Isozyme-specific Interaction of Protein Kinase Cδ with Mitochondria Dissected Using Live Cell Fluorescence Imaging

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PKCδ signaling to mitochondria has been implicated in both mitochondrial apoptosis and metabolism. However, the mechanism by which PKCδ interacts with mitochondria is not well understood. Using FRET-based imaging, we show that PKCδ interacts with mitochondria by a novel and isozyme-specific mechanism distinct from its canonical recruitment to other membranes such as the plasma membrane or Golgi. Specifically, we show that PKCδ interacts with mitochondria following stimulation with phorbol esters or, in L6 myocytes, with insulin via a mechanism that requires two steps. In the first step, PKCδ translocates acutely to mitochondria by a mechanism that requires its C1A and C1B domains and a Leu-Asn sequence in its turn motif. In the second step, PKCδ is retained at mitochondria by a mechanism that depends on its C2 domain, a unique Glu residue in its activation loop, intrinsic catalytic activity, and the mitochondrial membrane potential. In contrast, these determinants, only the C1B domain is required for the phorbol ester-stimulated translocation of PKCδ to other membranes. PKCδ also basally localizes to mitochondria and increases mitochondrial respiration via many of the same determinants that promote its agonist-evoked interaction. PKCδ localized to mitochondria has robust activity, as revealed by a FRET reporter of PKCδ-specific activity (δCKAR). These data support a model in which multiple determinants unique to PKCδ drive a specific interaction with mitochondria that promotes mitochondrial respiration.

PKC is a major family of serine/threonine kinases that plays crucial roles in cellular signal transduction, regulating diverse cellular behaviors such as survival, growth and proliferation, migration, and apoptosis (1–5). The ten full-length mammalian isozymes of the PKC family serve different biological roles and are regulated differently (1–5). These isozymes are classified as either conventional (α, the alternatively spliced βI and βII, and γ), novel (δ, ε, θ, η), or atypical (ζ, ι/λ) based on the nature of their N-terminal regulatory domains (1–5). Conventional and novel PKC isozymes possess tandem C1A and C1B regulatory modules that enable them to translocate to membranes in response to receptor-mediated generation of the lipid second messenger diacylglycerol (DAG)3 or in response to stimulation with phorbol esters, a class of tumor-promoting compounds that are potent functional DAG analogues. For conventional isozymes, translocation to membranes is also facilitated by a Ca2+–binding C2 regulatory module. The C2 domain of novel isozymes does not bind Ca2+ or directly enable translocation to membranes, but their C1B domain has a 100-fold higher affinity for DAG that compensates in membrane recruitment (6, 7). Additionally, the C2 domain of PKCδ serves as a phosphotyrosine protein-binding domain (8) and is itself tyrosine-phosphorylated at several key sites that drive nuclear localization of PKCδ (4, 9–11). Atypical isozymes are not regulated by either DAG or Ca2+ but do bind anionic phospholipids and bind other proteins via a PB1 domain. Constitutive phosphorylation at three conserved sites in the kinase domain of PKCs process them into a catalytically competent but inactive conformation (1–5, 12). These sites are the activation loop, phosphorylated by

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3 The abbreviations used are: DAG, diacylglycerol; CKAR, C kinase activity reporter; δCKAR, PKCδ-specific CKAR; PDHC, pyruvate dehydrogenase complex; PDBu, phorbol-12,13-dibutyrate; BisIV, bisindolylmaleimide IV; NaPpI, 4-amino-1-tert-butyl-3-(1′-naphthyl)-pyrazolo[3,4-d]pyrimidine; FCCP, carbonyl cyanide-4-(trifluoromethoxy)phenylhydrazone; RFP, red fluorescent protein; CFP, cyan fluorescent protein; YFP, yellow fluorescent protein; TOM20, translocase of the outer membrane 20; Mito-, mitochondrially targeted; PM, plasma membrane; OCR, oxygen consumption rate; RF, regulatory fragment; CF, catalytic fragment.
the upstream kinase phosphoinositide-dependent kinase-1, and two C-terminal autophosphorylation sites: the turn motif and hydrophobic motif. The kinase complex mTORC2 promotes the priming phosphorylation of some PKC isozymes but not of PKCd (13). The hallmark for PKC activation is its translocation to intracellular membranes, notably plasma membrane and Golgi, where second messenger- or phorbol ester-mediated engagement of its regulatory domains allosterically releases an autoinhibitory pseudosubstrate peptide sequence from its active site (1–5, 14).

PKCd, a novel PKC isozyme ubiquitously expressed in mammalian tissues, has been implicated in both of the two main functions of mitochondria: mitochondrial apoptosis and metabolism. PKCd knock-out mice exhibit various phenotypes, such as increased proliferation of B cells (15), exacerbated vein graft arteriosclerosis (16), and suppression of radiation-induced apoptosis in salivary parotid cells (17), that are consistent with a proapoptotic role for PKCd. A large body of literature has addressed the role of PKCd in apoptotic signaling to mitochondria, with cell fractionation and immunofluorescence experiments showing the activity-dependent translocation of PKCd to mitochondria in a variety of cell lines upon induction of apoptosis (18–21). In particular, phorbol esters have been observed to induce the translocation of endogenous and exogenous PKCd to mitochondria in multiple cell lines, with prolonged phorbol ester treatment leading to apoptosis (22, 23). PKCd knock-out mice also exhibit cardiac phenotypes, which tend to show that PKCd is cardioprotective (24, 25), and proteomic analysis of cardiac tissue from PKCd knock-out mice shows changes in the levels of various metabolic proteins (24, 26). Importantly, PKCd has been implicated in the regulation of the pyruvate dehydrogenase complex (PDHC), which controls the flux of fuel entering the citric acid cycle by converting pyruvate to acetyl CoA in the mitochondrial matrix, thereby regulating mitochondrial respiration. PKCd does this either by activating the phosphatases that dephosphorylate and activate the PDHC in response to insulin in muscle and liver cells (27) or by indirectly inhibiting one of the kinases (PDH kinase 2) that phosphorylate and inactivate the PDHC in response to vitamin A in mouse embryonic fibroblasts (28–30). The precise mechanism by which PKCd interacts with mitochondria remains to be established.

Here, we show that PKCd binds to and is active at mitochondria by an isozyme-specific mechanism distinct from that driving the canonical PKC interaction with plasma membrane or Golgi. We examine signaling at mitochondria by visualizing and quantifying PKCd translocation to, retention at, basal interactions with, and activity at mitochondria using FRET-based reporters in real time in live cells. We demonstrate that multiple determinants unique to PKCd drive a highly specific interaction with mitochondria that promotes mitochondrial respiration.

**EXPERIMENTAL PROCEDURES**

Materials—Phorbol-12,13-dibutyrate (PDBu), G66976, G66983, bisindolylmaleimide (BisIV), and rottlerin were obtained from Calbiochem. 4-Amino-1-(tet-butyl-3-(1′-naphthyl)-pyrazolo[3,4-d]pyrimidine (NaPP1) was a generous gift from Dr. Kavita Shah. Insulin, oligomycin, carbonyl cyanide-4-(trifluoromethoxy)phenylhydrazone (FCCP), and MeSO (as a solvent for drugs and as a vehicle control) were obtained from Sigma-Aldrich. The PKCd phospho-activation loop, PKCd/θ phospho-turn motif, and pan-PKC phospho-hydrophobic motif antibodies were purchased from Cell Signaling. The DsRed antibody, which recognizes the DsRed variant red fluorescent protein (RFP), and the GFP antibody, which recognizes the GFP variants cyan fluorescent protein (CFP) and yellow fluorescent protein (YFP), were obtained from Clontech.

**Plasmids**—All of the plasmids were constructed in the mammalian expression vector pcDNA3 (Invitrogen). Targeting of CFP and activity reporters to the outer membrane of mitochondria (Mito-CFP and Mito-δCKAR) was achieved by the addition of the sequence encoding the N-terminal 35 amino acids of TOM20 (translocase of the outer membrane 20) to the 5′ end of constructs (31–34). Targeting of CFP to Golgi (Golgi-CFP) was achieved by the addition of the sequence encoding the N-terminal 35 amino acids of endothelial nitric-oxide synthase to its 5′ end (31, 33). Targeting of CFP to the plasma membrane was achieved by the addition of the sequence encoding the N-terminal 7 amino acids of Lyn kinase to its 5′ end (31, 35). Full-length mouse PKCd, rat PKCdII, and human PKCdε and most of the PKCd mutants were C-terminally YFP tagged (PKCd-YFP) for the intermolecular FRET translocation assay. An N-terminally YFP-tagged mouse PKCd (YFP-PKCd) was also constructed and used to generate the PKCdδE500G/D/Q and PKCdL636T/N637R kinase domain mutants. For PKCdΔC2, the first 126 residues were deleted. PKCdC2 consists of the first 88 residues of PKCd. PKCdΔC1A, PKCdΔC1B, and PKCdΔC1AΔC1B were constructed by inserting KpnI restriction sites flanking the domain(s) of interest, digesting out the domain(s), and religating the plasmids. For PKCdΔC1A, residues His-159 to Cys-208 were deleted. For PKCdΔC1B, residues His-231 to Cys-280 were deleted. For PKCdΔC1AΔC1B, residues His-159 to Cys-280 were deleted. PKCdΔC1B consists of residues 225–281. PKCdε(δC1B) consists of PKCdε with its C1B domain (resides His-243 to Cys-292 on PKCdε) replaced with the C1B domain of PKCdδ (resides His-231 to Cys-280 on PKCdδ). PKCdCF consists the C-terminal half of PKCd beginning with residue Asn-328. PKCdRF/εCF consists of the N-terminal 327 residues of PKCdδ fused directly to the C-terminal half of PKCd beginning with PKCdε residue Ser-388. The PKCdδE500G/D/Q, PKCdL636T/N637R, and PKCdM425A mutants were generated by site-directed mutagenesis following the QuikChange protocol (Stratagene). The C kinase activity reporter (CKAR) (31, 35, 36) and δCKAR (PKCd-specific CKAR) (34) constructs have been described previously. The negative control Thr/Ala mutant of Mito-δCKAR (Mito-δCKAR/A) was generated by site-directed mutagenesis following the QuikChange protocol. All of the constructs imaged in conjunction with the intramolecular FRET activity reporters were C-terminally RFP tagged.

**Cell Culture and Transfection**—COS-7 cells were cultured in DMEM (Cellgro) supplemented with 5% fetal bovine serum (HyClone or Invitrogen) and 1% penicillin/streptomycin (HyClone) at 37 °C in 5% CO2. COS-7 cells were transiently transfected using FuGENE 6 (Roche Applied Science).
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~24 h after the cells were plated. L6 myocytes were cultured in α-minimal essential medium (Invitrogen) supplemented with 10% fetal bovine serum and 1% penicillin/streptomycin at 37 °C in 5% CO₂. L6 myocytes were transiently transfected using Lipofectamine 2000 (Invitrogen) ~24 h after the cells were plated.

Live Cell Fluorescence Imaging—The cells were plated onto sterilized glass coverslips in 35-mm imaging dishes and co-transfected with the indicated constructs to monitor either translocation via intermolecular FRET or activity via intramolecular FRET. Approximately 48 h post-transfection, the cells were washed twice with and subsequently imaged in Hanks’ balanced salt solution (Cellgro) containing 1 mM CaCl₂ in the dark at room temperature. For some experiments, the cells were pretreated with the specified drugs in Hanks’ balanced salt solution for at least 20 min at 37 °C or as indicated. Otherwise, the indicated drugs were introduced during live cell imaging.

The images were acquired via a 100× objective on a Zeiss Axiovert microscope (Carl Zeiss Microimaging, Inc.) using a MicroMax digital camera (Roper-Princeton Instruments) controlled by MetaFluor software version 3.0 (Universal Imaging Corp.). Optical filters were obtained from Chroma Technologies. Time lapse images of CFP, FRET, YFP, and RFP were collected every 15 s through either a 10% or a 25% neutral density filter. CFP and FRET images were obtained through a 420/20-nm excitation filter, a 450-nm dichroic mirror, and either a 475/40-nm emission filter (CFP), or a 535/25-nm emission filter (FRET). YFP images monitored as a control for photobleaching were obtained through a 495/10-nm excitation filter, a 505-nm dichroic mirror, and a 535/25-nm emission filter. RFP images were obtained through a 560/25-nm excitation filter, a 505-nm dichroic mirror, and a 629/53-nm emission filter. Excitation and emission filters were switched in the filter wheel (Lambda 10-2; Sutter). Integration times were 200 ms for CFP and FRET and 100 ms for YFP and RFP.

For each cell imaged, a region encompassing the cellular organelle of interest was manually selected on the CFP image, and MetaFluor calculated a FRET ratio consisting of the average FRET/CFP emission ratio (for intermolecular FRET translocation reporters) or CFP/FRET emission ratio (for intramolecular FRET activity reporters) for this region. Base-line FRET ratios were acquired for 5 min before introduction of drugs, and the trace for each cell was slope-adjusted for any base-line drift (31) and normalized to the average base-line value. Throughout this paper, normalized FRET ratios are plotted as the means ± S.E. of data from the specified n number of cells (typically 10 cells/group) imaged over at least three and more typically five independent experiments.

The cells were stained with MitoTracker Deep Red (Invitrogen Molecular Probes) by being incubated with a 167 nM working solution of MitoTracker in prewarmed culture medium for 15 min at 37 °C and 15 min at room temperature before being washed and imaged in Hanks’ balanced salt solution. The images were pseudocolored and overlaid to visualize co-localization using ImageJ.

Immunoblotting—COS-7 cells grown in 6-well dishes were collected for immunoblotting ~24 h after transfection, when cells had grown to confluency. The cells were washed with ice-cold PBS; then lysed on ice in a buffer containing 1% Triton X-100, 50 mM Tris (pH 7.5), 10 mM Na₄P₂O₇, 50 mM NaF, 100 mM NaCl, 5 mM EDTA, 2 mM benzamidine, 50 μg/ml leupeptin, 1 mM PMSF, and 1 mM sodium vanadate; and then cleared by centrifugation at 16,000 × g for 2.5 min. Detergent-solubilized lysates were separated on SDS-PAGE gels, transferred onto PVDF membranes, and probed using the indicated antibodies. The blots were visualized via chemiluminescence on a FluorChem imaging system (Alpha Innotech).

Analysis of Mitochondrial Respiration—COS-7 cells transfected with each of the indicated constructs were plated in quintuplicate at 40,000 cells/well in growth medium in an XF-96-well plate (Seahorse Bioscience) for each experiment. Approximately 24 h later, the medium was changed to unbuffered DMEM containing 10 mM glucose, 10 mM pyruvate, and 2 mM GlutaMAX (Invitrogen), and a Seahorse Extracellular Flux Analyzer XF96 (Seahorse Bioscience) was used to measure the oxygen consumption rate (OCR). Rates of basal, State 4 (in the presence of 2 μM oligomycin), and uncoupler-stimulated (in the presence of a titrated concentration of FCCP) respiration were assessed. The cumulative data over multiple experiments are presented as the means ± S.E. relative to the vector control group. The same cells were plated in parallel in 6-well dishes to monitor the rates of cell growth and collected for immunoblotting to confirm expression of the constructs.

Statistical Analysis—Statistical analyses were performed using Prism 6 (GraphPad Software).

RESULTS

Intermolecular FRET Demonstrates that PKCα Interacts with Mitochondria via an Isozyme-specific Mechanism—To investigate the possibility that PKC signals at and interacts directly with mitochondria, we employed an intermolecular FRET reporter assay to monitor in real time and in live cells the interactions between an energy donor CFP targeted to the outer membrane of mitochondria using a mitochondrial localization sequence (31–34) and a co-transfected energy acceptor YFP fused to PKC (Fig. 1A). An increase in the FRET signal upon stimulation indicated translocation by a particular PKC isozyme to mitochondria. Proper localization of CFP to mitochondria in COS-7 cells was confirmed by the strandlike appearance of CFP consistent with the appearance of mitochondria in healthy cells and by its co-localization with MitoTracker Deep Red (Fig. 1B). YFP-tagged PKC isozymes tended to be excluded from the nucleus but were otherwise basally localized diffusely throughout the cytoplasm, with a basal concentration of PKCα in particular at Golgi (Fig. 1D). Acute activation of PKCs by stimulation with 200 nM of the phorbol ester PDBu induced all the PKCs to translocate rapidly to various intracellular membranes (Fig. 1D), but only PKCα translocated to and was retained at mitochondria (Fig. 1, C, black, and D). This was both quantified by a robust real time increase in the FRET signal aggregated over many cells and multiple experiments (Fig. 1C) and corroborated in time lapse YFP images (Fig. 1D). Despite phorbol esters being potent PKC agonists that activate all conventional and novel PKC isozymes, this translocation by PKCα to mitochondria proved to be an isozyme-spe-
cific phenomenon. PDBu did not induce a similar FRET response in either the conventional PKC isozyme PKC\(_{\alpha}/\)H9252 or the novel PKC isozyme PKC\(_{\alpha}/\)H9280 (Fig. 1C, red), despite the fact that both these isozymes clearly responded to PDBu by translocating to other cellular membranes (Fig. 1D). The only other PKC isozyme that was observed to translocate to mitochondria was PKC\(_{\alpha}/\)H9258, the PKC isozyme most closely related to PKC\(_{\alpha}/\)H9254 (37) but whose expression is much more limited (primarily to hematopoietic cells (38) and skeletal muscle (39)), which translocated to mitochondria at the same initial rate as PKC\(_{\alpha}/\)H9254 but was retained to a lesser degree (Fig. 1C, gray). These data reveal that PKC\(_{\alpha}\) interacts with mitochondria via a unique mechanism distinct from the canonical C1-mediated binding to DAG or phorbol esters in the membrane.

**FIGURE 1.** Intermolecular FRET demonstrates that PKC\(_{\alpha}\) interacts with mitochondria via an isozyme-specific mechanism. A, diagram of intermolecular FRET translocation reporter, in which an increase in the FRET signal between mitochondrially targeted CFP and YFP-tagged PKC constructs indicates PKC translocation to mitochondria. B, pseudocolored images of a COS-7 cell transfected with mitochondrially targeted CFP and YFP-tagged PKC constructs indicates PKC translocation to mitochondria. C, COS-7 cells co-transfected with Mito-CFP and the indicated YFP-tagged PKC isozyme were stimulated with 200 nM PDBu during FRET imaging, which induced a rapid and sustained increase in the FRET/CFP ratio from PKC\(_{\alpha}\), a transient increase from PKC\(_{\beta}\), and no significant change from either PKC\(_{\alpha}/\)II or PKC\(_{\alpha}/\)I. Throughout this paper, FRET ratios are plotted as the means ± S.E. of data from the specified n number of cells imaged over at least three independent experiments. D, representative CFP and YFP channel time lapse images of COS-7 cells co-transfected with Mito-CFP and the indicated YFP-labeled PKC constructs and monitored for FRET ratio changes in response to PDBu stimulation. The specified times correspond to time points on the FRET ratio plots in C and indicate images captured at the beginning (0 min), the end (15 min), and, for PKC\(_{\beta}\), the FRET ratio high point (7.25 min). The images for PKC\(_{\alpha}\) and PKC\(_{\beta}\) visually show their translocation to mitochondria after PDBu stimulation. The images for PKC\(_{\alpha}/\)II and PKC\(_{\alpha}/\)I show their PDBu-responsive translocation to other intracellular membranes, despite their lack of FRET responses at mitochondria. E, schematic of the PKC isoforms and PKC\(_{\alpha}\) constructs used throughout the paper.

Regulatory Domains of PKC\(_{\alpha}\) Are Necessary but Not Sufficient for PDBu-induced Translocation to and Retention at Mitochondria—To address the isozyme-specific mechanism by which PKC\(_{\alpha}\) binds mitochondria, we investigated the determinants necessary for the interaction (Fig. 1E). Deletion of the C2 domain of PKC\(_{\alpha}\) resulted in a transient association of PKC\(_{\alpha}\) with mitochondria: the half-time for mitochondrial translocation (~30 s) was similar to that for wild-type enzyme, but the construct was released from the membrane with a half-time of ~2 min (Fig. 2A, closed symbols, and supplemental Fig. S1). Thus, the C2 domain of PKC\(_{\alpha}\), although not required for its initial translocation to mitochondria, is necessary for its retention there. In contrast, deletion of the C2 domain had no effect on PKC\(_{\alpha}\) translocation to and retention at the Golgi (Fig. 2A,
open circles and triangles). The isolated δ2 domain did not translocate to mitochondria (Fig. 2A, diamonds), indicating that the role of the δ2 domain in retaining PKCδ at mitochondria is not sufficient to bring it to mitochondria. These data reveal that the C2 domain of PKCδ contributes to the retention of PKCδ at mitochondria.

Deletion of the C1A domain, the C1B domain, or both C1 domains abolished the translocation of PKCδ to mitochondria (Fig. 2B, upper panel, and supplemental Fig. S1), indicating that both C1 domains of PKCδ are necessary for its translocation to mitochondria. In contrast, only the C1B domain is required for the translocation of PKCδ to plasma membrane: deletion of the C1A domain alone had no effect on the interaction of PKCδ with plasma membrane (Fig. 2B, lower panel). Neither the isolated δC1B domain nor a chimera of PKCɛ with its εC1B domain substituted with the δC1B translocated to mitochondria (Fig. 2C and supplemental Fig. S1), indicating that the δC1B domain, despite being the primary phorbol ester-binding domain of PKCδ (40), is not sufficient to promote PKCδ PDBu-induced interaction with mitochondria. All of these regulatory domain deletion mutants of PKCδ were properly primed by phosphorylation at the activation loop, turn motif, and hydrophobic motif (Fig. 2D). Finally, neither the regulatory fragment (RF) of PKCδ (fused with the catalytic fragment of PKCɛ to form a properly folded and phorbol ester-sensitive full-length PKCδ/ɛ chimera) nor its catalytic fragment (CF) alone were capable of translocating to mitochondria (Fig. 2E and supplemental Fig. S1), indicating that at least one determinant outside the RF is necessary for the interaction of PKCδ with mitochondria.

Critical Residues in the Activation Loop and Turn Motif—PKCδ and PKCθ have a unique acidic residue (E and D, respec-
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five amino acids N-terminal of the activation loop phosphorylation site, where all the other PKC isozymes as well as PKA and Akt possess a glycine (Fig. 3A). Mutation of this glutamate to glycine in PKCδ/E500G modestly delayed its translocation to mitochondria by 30 s but significantly decreased its retention at mitochondria to ∼60% by 10 min after PDBu stimulation (Fig. 3B, solid symbols, and supplemental Fig. S1). In contrast, E500G had no effect on the interaction of PKCδ with other membranes such as Golgi (Fig. 3B, open symbols).

To determine whether the charge or the size of this Glu plays the most critical role in PKCδ interactions with mitochondria, we also mutated this residue to either an aspartate (E500D) or a glutamine (E500Q). The E500D mutation, which retains the negative charge of Glu but slightly decreases the size of the residue, caused little loss of retention at mitochondria, although, like E500G, it also translocated after a 30-s delay (Fig. 3C, squares). Thus, the smaller but negatively charged Asp can mostly compensate for the role of Glu-500 in retention. This is consistent with the ability of PKCδ, which possesses an Asp at this position, to interact somewhat with mitochondria (Fig. 1C). Also, mutation of the Asp on PKCδ to a Glu did not improve its retention at mitochondria (data not shown). The E500Q mutation, which retains the size of Glu but loses the charge at this residue, caused a loss of retention similar to that of E500G, without delaying translocation to mitochondria (Fig. 3C, diamonds). Thus, the negative charge of Glu-500 is responsible for its role in retention of PKCδ at mitochondria, whereas the bulk of Glu-500 contributes slightly to the initial affinity of PKCδ for mitochondria. When both the negative charge and bulk of Glu-500 were lost in E500G, PKCδ both translocated...
with a slight delay and partially lost retention at mitochondria (Fig. 3C, triangles).

PKCβ and PKCθ also possess a unique Leu-Asn sequence near the turn motif, where most of the other PKC isozymes possess a Thr-Arg sequence (Fig. 3A). Mutation of these two residues (denoted L636T/N637R or LN/TR) resulted in a complete loss of PDBu-induced interaction by PKCβ/H9254 with mitochondria (Fig. 3D, closed symbols, and supplemental Fig. S1), highlighting the importance of these two residues in the isozyme-specific interaction of PKCβ/H9254 with mitochondria upon activation. Indeed, FRET between mitochondria and the LN/TR mutant actually decreased upon stimulation, indicating the translocation of PKCβLN/TR away from mitochondria in preference for other PDBu-containing membranes and therefore implying the basal localization of PKCβLN/TR to mitochondria. In contrast, the LN/TR mutation did not affect the interaction of PKCβ/H9254 with Golgi (Fig. 3D, open symbols).

Despite the proximity of these mutations to conserved phosphorylation sites, these activation loop and turn motif mutants were fully phosphorylated at the activation loop, turn motif, and hydrophobic motif (Fig. 3E). Thus, their loss of interactions with mitochondria cannot be attributed to the loss of processing phosphorylations.

Pharmacological Inhibition Demonstrates That PKCβ Interaction with Mitochondria Is Dependent on Novel PKC Activity—We next addressed whether the catalytic activity of PKCβ is necessary for its interaction with mitochondria. Pretreatment of cells with either the active site inhibitor Gö6983 or the substrate-uncompetitive inhibitor BisIV (41), both general PKC inhibitors that inhibit conventional and novel PKC isozymes, including PKCβ/H9254, caused both a dramatic reduction in the PDBu-induced translocation of PKCβ to mitochondria and a loss of retention reminiscent of deletion of the C2 domain (Fig. 4A, light blue symbols, and supplemental Fig. S1, Gö6983). In contrast, pretreatment with the conventional PKC isozyme-specific inhibitor Gö6976, which does not inhibit PKCβ, had no effect on mitochondrial translocation (Fig. 4A, dark blue). In contrast to its effects on mitochondrial translocation, pharmacological inhibition with Gö6983 did not affect the interaction of PKCβ with Golgi membranes (Fig. 4B). Pretreatment of cells with increasing concentrations of Gö6983 unveiled a dose-responsive inhibition of the phorbol ester-stimulated interaction
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Specific Inhibition of PKCδ Activity Prevents Its Retention at Mitochondria—To determine whether the intrinsic catalytic activity of PKCδ, rather than the activity of another inhibitor-sensitive novel PKC isoform, is necessary for its interaction with mitochondria, we took advantage of chemical genetics. We generated a PKCδ gatekeeper mutant by mutating a conserved bulky methionine residue in the ATP-binding pocket of PKCδ to a smaller alanine residue (PKCδ-M425A) (42, 43). This enlarges the active site to accommodate the small molecule inhibitor NaPP1, which cannot bind wild-type kinases (44). We then used CKAR, a genetically encoded, FRET-based general PKC activity reporter (35, 36), and δCKAR, a similar reporter specific for PKCδ activity (34), to monitor the activity of endogenous and overexpressed PKCδ (42). We generated a PKCδ gatekeeper mutant by mutating a conserved bulky methionine residue in the ATP-binding pocket of PKCδ to a smaller alanine residue (PKCδ-M425A) (42, 43). 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subsequent inhibition by NaPP1. Additionally, as measured using \( \text{H9254} \CKAR \), pretreatment with NaPP1 completely blocked the PDBu-stimulated activity of the gatekeeper mutant of \( \text{PKC}/\text{H9254} \), without preventing the activation of exogenous wild-type \( \text{PKC}/\text{H9254} \) (Fig. 5C). Having ascertained the specificity of our approach, we then used the gatekeeper mutant of \( \text{PKC}/\text{H9254} \) in our intermolecular FRET translocation assay. Specific inhibition of \( \text{PKC}/\text{H9254} \) gatekeeper mutant activity by pretreatment of cells with NaPP1 prevented its retention at mitochondria (Fig. 5D) without affecting its interaction with the plasma membrane (Fig. 5E).

We chose to use this chemical genetics approach to specifically inhibit \( \text{PKC}/\text{H9254} \) activity because of the lack of reliable \( \text{PKC}/\text{H9254} \)-specific inhibitors. Rottlerin has been used commonly in the literature as a \( \text{PKC}/\text{H9254} \)-specific inhibitor, but this has been refuted, and the effects of rottlerin have been attributed to its role as a mitochondrial uncoupler rather than as a \( \text{PKC}/\text{H9254} \) inhibitor (45, 46). Here, we provide further evidence using \( \text{CKAR} \) that, in contrast to Gö6983, a bona fide PKC inhibitor capable of inhibiting \( \text{PKC}/\text{H9254} \), pretreatment with rottlerin does not inhibit phorbol ester–stimulated \( \text{PKC}/\text{H9254} \) activity in cells (Fig. 5F).

**FIGURE 6.** Mitochondrially targeted \( \text{CKAR} \) reveals that \( \text{PKC}/\text{H9254} \) is active at mitochondria. A, COS-7 cells co-transfected with either Mito-\( \text{CKAR} \) or the negative control reporter Mito-\( \text{CKART/A} \) and either \( \text{PKC}/\text{H9254}-\text{RFP} \) or RFP were monitored for their FRET ratio responses to stimulation with 200 nM PDBu and inhibition with 250 nM Gö6983. B, C, and E, COS-7 cells co-transfected with Mito-\( \text{CKAR} \) and the indicated RFP-tagged \( \text{PKC}/\text{H9254} \) constructs were monitored for their FRET ratio responses to stimulation with 200 nM PDBu. D, COS-7 cells co-transfected with either Mito-\( \text{CKAR} \) or Mito-\( \text{CKART/A} \) and either \( \text{PKC}/\text{H9254}-\text{RFP} \) or \( \text{PKC}/\text{H9254/C1A-RFP} \) were titrated with the indicated increasing net concentrations of Gö6983.
Isozyme-specific Interaction of PKCδ with Mitochondria

FIGURE 7. Mitochondrial membrane potential modulates the retention of PKCδ at mitochondria. A, COS-7 cells co-transfected with either Mito-CFP or Golgi-CFP and PKCδ-YFP were pretreated with 500 nM of the mitochondrial uncoupler FCCP for 15 min at room temperature and compared with unpretreated controls in their FRET ratio responses to 200 nM PDBu stimulation. B, COS-7 cells co-transfected with Mito-δCKAR and PKCδ-RFP were pretreated with 500 nM FCCP for 30 min at 37 °C and compared with the unpretreated control in their FRET ratio responses to 200 nM PDBu stimulation.

caused by phosphorylation of the phospho-acceptor threonine in the substrate of the reporter peptide by exogenous PKCδ, because the negative control Thr to Ala (denoted T/A) version of the reporter did not respond significantly to phorbol ester treatment (Fig. 6A, red triangles) and because the reporter could not detect any significant endogenous PKCδ activity (Figs. 6A, gray circles, and 5, B and C). The portion of the Gö6983-induced reversal in FRET ratio that drops below baseline must be attributed to the reddish-yellow color of this inhibitor, which produces an artifactual FRET response observed even with the negative control reporter Mito-δCKART/A (Fig. 6A, red triangles).

Perturbations that had caused a loss of retention by PKCδ at mitochondria, such as deletion of the C2 domain, the mutation E500G, or inhibition of PKC activity by Gö6983, resulted in overall lower levels of PDBu-stimulated activity at mitochondria (Fig. 6B) and, in the case of Gö6983 pretreatment, a loss of sustained activity there (Fig. 6B, open blue circles). Deletion of either the C1B domain alone or both C1 domains, which had abolished PDBu-stimulated translocation by PKCδ to mitochondria, similarly abolished PDBu-stimulated PKCδ activity at mitochondria (Fig. 6C, orange diamonds and dark orange squares). PKCδ with the C1A domain alone deleted, however, unexpectedly retained some activity at mitochondria (Fig. 6C, light orange triangles), despite its lack of translocation there. This activity can be attributed to a higher basal localization by PKCδΔC1A to mitochondria, which was revealed by directly using Gö6983 to inhibit basal levels of PKCδ activity during FRET imaging. As previously noted, the addition of the reddish-yellow Gö6983 during imaging produces an artifactual below base-line drop in the FRET ratio, and increasing concentrations of Gö6983 produced the same magnitude of artifactual drops in FRET ratio in both Mito-δCKAR and Mito-δCKART/A in the presence of wild-type PKCδ-RFP (Fig. 6D, black and red circles). Nonetheless, the significantly greater inhibitor-induced drops in FRET ratio in the presence of PKCδΔC1A (Fig. 6D, light orange triangles) compared with wild-type PKCδ indicate that deletion of the C1A domain increases the basal activity of PKCδ at mitochondria, which is also corroborated by the greater basal localization of PKCδΔC1A at mitochondria evident in images of the fluorescently labeled construct (supplemental Fig. S1).

Similarly, the LN/TR mutant of PKCδ, which had failed to translocate to mitochondria, exhibited some activity at mitochondria (Fig. 6E), which can also be attributed to its basal localization at mitochondria, as evidenced from its translocation away from mitochondria in preference for other membranes in response to PDBu stimulation (Fig. 3D).

Mitochondrial Membrane Potential Modulates the Retention of PKCδ at Mitochondria—Pretreatment of cells with 500 nM FCCP, a mitochondrial uncoupler that dissipates the mitochondrial membrane potential by acting as a protonophore, resulted in a partial loss of retention by PKCδ at mitochondria following PDBu-stimulated translocation (Fig. 7A, closed symbols, and supplemental Fig. S1), whereas it had no effect on the retention of PKCδ at Golgi (Fig. 7A, open symbols). Similarly, FCCP pretreatment caused PDBu-stimulated activity by PKCδ at mitochondria to be lower and less sustained (Fig. 7B). These data are consistent with the mitochondrial membrane potential contributing to PKCδ retention at mitochondria.

Independent versus Coincident Mechanisms of PKCδ Interaction with Mitochondria—The foregoing results identified four perturbations that affect the retention of PKCδ at mitochondria: deletion of the C2 domain (PKCδΔC2), Gö6983 pretreatment, FCCP pretreatment, or removal of the negative charge at Glu-500 (PKCδE500G). To resolve whether these perturbations affect the same mechanism for retaining PKCδ at mitochondria or whether they affect independent mechanisms, we tested every one of the six possible perturbation pairings to determine whether the combined effect of each pair produced either an additive/synergistic or nonadditive effect on the retention of PKCδ at mitochondria. An additive or synergistic effect of two perturbations would suggest simultaneous disruption of two independent determinants driving the interaction of PKCδ with mitochondria. A lack of additivity or synergism would indicate that the two perturbations had coincided in affecting a single determinant driving the interaction of PKCδ with mitochondria. Using this type of analysis, we determined that the mechanism of retention regulated by Glu-500 is part of the same mechanism regulated by the C2 domain (Fig. 8A), PKCδ activity (Fig. 8B), or the mitochondrial membrane potential (Fig. 8C). In contrast, the mechanisms of retention regulated by the C2 domain, PKCδ activity, and the mitochondrial
membrane potential are all independent mechanisms (Fig. 8, D–F). By combining the latter three independent mechanisms of retention, we completely abolished the PDBu-stimulated interaction of PKC\(\delta/H9254\) with mitochondria, actually causing PKC\(\delta/H9254\) basally localized to mitochondria to translocate away from mitochondria in preference for other membranes (Fig. 8 G, red circles).

**Functional Effects of PKC\(\delta/H9254\) Interaction with Mitochondria**

We next examined whether the translocation of PKC\(\delta/H9254\) to mitochondria is triggered by a natural agonist. Stimulation of L6 skeletal muscle cells with 100 nM insulin resulted in the translocation of PKC\(\delta/H9254\), but not PKC\(\epsilon/H9252\) or PKC\(\theta\), to mitochondria; this translocation was further increased following phorbol ester stimulation, but again only for PKC\(\delta\) (Fig. 9A). These data reveal that a natural agonist also causes the isoform-specific recruitment of PKC\(\delta\) to mitochondria.

To investigate the physiological relevance of the PKC\(\delta\)-specific interaction with mitochondria, we measured the effect of PKC\(\delta\) overexpression on the OCR of intact COS-7 cells. Overexpression of PKC\(\delta/H9254\) but not of PKC\(\epsilon/H9252\), PKC\(\theta\), or PKC\(\delta\) resulted in a significant increase in the OCR and thus the mitochondrial respiration of these cells relative to a vector control (Fig. 9B). In contrast, PKC\(\theta\) overexpression significantly decreased the cellular OCR (Fig. 9B). Differences in OCR between groups were observed regardless of whether the basal (data not shown), State 4 (data not shown), or uncoupler-stimulated rates (Fig. 9B) were measured, but differences were most pronounced in the uncoupler-stimulated rates.

Structural alterations that had perturbed the interaction of PKC\(\delta\) with mitochondria all resulted in a loss of the increase in OCR produced by overexpression of wild-type PKC\(\delta\) (Fig. 9B). Mitochondrial respiration upon overexpression of PKC\(\delta/\Delta C2\)
was no different from that of vector control (Fig. 9B). Overexpression of PKCΔC1B, PKCΔC1A, or PKCδE500G actually had the dominant negative effect of reducing mitochondrial respiration below vector control levels (Fig. 9B). Overexpression of PKCΔC1B or PKCδLN/TR also tended to reduce mitochondrial respiration below that of vector control but did not reach statistical significance (Fig. 9B). All of the constructs were successfully overexpressed in these cells, as determined by Western blotting (data not shown). Differences in mitochondrial respiration did not result from differences in cellular proliferation, because the same cells plated in triplicate in parallel with the OCR measurements did not exhibit significant differences in proliferation, as determined by cell counts (data not shown). Thus, the isozyme-specific structural ability of PKCδ to interact with mitochondria has the functional consequence of increasing mitochondrial respiration.

DISCUSSION

Using an intermolecular FRET translocation reporter and an intramolecular FRET PKCδ activity reporter to monitor PKC translocation to and PKCδ activity at mitochondria in real time in live cells, we show that PKCδ translocates rapidly to and is active at the outer membrane of mitochondria upon activation with phorbol esters or, in L6 myocytes, with insulin. The mechanism of this translocation is distinct from that of the canonical C1 domain-mediated recognition of DAG or phorbol esters that drives conventional and novel PKC isozymes to the plasma membrane and Golgi. First, the interaction of PKCδ with mito-
PKCδ stands out among the PKC isozymes in two ways. First, it is structurally different: it has a unique activation loop containing a phosphomimetic Glu five residues N-terminal to the phosphorylated Thr that may partly account for its ability to be activated without phosphorylation by phosphoinositide-dependent kinase-1 (47, 48) and whose phosphorylation is agonist-dependent as well as constitutive (49, 50), and it has a unique turn motif that may account for its lack of regulation by mTORC2 (13). These structural features are shared by only one other isozyme, PKCθ. Second, it translocates and is activated by two mechanisms: the canonical lipid-dependent activation at intracellular membranes (Fig. 10, arrow 1) and a novel tyrosine kinase-dependent (and phorbol ester-independent) activation (50–53) that causes translocation into the nucleus (4, 9–11, 34) and plays a role in apoptosis (4, 9–11, 52) (Fig. 10, arrow 2). Here, we unveil a third unique property of PKCδ: this isozyme uniquely interacts with and is activated at mitochondria, where it promotes mitochondrial respiration (Fig. 10, arrow 3).

The phorbol ester-stimulated interaction of PKCδ with mitochondria occurs in two steps and depends on multiple parameters (Fig. 10). First, both C1 domains and a unique LN sequence in the turn motif are required for the initial recruitment of PKCδ to mitochondria (Fig. 10, Step A). Fluorescently tagged phorbol esters have been visualized to distribute to mitochondria (54), and we speculate that the interaction between the C1B domain of PKCδ and phorbol esters distributing to mitochondria helps to bring other determinants on PKCδ into contact with mitochondria. Alternatively, PKCδ is capable of binding PDBu in the absence of membrane lipids, albeit with lower affinity (55), and this interaction may directly drive the interaction of PKCδ with nonmembrane targets at mitochondria; there is precedent for phorbol ester-mediated binding by other PKC isozymes to nonmembrane structures (56) and for C1 domains to mediate protein, as well as lipid, interactions (57). Second, the C2 domain, its intrinsic catalytic activity, and the mitochondrial membrane potential independently regulate the sustained interaction of PKCδ at mitochondria, whereas the charge of Glu-500 on PKCδ cooperates with all of the latter three mechanisms to retain PKCδ at mitochondria (Fig. 10, Step B). Interestingly, phorbol esters have been shown to alter the mitochondrial membrane potential in PKCδ-overexpressing keratinocytes (23) and to potentiate and accelerate the effect of a mitochondrial ATP-dependent K⁺ channel opener (58). In contrast, only the C1B domain among all of these is required for the phorbol ester-stimulated translocation of PKCδ to other intracellular membranes such as the plasma membrane and Golgi (this paper and Ref. 40).

Stimulation with phorbol esters or with insulin in L6 myocytes enables us to observe an interaction between PKCδ and mitochondria because it produces an acute change in the FRET ratio. However, our data also support a model in which a significant amount of PKCδ associates basally with mitochondria via many of the same determinants. First, the C1B domain of PKCδ is basally localized to mitochondria, because FRET between Mito-CFP and ΔC1B-YFP decreases upon phorbol ester stimulation (Fig. 2C), reflecting translocation of the isolated domain basally localized to mitochondria away to other membranes. Second, full-length PKCδ also appears to be basally localized to mitochondria, because pretreatment with greater than 750 nM Gö6983 (Fig. 4C) or combining multiple perturbations that disturb the retention of PKCδ at mitochondria (Fig. 8) not only brings its phorbol ester-stimulated FRET ratio with Mito-CFP back down to base line but further below base line, indicating disturbance of its basal interaction with mitochondria. In contrast, the C1A domain of PKCδ may block its basal interaction with mitochondria, because deletion of the C1A domain

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FIGURE 10. Model showing three mechanisms for the agonist-evoked redistribution of cytosolic PKCδ to intracellular compartments and summarizing the determinants involved in its interaction with mitochondria. Arrow 1, canonical binding of PKCδ to DAG-containing membranes such as the plasma membrane (shown) and Golgi is mediated by the C1B domain. Arrow 2, Src-dependent translocation of PKCδ to the nucleus depends on Tyr phosphorylation (indicated by red circles) of PKCδ. Arrow 3, the novel, isozyme-specific interaction of PKCδ with mitochondria occurs via two steps: step A, an initial recruitment step that depends on the C1A and C1B domains and on Leu-Asn (LN) in the turn motif, and step B, a retention step that depends on four elements: the C2 domain, an acidic residue in the activation loop (Glu-500, E500), intrinsic catalytic activity (represented by ATP → ADP), and the mitochondrial membrane potential (represented by plus and minus signs). One or more protein scaffolds (gray rectangles) at mitochondria are hypothesized to mediate this interaction, which promotes mitochondrial respiration.
increases the extent of basal PKCδ activity at mitochondria, as revealed by titration with inhibitor (Fig. 6D) and by the low level of activity induced by phorbol esters (Fig. 6C) despite the lack of PKCδΔC1A translocation (Fig. 2B). By comparison, depletion of the C1B domain alone or in combination with C1A results in no translocation to mitochondria (Fig. 2B) and no basal phorbol ester-induced activity at mitochondria (Fig. 6C). The C1A domain may basally recruit PKCδ to other membranes or mask the basal interactions of the neighboring C1B domain with mitochondria. In a manner similar to PKCδΔC1A, the turn motif mutant PKCδLN/TR also appears to be basally localized to mitochondria, translocating in net away from mitochondria upon PDBu stimulation (Fig. 3D) yet still exhibiting some activity at mitochondria from the basally localized pool (Fig. 6E). Thus, the C1B and C2 domains, its intrinsic catalytic activity, as well as the mitochondrial membrane potential may contribute to the basal interaction of PKCδ with mitochondria, whereas the C1A domain and the LN motif may interfere with it.

Consistent with PKCδ localizing basally to mitochondria, we found that unstimulated PKCδ overexpression increases mitochondrial respiration (Fig. 9B). This is consistent with reports that PKCδ activates the PDHC, which converts pyruvate to acetyl CoA, thereby increasing mitochondrial respiration. Caruso et al. (27) showed that insulin stimulation of L6 skeletal muscle cells and immortalized hepatocytes induces PKCδ to translocate to mitochondria and to phosphorylate PDH phosphatases 1 and 2, which then dephosphorylate and activate the PDHC, increasing mitochondrial respiration. The work of Hammerling and co-workers (28, 29) demonstrated in mouse embryonic fibroblasts that retinol, or vitamin A, induces a PKCδ signalosome composed of retinol, PKCδ, cytochrome c, and the adapter protein p66Shc to dephosphorylate and inhibit PDH kinase 2, which usually phosphorylates and inactivates the PDHC, thereby leading also to the activation of the PDHC and increased mitochondrial respiration. In addition, we found that overexpression of PKCe, which has been shown to oppose both the apoptotic (2) and metabolic (30) effects of PKCδ, actively decreases mitochondrial respiration relative to vector control (Fig. 9B), consistent with its proposed role of inhibiting the PDHC (30). Finally, none of the PKCδ mutant constructs that had been defective in phorbol ester-induced interaction with mitochondria produced the same increase in oxygen consumption rate as wild-type PKCδ (Fig. 9B), indicating that the structural elements important for the interaction of PKCδ with mitochondria are also important for its effect on mitochondrial respiration. Indeed, PKCδΔC1AΔC1B, PKCδΔC1A, and PKCδE500G actually significantly reduced mitochondrial respiration below vector control levels in a manner similar to PKCe (Fig. 9B). One possibility is that these constructs that interact imperfectly with mitochondria produce dominant negative effects by mislocalizing or displacing endogenous PKCδ from mitochondria.

It remains to be determined whether PKCδ produces its effect on mitochondrial respiration by remaining on the cytoplasmic surface of the mitochondrial outer membrane and simply transducing signals to substrates on the surface of mitochondria (e.g., by phosphorylating intermediate phosphatases that are then translocated into the mitochondrial matrix, where they regulate the PDHC) or whether PKCδ is itself subsequently translocated into the mitochondria, possibly by protein partners that mediate the retention step of its interaction with mitochondria. In addition to the kinase and phosphatases that regulate the PDHC, other potentially relevant metabolic substrates of PKCδ that have been noted in the literature include the d subunit of ATP synthase in the inner membrane of mitochondria (in neonatal cardiac myocytes exposed to hypoxia or phorbol ester (59)) and the M2 splice isoform of pyruvate kinase (in MCF-7 cells (60)), which plays an important role in the metabolic switch from the use of glucose in oxidative phosphorylation to its use in aerobic glycolysis in tumors (61). These latter two mechanisms would not likely be at play in the increased rates of uncoupler-stimulated respiration that we observed.

Past studies of PKCδ signaling to mitochondria have used kinase-dead constructs of PKCδ and pharmacological inhibition with rottlerin to attempt to establish the need for catalytic activity. Some studies have even used Gö6976 as a PKCδ inhibitor (28, 29), whereas Gö6976 is actually a conventional-selective PKC inhibitor that does not inhibit novel PKCs even at micromolar concentrations (62). Our data and others in the literature show rottlerin not to be a PKCδ-specific inhibitor, with its effects being attributable to its role as a mitochondrial uncoupler (45, 46). We have also found that kinase-dead constructs of PKCδ, especially the K376R mutant used predominantly in the literature, are not fully primed by phosphorylations (data not shown) and therefore may not be properly folded and competent for activation. This study uses the more reliable pharmacological tools of Gö6983 and BisIV as general PKC inhibitors (Fig. 4) and the chemical genetics approach of a gatekeeper mutant of PKCδ shown by our data to be specifically inhibited by NaPP1 (Fig. 5). Furthermore, live cell FRET imaging is a more sensitive and informative technique for evaluating molecular interactions than biochemical techniques such as pull downs, fractionations, and immunofluorescence because it can detect and quantify transient, low affinity, dynamic interactions in real time and in a more physiological cellular context and can be used to determine the kinetics of an interaction.

Although this study was performed in a simplified system that employed overexpression of PKCδ and primarily phorbol ester stimulation, the mechanisms revealed by this study should be generalizable to more physiological systems. First, PKCδ overexpression was ~4-fold that of endogenous PKCδ levels in transfected COS-7 cells by Western blot analysis (data not shown). Second, PKCδ is basally localized to mitochondria via many of the same determinants that drive its phorbol ester-stimulated translocation. Third, unstimulated PKCδ overexpression increases mitochondrial respiration, and mutation of any of the structural elements that play a role in the phorbol ester-induced interaction of PKCδ with mitochondria abolishes this basal effect, with some mutations even further reducing respiration. Fourth, stimulation of myocytes with insulin, a physiological agonist, produces the same isozyme-specific interaction of PKCδ with mitochondria.

Our data support a model in which multiple determinants unique to PKCδ, involving its C2, C1A, and C1B domains, critical residues in its activation loop and turn motif, its kinase activity, and the mitochondrial membrane potential, contrib-
ute to a specific interaction with the outer membrane of mitochondria, where PKCδ is active, thereby regulating mitochondrial respiration (Fig. 10). Given the noncanonical nature of the interaction between PKCδ and the mitochondrial outer membrane and the participation of multiple determinants unique to PKCδ, we speculate that one or more protein scaffolds at the mitochondria mediate this isoform-specific interaction.

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