Intracellular zinc increase affects phosphorylation state and subcellular localization of protein kinase C delta (δ)

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A B S T R A C T
Protein kinase C delta (PKCδ) is a Ser/Thr-specific kinase involved in many fundamental cellular processes including growth, differentiation and apoptosis. PKCδ is expressed ubiquitously in all known cell types, and can be activated by diacylglycerol, phorbol esters and other kinases. Multiple lines of evidence have indicated that the mode of activation greatly influences the role PKCδ plays in cellular function. Divalent metal ions, such as zinc, are released as a response to cellular stress and injury, often resulting in oxidative damage and cell death. In this study, we evaluate the effect increased concentrations of intracellular zinc has on the phosphorylation state and subcellular localization of PKCδ. More specifically, we demonstrate that intracellular zinc inhibits the phosphorylation of PKCδ at Thr505 in a concentration-dependent manner and facilitates the translocation of PKCδ from the cytosol to the Golgi complex. Analysis of a PKCδ structural model revealed a potential His-Cys3 zinc-binding domain adjacent to residue Thr505 and suggests that interaction with a Zn2+ ion may preclude phosphorylation at this site. This study establishes zinc as a potent modulator of PKCδ function and suggests a novel mechanism by which PKCδ is able to “sense” changes in the concentration of intracellular zinc. These findings illuminate a new paradigm of metal ion-protein interaction that may have significant implications on a broad spectrum of cellular processes.

1. Introduction

Protein kinase C delta (PKCδ) is a member of the PKC family of Ser/Thr kinases that are expressed in brain, heart, spleen, lung, liver, ovary, pancreas and adrenal gland [1, 2]. PKCδ activity has been implicated in a wide range of cellular processes including proliferation [3, 4], differentiation [5] and apoptosis [6-8]. Structurally, PKCδ consists of a C-terminal catalytic domain and N-terminal regulatory domain [9, 10]. Together with PKC ε, η and θ, PKCδ is classified as a ‘novel kinase’, which can be activated by diacylglycerol (DAG) or phorbol esters and is calcium-independent [11]. PKCδ is the most well-studied novel protein kinase, and it has recently been discovered that PKCδ can have both pro- and anti-apoptotic function [6, 12]. Despite these advancements however, a consensus on the functional nature of PKCδ remains to be established. This disparity likely exists because the activity of PKCδ is highly dependent on cell type, phosphorylation state, external stimuli and intracellular localization [10, 13, 14].

The biological functions of proteins within the PKC family are primarily controlled by their subcellular localization [13, 15, 16] and phosphorylation state [17]. Aside from being activated by DAG and phorbol esters, PKCδ can also be activated by other kinases. Regulation of PKCδ function by phosphorylation is quite complex [17]; PKCδ has been shown to be phosphorylated at up to 17 sites, including eight Ser/Thr residues and nine Tyr residues [18]. Several of these sites, such as Ser43 and Ser462, remain constitutively phosphorylated and have little known effect of PKCδ function [18]. Conversely, phosphorylation of PKCδ within its C-terminal domain activation loop at Thr505 has been associated with increased kinase activity [18]. PKCδ is unique from all the other kinases in the novel kinase subfamily, because it does not require phosphorylation at Thr505 for catalytic function [19]. These observations have led to speculation that phosphorylation at Thr505 (Thrφ505) is used to fine-tune PKCδ substrate specificity [19, 20]. The complex nature of PKCδ phosphorylation may explain the multifaceted and stimulus-dependent functions of PKCδ, making it an interesting target for investigation.

Zinc is the most abundant intracellular trace mineral, being found in the cytosol, organelles and nucleus of all cell types [21]. Zinc is required for the function of hundreds of enzymes [22] and is crucial for

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stabilizing the folded structure of many non-enzymatic proteins [23]. Zinc is unique among transition metals in that it is redox inert in biological systems and has only one valence state: Zn(II) [24]. The electron donors such as oxygen, nitrogen and sulfur limit zinc interactions with proteins [24]. As a cofactor, zinc may remain bound during the lifetime of proteins that rely on it for catalytic function. Other proteins, however, may bind free zinc reversibly with dissociation rates corresponding with functional requirements. These dissimilar binding attributes allow zinc to play highly important roles in cell physiology [25, 26]. Despite its inert redox potential, concentrations of intracellular free zinc are tightly regulated through organelle compartmentalization [27] and binding to metallothioneins [25, 28]. This high level of control is achieved by transmembrane proteins that transport zinc in and out of intracellular compartments. Proteins such as Zrt and Irt-like protein function to transport zinc ions (Zn2+) into the cytosol from outside the cell or from vesicles, whereas proteins such as ZnT control the efflux of Zn2+ from the cytoplasm to outside the cell or into vesicles [29]. While zinc is considered an essential element required for normal physiological function, dysregulated zinc homeostasis has been implicated in pathogenesis of human disease. For example, zinc deficiencies can result in oxidative damage to DNA and lead to increased cancer risk [30]. Furthermore, inefficient zinc transport has been linked to the onset of neurodegenerative diseases [31, 32], depression [33, 34] and type II diabetes [35, 36]. Finally, numerous reports have shown that concentrations of intracellular zinc increase during hypoxia [37–40] and that such increases in zinc can precede cell death in neurons [39]. More recently, zinc has emerged as a potent modulator of protein phosphorylation in cells. Indeed, it has been shown that increases in intracellular zinc concentrations can facilitate the phosphorylation of proteins such as tau [41], insulin-like growth factor receptor [42], and insulin-like growth factor-1 [43]. The inhibitory effects of zinc on protein phosphorylation have also been reported [44] however, less is known about how zinc influences this process.

The current study was designed to investigate how increases in intracellular zinc concentrations affect the phosphorylation state and subcellular trafficking of PKCδ. We report here that increasing concentrations of intracellular free zinc significantly inhibits phosphorylation of PKCδ at a key phosphorylation site: Thr505, in a concentration-dependent manner; and influences PKCδ translocation from cytosol to the Golgi complex. We also employed molecular modeling software to gain additional insight into how zinc might affect the structure of PKCδ. These studies show that zinc may bind to a novel His-Cys3 binding pocket that is adjacent to Thr505. We therefore propose a mechanism by which PKCδ is able to ‘sense’ concentrations of intracellular zinc by coordinating it to a His-Cys3 binding site. Importantly, this binding event repositions His197 so that it blocks access to Thr505, effectively inhibiting phosphorylation at this site. This new paradigm enhances our understanding of zinc-PKCδ interactions and implicates zinc as a potent modulator of PKCδ activity.

2. Materials and methods

2.1. Reagents and chemicals

Unless otherwise stated, all chemicals were purchased from Sigma-Aldrich (St. Louis, MO). Fluorescent zinc dye, FluoZin-3, AM was purchased from Life Technologies (Waltham, MA). Microscopy supplies were purchased from VWR (Radnor, PA). HeLa cells and cell culture reagents were purchased from ATCC (Manassas, VA). Immunoblot materials were purchased from Bio-Rad (Hercules, CA) and Invitrogen (Waltham, MA). Antibodies were from Cell Signaling Technology (Danvers, MA) and Abcam (Cambridge, MA).

2.2. Cell culture

HeLa cells (laboratory passage 4–14) were maintained in Eagle's minimum essential medium (EMEM) supplemented with 5% fetal bovine serum (FBS) under a humidified atmosphere (95%) containing 5% CO2 at 37 °C. Cells were subcultured every other day using standard trypsinization methods, according to ATCC recommendations.

2.3. Detection of intracellular free zinc levels

HeLa cells were seeded at medium density onto glass-bottom Petri dishes (P35G-4.5-14-C; MatTek Corp, Ashland, MA). Following seeding, the cells were allowed to incubate under a humidified atmosphere (95%) containing 5% CO2 at 37 °C for at least 24 h before experimentation. Prior to treatments, the cells were washed three times with warm physiological buffer (25 mM HEPES, 125 mM NaCl, 3 mM KCl, 1.28 mM CaCl2, 1.1 mM MgCl2 and 5 mM glucose, pH 7.4). For zinc detection, FluoZin-3, AM was added to the cell media at final concentration of 1 μM and cells were incubated for 60 min at room temperature. After incubation with the dye, cells were washed three times with physiological buffer and allowed to “rest” at room temperature for 30 min. To ensure removal of incident extracellular dye, the cells were washed one final time with physiological buffer before observation by fluorescence microscopy. For treatments, either pyrithione alone or a combination of pyrithione and zinc chloride was added to the media and the cells were allowed to incubate for 10 min at 37 °C. Final concentrations of pyrithione and zinc chloride were 10 μM and 50 μM respectively. Images were collected using a Motic AE31 microscope outfitted with a QImaging Retiga 1300i camera with a 40×/0.75 Olympus objective. Image-Pro Plus 6.2 (Media Cybernetics) was used to collect and analyze the data. Images were collected every 30 s and zinc was added after 60 s of baseline fluorescence was recorded. To detect changes in intracellular fluorescence, regions of interest (ROI) were selected manually by highlighting areas of the cytosol, excluding the nucleus. Background fluorescence was measured from multiple regions of the same images that contained no cells. Changes in fluorescence (ΔF) were determined using Eq. (1):

$$\Delta F = \frac{F_{\text{measure}} - F_0}{F_0}$$

where $F_{\text{measure}}$ is measured fluorescence and $F_0$ is average fluorescence at baseline before the application of zinc. Each fluorescent measurement was background subtracted.

2.4. PKCδ subcellular localization (western blot)

HeLa cells were seeded at medium density onto 35 mm plastic Petri dishes in 2 mL EMEM supplemented with 5% FBS. Cells were allowed to attach to the plates under a humidified atmosphere (95%) containing 5% CO2 at 37 °C for 24 h before treatments. Immediately prior to treatment, the cells were washed three times with 1 mL of physiological buffer. Depending on experimental conditions, washed cells were then switched to physiological buffer supplemented with either 10 μM pyrithione or 10 μM pyrithione and 50 μM zinc chloride and allowed to incubate for 30 min at 37 °C. All pyrithione and zinc chloride solutions were made fresh before the experiments. Following incubation, the cells were washed with cold PBS and harvested by scraping. The cells were then counted and viable cells were identified using Trypan Blue. Cell number and volumes were adjusted to ensure that control and zinc-treated samples had similar numbers of viable cells. Cells were lysed in cold PBS supplemented with protease inhibitor cocktail (Sigma-Aldrich, P2714) by rapidly passing the cells 20 times through a 27-gauge needle. Lysis efficiency was monitored by observing samples under brightfield microscopy; lysis was deemed complete when no whole cells were observed. Cell homogenates were then centrifuged at 1500 rpm (900 × g) at 4 °C for 10 min to pellet unbroken cells and nuclei. The resulting supernatant was then spun at 55,000 rpm (100,000 × g) for 4 °C for 60 min using an Optima TLX Ultracentrifuge (Beckman Coulter, Brea, CA). The supernatant, which contained soluble cytosolic proteins, was heated to 95 °C for 5 min in 1 × Laemmli buffer to prepare the samples.
for SDS-PAGE. The resultant pellet, which contained organelles such as fragments of the endoplasmic reticulum, Golgi complex, mitochondria and plasma membrane, was washed with PBS supplemented with protease inhibitors and centrifuged again at 55,000 rpm (100,000 × g) for 60 min at 4 °C. Samples were then prepared for SDS-PAGE by heating the resuspended pellet to 95 °C for 5 min in 1 × Laemmli buffer. Subsequent washing and centrifugation of the pellet was performed to reduce contamination by cytosolic proteins. The samples containing whole cell, cytosol and organelle extracts were then loaded onto a 10% polyacrylamide gel and separated by SDS-PAGE using an XCell module (Bio-Rad). Separated proteins were transferred to a 0.45 μm PVDF membrane (Invitrogen) at 300 mA constant voltage for 2 h at room temperature. The membrane was then blocked with 5% bovine serum albumin in tris-buffered saline (TBS) supplemented with 0.1% Tween-20 (TBST) for 2 h at room temperature. Anti-PKCδ [EPR17075] rabbit monoclonal antibody (Abcam, #182126, 1:1000 dilution in 5% BSA/TBST) was used to label PKCδ by incubating the membrane in primary antibody solution overnight with gentle agitation at 4 °C. The membrane was then washed with TBST to remove unbound primary antibody and incubated in a solution containing horse radish peroxidase (HRP)-linked secondary anti-rabbit antibody (Cell Signaling Technology, #7074, 1:3000 dilution in 2% BSA/TBST) for 90 min at room temperature with moderate agitation. Membranes were developed using Clarity Western ECL Substrate (Bio-Rad) and signals were detected on a molecular imaging system, ChemiDoc XRS+ (Bio-Rad) using ImageLab 3.01 software (Bio-Rad). Total protein staining on the blotted membranes was achieved by incubating membranes in Coomassie brilliant blue R-250 staining solution (Bio-Rad) for 1 min at room temperature. Following staining the membrane was immediately washed 3 times with deionized water and subsequently washed 3 times with deionized water for 10 min each. The washed membrane was dried overnight at room temperature in the dark and imaged using a ChemiDoc XRS+ imaging system (Bio-Rad) set to automatic exposure on the colorimetric blot setting using ImageLab 3.01 software (Bio-Rad).

2.5. PKCδ subcellular localization (immunofluorescence)

HeLa cells were trypsinized and seeded at medium density onto sterilized glass coverslips (15 mm, #1, Corning) in 35 mm plastic Petri dishes; three coverslips were used per each dish. Cells were plated in 2 mL EMEM supplemented with 5% FBS and were allowed to attach for 24 h under a humidified atmosphere (95%) containing 5% CO₂ at 37 °C. Prior to treatments, the cells were washed 3 times with 1 mL of physiological buffer. Following washing, the cells were subjected to pyridine and zinc treatments as described above. Once the treatments were complete, the coverslips were removed from the Petri dish and the adhered cells were gently washed with physiological buffer. Following washing the cells were immediately fixed by submerging the cover slips in cold (−20 °C) methanol for 5 min. After fixation, the adhered cells were washed with room temperature PBS and permeabilized by submerging the cover slips in room temperature PBS supplemented with 0.1% Triton X-100 for 5 min. Following permeabilization the coverslips were submerged in blocking solution (1% BSA, 10% normal goat serum, 0.3 M glycine and 0.1% Tween-20 in PBS) for 1 h at room temperature. Once blocked, the cells were treated with antibodies against PKCδ: anti-PKC delta [EPR17075] rabbit monoclonal antibody (Abcam, #182126, lot GR189581-2, 1:100 dilution in blocking solution), or β-COP: anti-β-COP mouse monoclonal clone maD (Sigma, #G6160, 1:400 dilution in blocking solution). The fixed cells were incubated with primary antibodies overnight at 4 °C without agitation. Following incubation, the adhered cells were washed 4 times with PBS supplemented with 0.1% Tween-20 and incubated with appropriate secondary antibodies (1:400 dilution in blocking solution) for 1 h at room temperature in the dark without agitation. Goat anti-rabbit Alexa 488 (Invitrogen) was used for detection of PKCδ and goat anti-mouse Alexa 546 (Invitrogen) was used for detection of β-COP. Excess secondary antibodies were removed by washing the coverslips 4 times with PBS supplemented with 0.1% Tween-20 at room temperature. The cells were then incubated in PBS containing 1 μg/mL of DAPI (nuclear stain) for 5 min at room temperature. Cells were then washed 3 times with PBS supplemented with 0.1% Tween-20. Each coverslip was then mounted on a glass microscope slide (VWR) using 70% glycerol in PBS. Images were collected immediately on Nikon Fluorescent Microphot-SA microscope outfitted with a 100 ×/1.3 Nikon objective with ImageQ camera with Qcapture software.

2.6. PKCδ phosphorylation (western blot)

HeLa cells were plated and maintained as described above. Briefly, cells were plated on 35 mm culture dishes in 2 mL EMEM supplemented with 5% FBS and were allowed to attach for 24 h. Prior to treatments, cells were washed 3 times with physiological buffer and switched to media supplemented with 10 μM pyrithione and 0, 1, 10 or 50 μM zinc chloride. Cells were then allowed to incubate for 30 min at 37 °C. Following incubation, the treatment solutions were removed and the cells were washed once with physiological buffer. Then, 100 μL of 1.5 × Laemmli sample buffer (Bio-Rad) was added to each culture dish and the cells were scrapped to lift them off the plate. Suspended cells were then transferred to 1.5 mL centrifuge tubes and homogenized on ice using an Eppendorf homogenizer (15 rotations, 1 s each). Following homogenization, the samples were immediately heated to 95 °C for 5 min. The lysate was then loaded onto a 10% polyacrylamide gel and proteins were separated by SDS-PAGE electrophoresis using an XCell II module (Bio-Rad). Proteins were then transferred to a 0.45 μm PVDF membrane using techniques as described above. Following transfer, the membrane was blocked with 5% BSA/TBST for 2 h at room temperature. The membranes were then incubated in BSA/TBST containing antibodies against phospho PKCδ (Thr505): anti-phospho PKCδ (pThr505) rabbit polyclonal antibody (Cell Signaling Technology, #9374, lot 6, 1:1000 dilution in 5% BSA/TBST) or total PKCδ: anti-PKCδ rabbit polyclonal antibody (Cell Signaling Technology, #2058, lot 1, 1:1000 dilution in 5% BSA/TBST). The membranes were incubated with the primary antibody solution overnight at 4 °C with gentle agitation. Following incubation, the membranes were washed 3 times with TBST to remove unbound primary antibody. The membranes were then incubated with horse radish peroxidase (HRP)-linked secondary anti-rabbit antibody (Cell Signaling Technology, #7074, 1:3000 dilution in 2%BSA/TBST) for 90 min at room temperature with moderate agitation. The excess secondary antibody was removed by washing the membrane 3 times with TBST with moderate agitation. Membranes were developed and bands were detected as described above. Quantification of the band intensity was performed using ImageLab software (Bio-Rad). Each band was highlighted manually and density values were assigned by the program for each protein band. Density values were also quantified from equivalent areas of the blot containing no protein (background) and this value was subtracted from each density value obtained for protein bands. The ratio of total PKCδ to phospho PKCδ (Thr505) was calculated and plotted as a percent change of the control.

2.7. Molecular modeling

Zinc-binding sites were predicted by uploading the non-redundant human PKCδ sequence (UniProtKB - Q05655) to the ZincExplorer analysis server. Residues with high confidence of zinc-binding included prediction scores between 0.49 and 1 with the higher scores being of greater confidence. To predict the secondary and tertiary structure of the full-length protein, the non-redundant full-length human PKCδ primary sequence (residues 1–676, UniProtKB - Q05655) was uploaded to the PHYRE2 (protein homology/analogy recognition engine V 2.0) server. Zn²⁺ atoms were added to the PKCδ apo-structure using PyMOL.
molecular modeling software (Version 1.8 Schrödinger, LLC). The resulting PDB files were then uploaded into the Swiss PDB Viewer (DeepView/Swiss-PDBViewer, Swiss Institute of Bioinformatics) and energy minimizations were performed using a version of the GROMOS 43B1 force field. Minimized protein structures were then visualized and rendered using PyMOL molecular modeling software.

3. Results

3.1. Concentrations of intracellular zinc increase with addition of exogenous zinc

The capacity to study how zinc affects the biology of PKCδ relied on our ability to systematically raise levels of intracellular zinc in cultured cells. To accomplish this, we labeled HeLa cells with a fluorescent indicator of free zinc: FluoZin-3, AM. This small molecule fluorophore has high affinity for zinc with a dissociation constant (Ki) of approximately 15 nM [45]. FluoZin-3, AM exhibits a 50-fold increase in fluorescence in response to saturating levels of zinc. This indicator can be used to monitor changes in cytosolic free/labile zinc, where an increase in concentration of intracellular free zinc results in cytosolic zinc increase. Furthermore, this technique may be used to simulate cellular responses to pathological conditions that result in cytosolic zinc increase.

3.2. Changes in intracellular zinc affect the subcellular localization of PKCδ

It has been documented that the function of PKCδ is determined by its subcellular localization [13, 15, 46–48]. To investigate whether changes in intracellular free zinc affect the subcellular localization of PKCδ, we treated HeLa cells with 10 μM sodium pyrithione or a combination of 10 μM sodium pyrithione and 50 μM zinc chloride for 30 min and evaluated the subcellular localization of PKCδ by western blot (Fig. 2A). Cells were plated, maintained and treated as described in the Materials and methods. Following treatments, the cells were lysed and subjected to consecutive centrifugation to separate cellular components. For the purpose of this study, the whole cell lysate, before centrifugation was identified as “whole cell”; the supernatant that contained cytosolic proteins was labeled “cytosol”; and the pellet that contained fragments of organelles including the endoplasmic reticulum, Golgi complex, mitochondria and plasma membrane was designated “organelles”. Antibodies specific to PKCδ (Abcam, #182126), were used to visualize the amount of total PKCδ protein in each cellular fraction. As expected, the whole cell lysate showed no significant difference in amount of PKCδ in cells treated with or without zinc (Fig. 2A, upper image). On the other hand, different subcellular localization profiles of PKCδ emerged when the cytosolic and organelle fractions were compared. For example, PKCδ was found to localize to the cytosol in cells that were not exposed to zinc. However, the cytosolic concentration of PKCδ in cells treated with zinc was significantly reduced. It was also observed that PKCδ localizes to cellular organelles even in the absence of zinc, as was indicated by the presence of PKCδ in the organelle fraction of untreated cells. Interestingly, treatment with zinc resulted in a significant increase of PKCδ in the organelle fraction.
should be noted that the total protein concentration was equivalent between the control and zinc treated cells, as shown by the total protein stain (Coomassie) of the same blot (Fig. 2A, lower image). Taken together, these results suggest that increases in the concentration of intracellular zinc influence the trafficking of PKCδ from the cytosol to organelles or the plasma membrane.

The results from our cellular fractionation experiments strongly suggested that increases in intracellular zinc affect translocation of PKCδ translocates from cytosol to Golgi complex upon exposure to zinc (A) Western blot analysis of HeLa cell lysates. Whole cell: whole cell lysate; cytosol: supernatant recovered from high-speed centrifugation of whole cell lysate; organelles: pellet recovered from high-speed centrifugation of whole cell lysate. Size markers (kDa) are shown to the left of the blot images. (B) Immunofluorescence micrographs of HeLa cells. Green fluorescence indicates localization of primary recognizing PKCδ; blue indicates presence of cell nuclei (DAPI stain). Scale bar is 10 μm. (C) Immunofluorescence micrographs of HeLa cells. Green fluorescence indicates localization of primary antibodies recognizing PKCδ; red fluorescence indicates localization of primary antibodies recognizing β-COP; blue indicates presence of cell nuclei (DAPI). Scale bar is 10 μm.
PKCδ, however it was not clear from these studies which subcellular compartment PKCδ translocates to. To gain further insight into the zinc-mediated translocation of PKCδ, we treated HeLa cells with 10 μM sodium pyrithione and 50 μM zinc chloride for 30 min and evaluated the subcellular localization of PKCδ by immunofluorescence (Fig. 2B). For these experiments, cells were grown and treated on sterile coverslips used for subsequent immunofluorescent staining (see Materials and methods). Following treatment, the cells were fixed, permeabilized and incubated with primary antibodies against PKCδ (Abcam, #182126). Following treatments with primary antibodies, the cells were washed and incubated with fluorescently-labeled secondary antibodies against the antibodies targeting PKCδ. It was observed from these series of experiments that cells treated with no zinc contained faint and amorphous distribution of PKCδ in the cytosol. Cells treated with zinc contained no observable PKCδ in cytosol and the protein appeared as a bright collection of puncta in the perinuclear region (Fig. 2B). To address whether PKCδ was being targeted to the Golgi complex, HeLa cells were plated on glass coverslips and treated as described above (see Materials and methods). The cells were then fixed, permeabilized and incubated with primary antibodies targeting the Golgi-specific protein β-COP [49] and PKCδ. Following incubation, the cells were washed, and counterstained with fluorescently-labeled antibodies specific for antibodies that targeted β-COP (red) and PKCδ (green) (Fig. 2C). It was seen from these experiments that PKCδ localizes to the Golgi complex in HeLa cells that have been treated with zinc. Collectively, the compartmentalization immunoblot and immunofluorescent data suggest that short (less than 30 min) treatments of exogenous zinc cause PKCδ to traffic from cytosol to the Golgi complex.

3.3. Intracellular zinc increase inhibits phosphorylation of PKCδ at Thr505

It should be noted that the subcellular location is not the only factor in determining the functional activity of PKCδ. Indeed, phosphorylation is another well-documented modification that controls the activation of this kinase [10]. Several phosphorylation sites on PKCδ are known to affect kinase activity, including Thr505, a residue located within the C-terminal activation loop of PKCδ. While phosphorylation at this site has been associated with increased kinase activity, it is not required for catalytic function [18]. To assess how zinc affects the phosphorylation of PKCδ at Thr505, we treated cultured HeLa cells with varying concentrations of zinc and assessed pThr505 levels by western blot (Fig. 3). HeLa cells were cultured and maintained as described in the Materials and methods. Cells were treated with 10 μM sodium pyrithione and either 0, 1, 10 or 50 μM zinc chloride for 30 min. Following treatments, the cells were lysed and intracellular proteins were separated by SDS-PAGE. Proteins were then transferred to a PVDF membrane and subjected to western blot using antibodies against phosphorylated PKCδ (pThr505) or total PKCδ (Fig. 3A). Western blots from these experiments revealed that cells treated with zinc contained significantly less phosphorylated PKCδ. Notably, inhibition of phosphorylation at Thr505 was concentration dependent, with 50 μM zinc treatments resulting in the highest level of inhibition. Percent inhibition was quantified by calculating the ratio of normalized phosphorylated PKCδ band intensity between treated and untreated (control) samples (Fig. 3B). Cells treated with 50 μM zinc showed 50.8 ± 4.3% inhibition of phosphorylation at Thr505 compared to control cells, whereas 10 and 1 μM zinc treatments showed 43.2 ± 1.6% and 5.8 ± 6.7% inhibition respectively. These data suggest that intracellular zinc increase strongly inhibits phosphorylation of PKCδ at Thr505 and that this effect is concentration dependent, with higher zinc concentration showing more pronounced inhibition.

3.4. Modeling of PKCδ three dimensional structure complexed with zinc

To gain further insight into which amino acids within the PKCδ sequence can potentially bind zinc, we employed the ZincExplorer web engine [50]. This modeling software can be used to predict potential zinc-binding sites within proteins from their primary sequence. The ZincExplorer web engine consists of a SVM-based predictor, a cluster-based predictor and a template-based predictor. This software assigns zinc-binding prediction scores (0–1) to amino acids based on charge, environment and proximity to other zinc-binding residues. Higher prediction scores are interpreted as having higher probability of binding zinc, where confidence scores of 0.49–1 are considered 66.3% sensitive and 99% specific. To initiate these studies, the primary sequence of full-length PKCδ (residues 1–676, UniProtKB - Q05655) was uploaded to the ZincExplorer web server. The resultant prediction scores were then plotted as a function of amino acid residue (Fig. 4A). Overall, the program selected 16 amino acids in the PKCδ sequence that
had high confidence for zinc binding.

To further understand the influence of zinc on the phosphorylation state of PKCδ at Thr505 we generated a three-dimensional model of PKCδ. To the best of our knowledge, no crystal structure of PKCδ currently exists; we therefore employed the protein fold recognition server PHYRE2 to develop a three-dimensional molecular model of PKCδ. The PHYRE2 program uses advanced remote homology detection methods to build 3D models of proteins with known primary sequences [51]. Our model was generated by uploading the sequence of full-length PKCδ (residues 1–676, UniProtKB - Q05655) to the PHYRE2 server. The resulting model was rendered in PyMOL and a Zn2+ pseudoatom was added near the Thr505 residue. Interestingly, this site contains proximal residues including His197, Cys200, Cys393 and Cys509, with zinc binding predictions scores of 0.91 for His197, 0.79 for Cys200, 0.46 for Cys509 and 0.23 for Cys393; with His197 and Cys200 receiving high-confidence zinc-binding scores (Fig. 4A). These amino acids represent a potentially novel His-Cys3 zinc-binding domain [52]. Following addition of the pseudoatom, the resulting apo- and holo-protein models were subjected to energy minimization using the Swiss PDB Viewer server [53]. The minimized protein structures were then rendered and visualized using PyMOL molecular modeling software (Fig. 4B and C). Distance measurements on the zinc-bound model revealed that the zinc atom sits 4.6 Å from His197, 3.7 Å from Cys200, 4.3 Å from Cys393 and 5.1 Å from Cys509 (Fig. 4C, panel on the right). It should be noted that these are...
approximate distance values generated using a computer model system and may not reflect actual interatomic distances found in the natural protein. Nevertheless, these results provide intriguing evidence of a potential PKCδ zinc-binding domain that has not been reported previously. Analysis of the solvent exposed surface of the apo-protein (no zinc) revealed that Thr505 was exposed to the solvent, which exposes this site to the cytosol, and give access to other kinases to phosphorylate that site (Fig. 4B). However, the holo-protein (zinc-bound) displayed a markedly different structure near the Thr505 residue with His197 pointed toward the zinc atom, effectively occluding Thr505 from the solvent exposed surface (Fig. 4C). These results strongly suggest that when PKCδ is bound to zinc at the His197, Cys200, Cys393 and Cys509 site, the ability of other kinases to phosphorylate Thr505 may be significantly diminished. This model shows a novel mechanism by which zinc can directly influence the phosphorylation state of a protein.

4. Discussion

Despite extensive research, many fundamental aspects of PKCδ activation remain to be elucidated. This is likely due to complex regulatory mechanisms, including phosphorylation state and subcellular localization, which control activation of this enzyme. In this study, we applied a method to treat cultured HeLa cells with exogenous zinc and showed that the phosphorylation state and subcellular localization of PKCδ is influenced by the presence of intracellular free zinc. More specifically, we demonstrated that intracellular free zinc inhibits phosphorylation of PKCδ at Thr505 in a concentration-dependent manner and that the presence of zinc facilitates targeting of PKCδ from the cytosol to the Golgi complex. Furthermore, we employed molecular modeling to elucidate how zinc binds PKCδ and inhibits phosphorylation at Thr505. These studies indicated that zinc may bind to a previously undocumented His-Cys3 binding site that is proximal to Thr505. Upon zinc binding, it was observed that the imidazole ring of His197 may orient itself in such a way that precludes phosphorylation at Thr505. These studies indicated that intracellular zinc has a two primary effects on kinase activity: 1) inhibiting phosphorylation of PKCδ at Thr505 and 2) facilitating PKCδ translocation to the Golgi complex. These new findings may help to enhance our understanding of diseases that are marked by changes in normal zinc homeostasis, including how zinc acts to modulate PKCδ phosphorylation and how cells respond to pathophysiological levels of intracellular zinc.

The detrimental effects and pathophysiological responses of intracellular zinc increase are well-documented [37–40, 54, 55]. However, many disease states induce cellular signaling events involving many independent factors that are often difficult to interpret in the context of a complex physiological response. Nevertheless, by increasing only intracellular zinc levels, we can begin to tease apart the many independent factors that are often difficult to interpret in the context of a complex physiological response. Nevertheless, by increasing only intracellular zinc levels, we can begin to tease apart the many independent factors that are often difficult to interpret in the context of a complex physiological response. Nevertheless, by increasing only intracellular zinc levels, we can begin to tease apart the many independent factors that are often difficult to interpret in the context of a complex physiological response. 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Nevertheless, by increasing only intracellular zinc levels, we can begin to tease apart the many independent factors that are often difficult to interpret in the context of a complex physiological response. Nevertheless, by increasing only intracellular zinc levels, we can begin to tease apart the many independent factors that are often difficult to interpret in the context of a complex physiological response.

Another possibility is that intracellular free zinc may directly hinder the phosphorylation of Thr505 by binding to a zinc-binding site adjacent to the Thr505 residue. To further explore the latter possibility, we employed molecular modeling software such as PHYRE2 and PyMOL to generate a folded structure of full-length PKCδ and evaluated how zinc affects cellular signaling networks that include PKCδ and will help elucidate the complex role zinc may play in regulating PKCδ activity and the cells stress response. It is likely that the subcellular localization of PKCδ influences its function because the kinase has access to different protein substrates within different areas of the cell. In this study, we evaluated how levels of intracellular zinc affect the subcellular location of PKCδ. To accomplish this, we compared cells that had normal intracellular zinc levels to cells that were treated with exogenous zinc. We used a similar method to induce intracellular zinc increase as in the Fluozin-3, AM fluorescence experiments by application of exogenous zinc in combination with a zinc ionophore. For these studies, we employed a centrifugal fractionation method to separate cytosolic proteins from organelles and plasma membrane proteins. This fractionation method used consecutive centrifugations of whole cell lysates to separate proteins by subcellular localization. These experiments revealed that exposure to zinc causes PKCδ to translocate from the cytosol to organelle and membrane compartments. On the other hand, cells that were not exposed to zinc had the majority of PKCδ localized to the cytosol, with very little found in the organelles or membranes. While this method clearly showed that intracellular zinc influences trafficking of PKCδ, it was unclear what specific organelle PKCδ translocates to following zinc exposure. To determine precisely where PKCδ translocates under these conditions, we evaluated similarly treated cells by immunofluorescence. Treated cells were fixed and labeled with antibodies specific for PKCδ and then counter-stained with fluorescently-labeled antibodies specific for those targeting PKCδ. These experiments showed that PKCδ localized to the cytosol in the absence of zinc and to a distinct perinuclear compartment in cells treated with zinc. Subsequent immunofluorescence experiments using antibodies against the Golgi-specific protein β-COP revealed that PKCδ localizes to the Golgi following treatment with zinc. These data suggest that intracellular zinc increase influences PKCδ targeting to the Golgi complex. Previous studies have shown that PKCδ translocates to the Golgi complex in response to apoptotic signals [15], indicating that increases in intracellular zinc concentrations may stimulate apoptosis in HeLa cells. However further studies are need to address whether such zinc exposure induces apoptosis in HeLa cells.

In addition to its subcellular localization, the phosphorylation state of PKCδ is also known to play a critical role in the activity and function of this kinase. In this study, we investigated the effect zinc had on the phosphorylation state of Thr505. This particular site is notable because it has been suggested to influence substrate specificity [19]. Furthermore, this site distinguishes PKCδ from other members of the ‘novel kinase’ family in that PKCδ is the only member that does not require this site to be phosphorylated for activation [56]. To evaluate the influence that intracellular zinc increase has on the phosphorylation state of Thr505, we treated HeLa cells as described above and quantified levels of phosphorylated PKCδ (pThr505) by western blot. Our results indicate that when intracellular zinc increases, the phosphorylation of PKCδ is inhibited. This result was concentration-dependent, with higher zinc treatment concentrations translating into greater inhibition of phosphorylation. Other groups have proposed that phosphorylation at Thr505 can influence substrate specificity [19]. Therefore, we suggest that zinc plays a role in regulating the activity and specificity of PKCδ, and may ultimately change cell fate.

We envision several independent, or perhaps concerted, scenarios where the phosphorylation status of Thr505 could be influenced by intracellular concentrations of free zinc. For example, increases in intracellular zinc may inhibit kinases that phosphorylate Thr505 or stimulate the activation of phosphatases that dephosphorylate Thr505. Another possibility is that intracellular free zinc may directly hinder the phosphorylation of Thr505 by binding to a zinc-binding site adjacent to the Thr505 residue. To further explore the latter possibility, we employed molecular modeling software such as PHYRE2 and PyMOL to generate a folded structure of full-length PKCδ and evaluated how
possible zinc binding could influence phosphorylation at Thr105. Our modeling efforts were initiated by assigning zinc binding prediction scores to each residue of the full-length PKCδ protein using the zinc binding prediction engine ZincExplorer. A total of 16 amino acids were determined to have zinc binding potential, interpreted as having zinc binding prediction scores greater than or near 0.49. We then generated a folded model of PKCδ by uploading its primary sequence (residues 1–676, UniProtKB - Q05655) to the PHYRE2 server. Interestingly, this folded model was observed to contain a potential His-Cys3 zinc binding site near Thr105, consisting of His197, Cys200, Cys393 and Cys509. Three of the residues His197, Cys200 and Cys509 each had zinc binding prediction scores greater than 0.49 indicating a high probability for binding zinc. When a zinc pseudoatom was placed near this binding pocket and the structure was energy minimized, it was observed that the donor residues were repositioned at distances between 4.6 and 5.4 Å from the zinc ion. Notably, when the solvent exposed surface was analyzed, it appeared that the presence of the zinc ion causes the His197 residue to essentially close the zinc binding pocket and occlude Thr105 from the solvent. The zinc-mediated reorientation of His197 provides a rational mechanism for how zinc binding inhibits phosphorylation of Thr105 in PKCδ. We therefore hypothesize that when intracellular zinc concentration increases, PKCδ will bind zinc at His197, Cys200, Cys393 and Cys509, and this interaction will directly prevent further phosphorylation of Thr105. This modification may also influence the intracellular targeting and function of PKCδ. Indeed, it has been shown previously that T505A mutations cause PKCδ to translocate to cellular compartments similar to the Golgi complex [57].

In summary, we have studied the effect intracellular zinc increase has on the phosphorylation state and subcellular localization of PKCδ. Our results indicate that short exposure (less than 30 min) of HeLa cells to exogenous zinc inhibits the phosphorylation of PKCδ at Thr105. We also showed that zinc influences the subcellular targeting of PKCδ, where increases in intracellular zinc facilitates translocation of PKCδ to the Golgi complex. It should be noted that these data alone are not wholly sufficient to elucidate whether intracellular zinc directly causes dephosphorylation and translocation of PKCδ to the Golgi complex. However, we propose here a novel mechanism by which PKCδ can bind zinc ions that would preclude phosphorylation of Thr105. Molecular modeling simulations have provided a rational for how PKCδ can use a novel His-Cys3 zinc-binding pocket to ‘sense’ intracellular concentrations of free zinc and respond to changes in zinc homeostasis. These findings indicate that zinc, and possibly other transition metals, may play a distinct role in regulating the activity of ‘novel kinases’ and should provide a basis for future studies investigating cellular processes that rely on metal ion-protein interactions.

5. Conclusions

We have identified a novel effector of PKCδ phosphorylation to be free zinc. This work demonstrated the intracellular zinc increase causes concentration dependent inhibition of phosphorylation of PKCδ at activation loop site Thr105 and changes the subcellular localization of PKCδ from cytosolic to Golgi complex.

Author contributions

KGS conceived the study, performed experiments and researched data, wrote the manuscript, generated the figures and references. JMH researched and performed the computer modeling, generated the computer modeling figures, contributed to discussions, reviewed and edited the manuscript and added references. YVL coordinated the study, secured the funding, contributed to discussions and reviewed the manuscript. All authors approved the final version of the manuscript.

Conflict of interest

The authors declare that they have no conflict of interests.

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