Kroemer, G. (2004). The pathophysiology of mitosolution in carbonate buffer (0.1 ml/well). After washing with PBS, chondrial cell death. Science *305*, 626–629. different concentrations of recombinant Shc proteins were added Griparic, L., van der Wel, N.N., Orozco, I.J., Peters, P.J., and van to each well and incubated for 2 hr at 37°C. Binding was revealed to each well and incubated for 2 hr at 37°C. Binding was revealed der Bliek, A.M. (2004). Loss of the intermembrane space protein<br>using an anti SH2-SHC antibody (Transduction Laboratories) and Mam1/OPA1 induces swelling an using an anti SH2-SHC antibody (Transduction Laboratories) and<br>horseradish peroxidase-conjugated anti-rabbit antibodies. Bound<br>antibodies were detected using the TMB detection reagent (SIGMA).<br>LeBel, C.P., Ischiropoulos, H

gen species formation and oxidative stress. Chem. Res. Toxicol. 5,<br>Supplemental Data include four figures and can be found with this 227–231.<br>article online at http://www.cell.com/cgi/content/full/122/2/221/ Mayer, M., and article online at http://www.cell.com/cgi/content/full/122/2/221/ DC1.

for some of the mitochondrial fractionation experiments. F.O. is re- **Experimental Procedures** cipient of a FIRC fellowship. This work was supported by grants From A.I.R.C. and M.I.U.R. M.G.; S.M., P.B., and P.G.P. are share-<br>MEFs were treated with 2 uM staurosporine or 800 uM H<sub>2</sub>O<sub>2</sub>. Cyto-<br>MEFs were treated with 2 uM staurosporine or 800 uM H<sub>2</sub>O<sub>2</sub>. Cyto-<br>

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25°C. Protein midpoint potentials (*E1/2*) were calculated as average (1988). Enhanced uptake of spermidine and methylglyoxal-bis(gua-<br>between the potential of oxidation (*Epa*) and reduction (*Epc*) peaks. pulbydrazone) b nylhydrazone) by rat liver mitochondria following outer membrane lysis. Biochem. Pharmacol. *37*, 957–961.

Recombinant Proteins, ELISA, and In Vitro Binding Assay<br>
All recombinant proteins were produced in E. coli as GST fusion<br>
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