Protein Kinase D2 Modulates Cell Cycle By Stabilizing Aurora A Kinase at Centrosomes

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Abstract

Aurora A kinase (AURKA) is a master cell-cycle regulator that is often dysregulated in human cancers. Its overexpression has been associated with genome instability and oncogenic transformation. The protein kinase D (PKD) family is an emerging therapeutic target of cancer. Aberrant PKD activation has been implicated in tumor growth and survival, yet the underlying mechanisms remain to be elucidated. This study identified, for the first time, a functional crosstalk between PKD2 and Aurora A kinase in cancer cells. The data demonstrate that PKD2 is catalytically active during the G2–M phases of the cell cycle, and inactivation or depletion of PKD2 causes delay in mitotic entry due to downregulation of Aurora A, an effect that can be rescued by overexpression of Aurora A. Moreover, PKD2 localizes in the centrosome with Aurora A by binding to γ-tubulin. Knockdown of PKD2 caused defects in centrosome separation, elongated G2 phase, mitotic catastrophe, and eventually cell death via apoptosis. Mechanistically, PKD2 interferes with Fbxw7 function to protect Aurora A from ubiquitin- and proteasome-dependent degradation. Taken together, these results identify PKD as a cell-cycle checkpoint kinase that positively modulates G2–M transition through Aurora A kinase in mammalian cells.

Implications: PKD2 is a novel cell-cycle regulator that promotes G2–M transition by modulating Aurora A kinase stability in cancer cells and suggests the PKD2/Aurora A kinase regulatory axis as a new therapeutic target for cancer treatment.

Introduction

Cell cycle is propelled by well-coordinated complex signaling events by which cells grow and divide. Uncontrolled cell division, whereby cells go through the cycle unchecked, lies at the base of cancer formation and progression. Cell cycle occurs through progression of four distinct stages, namely G0–G1, S, G2, and M, which are monitored by a battery of cyclin-dependent kinases (CDK) and their partner cyclins (1). Mutations in the signaling pathways, aberrant activation of CDKs, genetic lesions in the genes encoding cell-cycle–regulated proteins result in genome instability, abnormal growth, and eventually cancer.

The protein kinase D (PKD) family of serine/threonine kinases belongs to the Ca⁺⁺/Calmodulin-dependent protein kinase (CaMK) superfamily and it consists of three isoforms in mammals, notably, PKD1, PKD2, and PKD3, which are highly conserved throughout evolution (2). Structurally, PKD possesses an N-terminal regulatory domain that contains a tandem cysteine-rich Zn-finger like motif (CRD, C1a, and C1b) and a plekstrin homology (PH) domain, and a C-terminal catalytic domain (3, 4). In a canonical pathway, diverse physiologic factors, such as GPCR agonists, bioactive peptides (5), lipids (6), and growth factors (7) converge to the activation of PKDs through the generation of diacylglycerol (DAG) by phospholipase C (PLC) and the activation of classical or novel protein kinase C (c/nPKC; ref. 8). Activation of PKD is marked by phosphorylation of two conserved serine residues in the activation loop of PKDs and concomitant autophosphorylation of PKD1Ser³⁶⁰⁵ or PKD2Ser³⁶⁰⁶ are widely used as a biomarker of PKD activation (3, 9). PKD regulates a wide range of cellular processes including cell proliferation, migration/invasion, angiogenesis, protein trafficking, and gene expression. Its functional importance has been implicated in major human cancers including carcinomas of breast, skin, pancreas, and prostate (8).

Aurora kinases are master regulators of the cell cycle (10–13). Human genome encodes three isoforms of Aurora kinases; namely, Aurora kinases A, B, and C (14). Aurora A localizes on centrosomes, spindle poles, and spindles during mitosis and regulates centrosome function, spindle assembly, and mitotic progression. Expression of Aurora A is cell-cycle–regulated, that is, the levels of mRNA and protein are low in G1 and S, increase during G2–M, and reduce during mitotic exit. Aurora A is ubiquitinated by APC/Cdh1 and Fbxw7 ubiquitin ligases and degraded in proteasome-dependent pathways (15, 16). A number of Aurora A substrates have been reported, such as TPX2, CDC25B, and p53 (17). Aurora A is frequently overexpressed in cancer. Aberrant expression of Aurora A promotes centrosome amplification and aneuploidy, leading to genome instability and consequently oncogenic transformation (18). Thus, Aurora A represents a well-established therapeutic target for cancer. In this study, we uncovered a novel crosstalk between PKD2 and Aurora A kinase. Our study defined the functional impact of this regulatory axis on G2–M transition and mitosis. We have shown that PKD2 is required for stabilization of Aurora A kinase, and that the abrogation of PKD2 expression or inhibition of its catalytic activity causes proteasome-mediated downregulation of Aurora A kinase displaying delayed G2, mitotic catastrophe and cell death. Moreover, we report that...
PKD2 localizes on the centrosomes during G2 and that it is critical for centrosome separation in early G2. Taken together, our study provides compelling evidence to support the role of PKD2 in G2–M transition through modulating Aurora A stability, and aberrant PKD2 activity may contribute to oncogenesis.

**Materials and Methods**

**Cell culture, synchronization, and treatments**

Authenticated HeLa, LNCap, and PC3 cells were obtained from ATCC. After purchase, the cell lines were expanded and frozen according to manufacturer's instructions after two to three passages. LNCap was used for no more than 8 passages and PC3/HeLa according to manufacturer's instructions after two to three passages. LNCap and PC3 cells were grown in 1 × Minimum Essential Medium (MEM) supplemented with Eagle salt and l-glutamine (Invitrogen), RPMI1640 (Invitrogen), or Ham F-12 (Thermo Fisher Scientific) media, respectively, supplemented with 10% FBS and 1 × penicillin/streptomycin (Thermo Fisher Scientific). MT30002CI and maintained at 37°C in a humidified incubator containing 5% CO2. Cells were synchronized at G1–S or the start of M phases by double thymidine or nocodazole, respectively. Briefly, for G1–S synchronization, cells were treated with 2 mmol/L thymidine for 16 hours, washed three times with 1 × PBS, and re-fed into fresh culture medium without thymidine for 8 hours. The second block was performed by treating the cells with 2 mmol/L thymidine and nocodazole for another 16 hours. Cell arrest at the start of M phase was performed by treating the cells with nocodazole (100 ng/mL) for 16 hours. PKD inhibitors (CRT0066101 and k-B-NB142-70) were used at a concentration of 2 μmol/L for indicated times. Turbofect (Thermo Fisher Scientific) and Lipofectamine 3000 (Invitrogen) transfection reagents were used to transfect plasmids and siRNAs, respectively, according to the manufacturer's protocol.

**Cell survival assay**

Cell survival assay was performed by Cell Counting Kit-8 (Dojindo Laboratories) according to the manufacturer's protocol. Briefly, HeLa and PC3 cells were plated at a density of 4,000 cells per well in 96-well plate. After treating the cells with different doses of drugs (CRT0066101 and k-B-NB142-70) for 72 hours, CCK-8 solution was added to cells in each well, followed by incubation for 2 hours. Cell proliferation/viability was determined by measuring the OD at 450 nm.

**Western blotting and densitometry analysis**

Cells were lysed in cell lysis buffer (50 mmol/L Tris-HCl pH 7.5, 150 mmol/L NaCl, 1.5 mmol/L MgCl2, 10% glycerol, 1% Triton X-100, 5 mmol/L EGTA, 1 mmol/L Na3VO4, 10 mmol/L NaF, 1 mmol/L β-glycerophosphate and protease inhibitor cocktail). Cell extracts were resolved by SDS-PAGE, transferred to nitrocellulose membrane, and probed with following antibodies: PKD2 (Thermo Fisher Scientific, PAS-64538), Cyclin B1 (Cell Signaling Technology, 12231), phospho CDC25The48 (Cell Signaling Technology, 12028), phospho-Histone H3Ser10 (Cell Signaling Technology, 3377), Aurora Kinase A (Cell Signaling Technology, 14475), Aurora Kinase B (Cell Signaling Technology, 3094), γ-tubulin (Santa Cruz Biotechnology, sc-17788), PARP (Cell Signaling Technology, 9542), ubiquitin (P4D1, Cell Signaling Technology 3936), Lamin A (Cell Signaling Technology, 2032), Cdh1 (Santa Cruz Biotechnology, sc-56312), Fbxw7 (Abcam, EPR8069), and GAPDH (Enzo ADI-CSA-335-E). Protein half-life was measured as described previously (19). Briefly, the band intensities were quantified by ImageJ software and the values of target proteins were normalized against the band intensity of GAPDH. The measured protein intensity values were log transformed, fitted to a linear regression plot, and decay rate constant (k) was determined. The half-life was calculated from the formula $T_{1/2} = \ln(2)/k$.

**Immunoprecipitation analysis**

For immunoprecipitation assay, whole-cell extract was prepared by lysing the cells with cell lysis buffer. The lysate was centrifuged, supernatant was collected, and protein concentration was estimated using BCA protein assay kit (Pierce) according to manufacturer's protocol. Cell lysate containing equal amount of total protein was incubated with primary antibodies at 4°C overnight. Protein A/G agarose beads were added later for another 2 hours. The immunoprecipitates were collected and washed with wash buffer containing 0.5% Triton X-100. Finally, the samples were diluted with 6 × SDS PAGE loading buffer, boiled for 10 minutes, and subjected to Western blotting.

**Subcellular fractionation**

Cells were washed with ice-cold PBS, collected by centrifugation at 1,000 rpm for 5 minutes, and resuspended in cytoplasm extraction buffer (10 mmol/L HEPES pH 7.5, 60 mmol/L KCl, 1 mmol/L EDTA, 0.1% NP-40, 1 mmol/L DTT, and protease inhibitor cocktail) for 5 minutes on ice. The cell lysate was centrifuged at 1,000 rpm for 5 minutes at 4°C and cytoplasmic fraction was separated in a new tube. The nuclei were washed in cytoplasm extraction buffer without NP-40 and resuspended in nucleus extraction buffer (20 mmol/L Tris-HCl pH 8, 400 mmol/L NaCl, 1.5 mmol/L MgCl2, 0.2 mmol/L EDTA, 25% glycerol and protease inhibitor cocktail). The extracts were incubated on ice for 10 minutes. Both cytoplasmic and nuclear extracts were centrifuged at 14,000 rpm for 15 minutes at 4°C and transferred into new tubes.

**Flow cytometry**

Trypsinized cells were pelleted, washed twice with ice-cold FACs buffer (0.1% glucose in PBS), and resuspended in ice cold 70% ethanol while vortexing. Cells were fixed by incubating overnight at 4°C, washed with PBS, and incubated with 1 mL propidium iodide (PI) staining solution (50 μg/mL PI, 0.1 mg/mL RNase A in FACs buffer) for 30 minutes. Cells were analyzed in a FACS C40 (BD Biosciences) flow cytometer and the data were analyzed by ModFit 3.0 software (BD Biosciences). A total of 10,000 events were analyzed for each sample.

**Apoptosis assay**

Trypsinized cells were washed twice with ice-cold PBS and stained with Annexin V-Alexa Fluor 488 antibody and propidium iodide (Molecular Probes, Thermo Fisher Scientific) according to the manufacturer's instructions. The samples were divided in two parts. One half was subjected to FACScalibur (BD Biosciences) flow cytometer and the other part was subjected to confocal fluorescence microscopy using Alexa Fluor 488 and PI filters.
Immunofluorescence staining and microscopy

HeLa and PC3 cells were grown on poly-D-lysine–coated coverslips, washed with PBS, fixed in 4% paraformaldehyde at room temperature for 30 minutes, washed three times with 1× PBS, and blocked in 5% normal goat serum containing 0.3% Triton X-100 for 1 hour at room temperature. The cells were incubated with the primary antibodies diluted in antibody dilution buffer (1% BSA, 0.3% Triton X-100 in 1× PBS) overnight at 4°C, washed three times in 1× PBS, and incubated with secondary antibodies diluted in antibody dilution buffer for 1 hour at room temperature. The cells were counterstained with DAPI (1 μg/mL) and the coverslips were mounted in ProLong Gold antifade reagent (Invitrogen). The cells were analyzed using an Olympus Fluoview (FV1000) confocal microscope using 60 X/1.45 objectives.

Data availability

The data that support the findings of this study are available from the corresponding author upon reasonable request.

Statistical analysis

Data analysis was done using the Student t test for comparison between two groups (two-tailed). All statistical analyses were done using GraphPad Prism IV software (GraphPad Software). All values are represented as mean ± SEM of at least three independent experiments. A P < 0.05 was considered statistically significant (*, P < 0.05; **, P < 0.01; ***, P < 0.001; ****, P < 0.0001; ns, not significant).

Results

Elevated PKD2 activity during mitotic entry of the cell cycle

First, we examined the expression and activity of PKD (refers to PKD2 and PKD3 throughout the text) in different cell-cycle stages. PKD2 and PKD3 are major isoforms expressed in HeLa cells (Supplementary Fig. S1), similar to the prostate cancer cell line PC3, but differed from LNCaP cells (20). HeLa and PC3 cells were treated with nocodazole, a microtubule-destabilizing agent that activates the spindle assembly checkpoint and causes cell arrest in start of M-phase (prometaphase), and the activity of PKD2 was assessed after releasing from nocodazole using the phospho-PKD2Ser876 (p-PKD2) antibody. As shown in Fig. 1A (top) and B, p-PKD2 was detected in G2–M–arrested HeLa cells following nocodazole synchronization and persisted till the end of mitosis. This was accompanied by increased cyclin B1, a marker for G2 and M phases of the cell cycle, and phospho-Histone H3Ser10 (pH3), which was detectable in G2, peaks during mitosis and disappears when cells enters in G1 (Fig. 1A; Supplementary Fig. S2). Similar results were obtained with PC3 cells where activation of PKD2 was marked by increased levels of p-PKD2 after release from nocodazole arrest (Fig. 1A, bottom). To determine whether the activation is specific to G2–M, PKD2 activation was measured in G2–S synchronized HeLa cells that progressed to M-phase. As shown in Fig. 1C and D, when cells were released from a double thymidine (DT) block that synchronizes cells at G1–S border, PKD2 phosphorylation gradually elevated, starting at 4 hours when cells entered in late S-phase peaked around 10 hours in late G2, and persisted throughout mitosis till 14 hours after thymidine release. The highest activity was detected at G2–M border and M-phase. This pattern correlated well with increased cyclin B1 and pH3. As controls, PKD2 remained the same under both conditions.

The intracellular distribution of PKD2 was then analyzed at different stages of the cell cycle by immunofluorescence (IF) staining. In HeLa cells, PKD2 was primarily localized in the cytoplasm and perinuclear zone in interphase. When cells progress through cell cycle, it exhibited scattered punctate staining that are minimally overlapped with mitotic bodies (spindle fibers and midbody) and excluded from condensed chromatin (Fig. 1E, top). PKD2 phosphorylation was low or undetectable during interphase and significantly increased during G2, sustained throughout prophase, metaphase, and anaphase, before returning to baseline when the cells entered telophase. To corroborate these results in cancer, PC3 cells were stained in similar fashion. Similar results were found when different stages of the cell cycle in PC3 cells were imaged using antibodies against phospho-PKD2Ser876 (green) and α-tubulin (red, Fig. 1F). It is well documented that PKD, upon activation, accumulates in the nucleus (21). Indeed, we observed elevated active PKD in the nucleus during G2 and early prophase when the nucleus was still intact, further attesting that PKD2 was activated during G2 and mitosis. We further validated that phospho-PKD2 was increased in the nucleus during early mitosis by cell fractionation (Fig. 1G, left). HeLa and PC3 cells were arrested at the start of M phase by nocodazole and cytoplasmic (Cyto) or nuclear (Nuc) fractions were prepared. The samples were analyzed by Western blotting using phospho-PKD2 antibody and the results showed that, indeed, active PKD2 levels were elevated in early mitosis (Fig. 1G, right). Lamin A was blotted as a marker for the nuclear fraction. Taken together, these results suggested that PKD is active during G2 and mitosis and may play an important role in cell-cycle regulation.

Inactivation or knockdown of PKD2 causes delay in G2–M transition

The activation profile of PKD2 suggests that PKD2 may regulate G2 and M-phase cell-cycle progression. To test this hypothesis, cells with altered PKD2 expression and activity were synchronized by DT at G2–S border and released (Figs. 2A and 3A); cell-cycle progression was analyzed by flow cytometry. As shown in Fig. 2B and C, after thymidine release, cells reached G2–M at 8 hours, and by 12 hours, cells resumed normal distribution. In contrast, cells treated with the PKD inhibitor CRT0066101 (CRT101) reached G2–M at 8 hours, stayed after 12 hours, and by 14 hours there remained a large G2–M population as compared with the control. Inactivation of PKD by another PKD inhibitor kb-NB142–70 (kb-NB) gave rise to similar results (Fig. 2B, kb-NB). Quantification analysis confirmed prolonged G2–M phase of cell cycle in cells treated with PKC inhibitors (Fig. 2C). Thus, inactivation PKD2 prolonged G2–M phase and resulted in delayed mitotic entry in cells.

To determine that PKD2 activity was indeed required for G2–M transition, cells transfected with PKD2 siRNAs were synchronized to G2–S border by DT and released; cell-cycle progression was monitored by flow cytometry (Fig. 3A). Our data showed that cells transfected with nontargeting siRNA reached G2–M around 8 hours and returned to normal distribution at about 12–14 hours. In contrast, cells transfected with two PKD2 siRNAs (Fig. 3B) progressed slower through the cell cycle and peaked at G2–M phase around 12 hours and remained till after 14 hours (Fig. 3C), indicating that PKD2-depleted cells progressed through G2–M boundary at much slower rate than control cells. Knockdown of PKD2 using two siRNAs in these samples were confirmed by Western blotting (Fig. 3D). To corroborate these findings, we...
PKD is activated during mitotic entry. A, HeLa and PC3 cells were synchronized with nocodazole (100 ng/mL) for 16 hours and mitotic cells were released into fresh medium for indicated times. Cell extracts from each sample were subjected to Western blot analysis to analyze activation of PKD2. B, Densitometric quantification of phospho-PKD2S876 from the immunoblot of HeLa was performed. C, Western blot analysis of indicated proteins after the HeLa cells were released from a double thymidine (DT) block was performed. D, Densitometric quantification of phospho-PKD2S876 from the immunoblot of C was performed. E, HeLa cells were subjected to immunofluorescence (IF) staining to visualize colocalization PKD2 or phospho-PKD2S876 with α-tubulin during different phases of cell cycle. F, PC3 cells were subjected to IF staining to visualize colocalization of phospho-PKD2S876 with α-tubulin during different phases of cell cycle. DAPI was used to stain the nucleus. Scale bar, 8 μm. GAPDH was used as loading control in A and C. G, HeLa and PC3 cells were synchronized with nocodazole (100 ng/mL) for 16 hours and mitotic cells were collected by shake-off. Cytoplasmic (Cyto) and nuclear (Nuc) fractions were prepared and the samples were analyzed by Western blotting with indicated antibodies (left). Relative amount of phospho-PKD2S876 was quantified (right). The graphs show average of three independent experiments with error bars representing SEM (*, P < 0.05; **, P < 0.001; ***, P < 0.0001; ns, not significant).
examined HeLa cells with overexpressed PKD2. As shown
in Fig. 3E, overexpression of PKD2 significantly increased
the number of cells entering mitosis. Taken together, our
data indicate that PKD2 was activated in G2–M and its
activity was required for mitotic entry.

Abrogation of PKD2 expression sensitizes cells to mitotic
catastrophe and cell death
Cells experience prolonged G2 arrest and delayed mitotic
entry may eventually enter into mitosis, however, they often suffer from
aberrant mitosis known as mitotic catastrophe, and die subse-
quently by apoptosis (22). Here, we examined whether mitotic
catastrophe was associated with loss/inactivation of PKD2 and
delayed G2–M transition. Knockdown or inactivation of PKD2
using two siRNAs or PKD inhibitor CRT101, respectively, induced
a series of mitotic defects including chromosomal misalignment
at metaphase plane, abnormal chromosomal arrangement
(Fig. 4A, white arrows), and formation of multiple centrosomes
(Fig. 4A, red arrows). Quantification analysis indicated that
depletion or inactivation of PKD2 significantly increased cell

Figure 2.
Inactivation of PKD by CRT101 and kb-NB induces a delay in G2–M progression. A, Schematic representation of experimental design is shown. B, Flow
cytometry analysis of cells treated with DMSO, CRT101, and kb-NB were performed according to A. C, Summarizing graph of the distribution of different phases
of cell cycle are shown (right). The graphs show average of three independent experiments with error bars representing SEM.

PKD2 Promotes G2–M Transition via Aurora A
population with defective mitosis (Fig. 4B). It is well known that mitotic catastrophe leads to apoptosis followed by PARP and Lamin A cleavage (23, 24). Analysis of PKD2-depleted or PKD-inactivated HeLa cells showed expression of cleaved PARP (Fig. 4C). To further this analysis, cells transfected with siNT and siPKD2s were stained with Annexin V-Alexa Fluor 488/propidium iodide (PI) and analyzed by flow cytometry (Fig. 4D, left). Knockdown of PKD2 caused significant cell death (PI-positive) with elevated apoptotic population (Annexin V-positive) as compared with the control samples (siNT; Fig. 4D, right). Treatment of both HeLa and PC3 cells using increasing concentrations of CRT101 and kb-NB reduced cell viability in a concentration-dependent manner (Fig. 4E). Taken together, our data indicated that G2-M deficiencies caused by PKD2 depletion or inhibition could result in mitotic catastrophe and cell death followed by apoptosis.

Inactivation of PKD2 causes degradation of Aurora A, an effect that can be suppressed by overexpression of Aurora A.

We showed that knockdown or inactivation of PKD2 caused prolonged G2-M transition, delay in mitotic entry, and mitotic catastrophe (Figs. 2–4). In the order to gain insights into the role of PKD in cell cycle, we sought to identify its downstream targets. Through a whole-genome RNA-seq analysis conducted on cancer cells treated with CRT101 and kb-NB, we identified AURKA/Aurora A kinase as one of the top candidate genes altered by inhibition of PKD. Aurora A is a master regulator of G2-M and mitosis (12, 25–27). First, using normal prostate epithelial cell line RWPE-1 and prostate adenocarcinoma (PAC) cell lines, we sought to identify whether expression level of PKD2 correlates with that of Aurora A kinase. As shown in Fig. 5A, we observed positive correlation between PKD2 and Aurora A in these cell lines. Next, we subjected HeLa cells to IF staining and checked whether PKD2 and Aurora A colocalize at different stages of cell cycle.

**Figure 3.**
Abrogation of PKD2 expression delays G2-M progression. A, Schematic representation of experimental design is shown. B, Flow cytometry analysis of cells transfected with control nontargeting siRNA (siNT) and two siRNAs against PKD2 was performed according to A. C, Summarizing graph of the distribution of different phases of cell cycle are shown. The graphs show average of three independent experiments with error bars representing SEM. D, Western blot analysis of the samples from A was performed to confirm knockdown of PKD2. GAPDH was used as loading control. E, PKD2 was ectopically overexpressed using a plasmid expressing Flag-PKD2 (top) and percentage of interphase and mitotic cell population was counted (bottom). The graphs show average of three independent experiments with error bars representing SEM (*, P < 0.5; **, P < 0.01; ***, P < 0.0001).
mitosis. Our confocal microscopy results showed that a significant portion of PKD2 (green) colocalizes with Aurora A (red) during mitosis (Fig. 5B), raising a strong possibility that PKD2 might contribute to cell cycle by regulating Aurora A. We examined the expression levels of Aurora A upon inactivation or depletion of PKD. HeLa cells were arrested at the start of M phase by nocodazole, released from the block by mitotic shake off and turnover of Aurora A protein levels was analyzed in the absence or presence of CRT101 by Western blotting. The results showed that CRT101 caused rapid downregulation of Aurora A with < 50% remaining 2 hours after CRT101 treatment (Fig. 5C). To corroborate this finding in the context of cancer, we treated PC3 prostate adenocarcinoma and COV362 ovarian epithelial-endometroid carcinoma cell lines as described above and checked Aurora A turnover by Western blotting (Fig. 5D). As expected, Aurora A was rapidly downregulated by CRT101 in these two cell lines.

Figure 4.
Knockdown of PKD2 causes mitotic catastrophe followed by cell death. A, IF staining of cells transfected with control nontargeting siRNA (siNT) and two siRNAs against PKD2 (top) or DMSO/CRT101 (bottom) was performed to visualize γ-tubulin (green). DAPI was used to stain the nucleus. Red and white arrow heads indicate defects in spindle pole formation and misaligned chromosomes, respectively. B, Quantification of cell population harboring mitotic defects after the cells were treated with either si-PKD2 (top) or CRT101 (bottom). C, Western blot analysis was done to show cleavage of PARP/Lamin A upon depletion or inactivation of PKD2 in HeLa. GAPDH served as loading control. D, Microscopic analysis of cells transfected with siNT or two siPKD2s after costaining with Annexin V-Alexa Fluor 488 and propidium iodide (left). Quantification of cell population positive for Annexin V-Alexa Fluor 488 and PI (right). E, HeLa and PC3 cells were treated with different concentrations of PKD inhibitors (CRT101, kb-NB) and cultured for 72 hours. MTT assay was performed to assess cell viability. The graphs show average of three independent experiments with error bars representing SEM (***, P < 0.001; ****, P < 0.0001).
Meanwhile, the downregulation of Aurora A coupled with time-dependent decline of cyclin B1, p-CDC25S/T48 and p-HH3, which are in line with a critical role of PKD activity in G2→M transition and mitotic progression (Fig. 5E, left). In comparison, the expression levels of Aurora B did not significantly change (Fig. 5D and E, Aurora B), indicating a selective effect of CRT101 on Aurora A degradation. Importantly, when ectopically overexpressed, Aurora A completely reversed the effect of CTR101 on cell-cycle markers (Fig. 5E, right; Supplementary Fig. S3). Thus, Aurora A mediated the effect of PKD inhibition on G2→M transition. To corroborate these findings, Aurora A protein stability was assessed in cells transfected with siPKD2-3 (siD2) and treated with cycloheximide (CHX) that blocks protein synthesis. In the control cells (siNT), the half-life (t1/2) of Aurora A was 49 hours. Partial knockdown of PKD2 reduced t1/2 of Aurora A to about 28 hours, while nearly complete knockdown further reduced it to 11 hours (Fig. 5F). Similarly, knockdown of PKD3 with two siRNAs also resulted in downregulation of Aurora A (Supplementary Fig. S5). These results identify Aurora A as a key mediator of PKD2-regulated cell-cycle progression, and the activation of PKD2 at G2→M stabilizes Aurora A kinase and promotes mitotic entry.

Figure 5.
Inactivation or knockdown of PKD2 causes downregulation of Aurora A kinase. A, Western blot analysis was performed to determine expression patterns of PKD2 and Aurora A in designated cell lines. B, HeLa cells were stained with antibodies against PKD2 (green) and Aurora A (red). IF staining was performed to visualize colocalization of PKD2 and Aurora A at different stages of cell cycle. Scale bar, 8 µm. DAPI was used to stain the nucleus. C, HeLa cells were synchronized at the start of M-phase by nocodazole, released from the block and mitotic cells were cultured for indicated times with or without CRT101. Downregulation of Aurora A was analyzed by Western blotting (top). Quantification of Aurora A protein levels by densitometry scanning from immunoblots was performed (bottom). D, PC3 and COV362 cells were treated as described in C and turnover of Aurora A protein levels were analyzed by Western blotting. The expression levels of Aurora B were analyzed as control. E, HeLa Cells were arrested at the start of M-phase by nocodazole (noc), released in fresh medium containing DMSO or CRT101 for indicated times and cell extracts from each samples were analyzed by Western blotting using indicated antibodies (left). Aurora A was ectopically overexpressed and the cells were released from nocodazole block in fresh medium containing CRT101 for indicated times. Cell extracts from each sample were analyzed by Western blotting using indicated antibodies. Overexpression of Aurora A was confirmed by Western blotting. F, Cells were transfected by si-NT (control) and two different concentrations of si-PKD2. Forty-eight hours post transfection, cells were treated with cycloheximide (CHX) that blocks protein synthesis. In the control cells (siNT), the half-life (t1/2) of Aurora A was 49 hours. Partial knockdown of PKD2 reduced t1/2 of Aurora A to about 28 hours, while nearly complete knockdown further reduced it to 11 hours (Fig. 5F). Similarly, knockdown of PKD3 with two siRNAs also resulted in downregulation of Aurora A (Supplementary Fig. S5). These results identify Aurora A as a key mediator of PKD2-regulated cell-cycle progression, and the activation of PKD2 at G2→M stabilizes Aurora A kinase and promotes mitotic entry.
PKD2 and Aurora A localize at the centrosome and knockdown of PKD2 inhibits centrosome separation during G2.

Aurora A has been shown to localize at mitotic apparatus and contribute to G2–M transition and mitotic progression (13, 22, 28, 29). Aurora A resides in the centrosome and regulates centrosome duplication and separation, inhibition of Aurora A arrests cells in G2 and blocks mitotic entry. Here, we sought to determine whether the regulation of Aurora A by PKD also took place at the centrosome and whether it affects centrosome function. HeLa cells were costained with PKD2 and γ-tubulin, a marker of the centrosome. As shown in Fig. 6A, PKD2 (red) was found to colocalize with γ-tubulin (green) at different stages of cell cycle, and the colocalization was most prominent in the G2 stage. Next, cells were synchronized to the start of M-phase by nocodazole and the interaction of PKD2 and γ-tubulin were examined by reciprocal immunoprecipitation. As shown in Fig. 6B, γ-tubulin was detected in immunoprecipitated PKD2 complex, and vice versa. Thus, a portion of PKD2 was present on the centrosome and PKD2 directly interacts with γ-tubulin at the centrosome during G2–M, which was further confirmed by IF staining (Fig. 6C). Next, we examined whether PKD2 might directly interact with Aurora A. Aurora A is known to localize at the centrosome (13, 27, 30, 31). Our data confirmed that Aurora A (red) colocalized with γ-tubulin (green) during G2 and mitosis (Supplementary Fig. S6). By costaining for PKD2 and Aurora A, we also found significant colocalization of PKD2 (green) and Aurora A (red) during G2 and mitosis (Fig. 5A). Colocalization of PKD2 and Aurora A on the centrosome raised a possibility that they might physically interact. However, our follow-up study indicated that PKD2 failed to coimmunoprecipitate with Aurora A in G2–M–synchronized cells (Supplementary Fig. S7). Despite the lack of direct interaction, the colocalization of PKD2 and Aurora A support the potential functional link between the two proteins in centrosome. Similar colocalization pattern was observed when the intracellular distributions of PKD3 and Aurora A were analyzed during G2 and mitosis (Supplementary Fig. S7). Aurora A is crucial for

![Image](https://example.com/image.png)

Figure 6. PKD2 positively regulates centriole separation during late G2 by colocalizing with centrosomes. A–C, HeLa cell was subjected to IF staining to visualize colocalization of PKD2 (red) and γ-tubulin (green). B, Coimmunoprecipitation assay was carried out to assess physical interaction between PKD2 and γ-tubulin. D, Knockdown of PKD2 by two siRNAs was confirmed by IF staining. E, Centrioles were visualized by IF staining of γ-tubulin in cells transfected with siNT or two siPKD2s (left). Cell population harboring unseparated centrioles was quantified (right). At least 100 cells were counted and graph shows average number of cells with error bars representing SEM (***, P < 0.0001). DAPI was used to stain the nucleus.
PKD2 interferes with Fbxw7-mediated ubiquitination and downregulation of Aurora A

To assess whether PKD regulates Aurora A degradation through the proteasome degradation pathway, cells were synchronized with nocodazole, released in fresh medium containing CRT101 in the presence or absence of a proteasome inhibitor MG132, and the levels of Aurora A were examined by immunoblotting. Our data showed that Aurora A levels were dampened within 3 hours of CRT101 treatment and MG132 completely inhibited CRT101-induced Aurora A degradation (Fig. 7A). The half-life of Aurora A was measured in the presence of cycloheximide (CHX). As shown in Fig. 7B, treatment of CRT101 reduced the half-life of Aurora A protein to about 5 hours, whereas MG132 restored its half-life like control sample (nocodazole only; Fig. 7B). These results demonstrated that CRT101 directed Aurora A to the proteasomal pathway for its degradation. To further corroborate this finding, G2–M–arrested cells were released in medium containing MG132 with or without CRT101, and cell extracts from the samples were subjected to IP using anti-Aurora A antibody, followed by immunoblotting with anti-ubiquitin antibody (Fig. 7C). We found that CRT101 increased ubiquitination of Aurora A (Fig. 7C, top) confirming that Aurora A is downregulated via ubiquitination-dependent proteasomal degradation pathway upon treatment with CRT101. Aurora A is targeted by Cdh1 and F-box protein Fbxw7 (15, 32) for degradation. Therefore, to gain insight into PKD-mediated downregulation of Aurora A, we knocked down PKD2, Cdh1, and Fbxw7 using site-specific siRNAs and analyzed the turnover of Aurora A protein levels. Knockdown of PKD2 downregulated Aurora A, whereas depletion of Cdh1 did not significantly alter Aurora A protein level (Fig. 7D). Interestingly, when Fbxw7 was depleted in PKD2 knocked down cells, Aurora A level was significantly restored as compared with siPKD2 (Fig. 7E). To further corroborate our findings, cells were transfected with nontargeting siRNA (siNT) or siFbxw7, synchronized at the start of M-phase by nocodazole, treated with CRT101 and cell extracts were analyzed by Western blotting to monitor Aurora A degradation (Fig. 7F). The immunoblot showed that Aurora A was downregulated within 3 hours of CRT101 treatment and this turnover was inhibited by Fbxw7 knockdown. These results suggested that PKD protects Aurora A from Fbxw7-mediated ubiquitination and proteasomal degradation to regulate G2–M transition.

Discussion

PKD as a key target of the second messenger DAG regulates many important cellular functions including cell proliferation. Aberrant expression and activation of PKD have been demonstrated in many cancers and are implicated in oncogenic transformation and tumor progression. PKD2 in particular has been shown to promote tumor cell proliferation in human cancer including carcinoma of the prostate (8). However, the mechanisms through which it regulates tumor cell proliferation have not been well defined. In this study, we identified PKD2 as a cell-cycle regulator that promotes G2–M transition and mitotic entry. PKD2 is activated during G2 and M phases of the cell cycle. It is localized at the centrosome and upon activation regulates centrosome duplication and separation, a key event for the cells to exit G2 and enter mitosis. Mechanistically, we further demonstrated that PKD activity was required for Aurora A stability during G2-M, and it acts through Fbxw7 to protect Aurora A from ubiquitination and proteasomal degradation. Our study has identified PKD2 as a potential G2–M checkpoint kinase. Its aberrant activation may contribute to premature mitotic entry and genome instability that drive tumorigenesis.

A key finding from our study is that PKD is active during G2, and its activity is sustained throughout mitosis (Fig. 1E). Increased PKD2 activity in G2–M was supported by: (i) immunoblotting results showing elevated active form of PKD2 in cells entering G2 and progressing through mitosis (Fig. 1A–D). (ii) IF staining results showing increased active PKD2 staining in cell nuclei during G2 (Fig. 1) and prophase of mitosis, indicative of PKD activation (21). Our finding that activation of PKD is crucial for G2–M cell-cycle progression agrees with the report by Papazyan and colleagues that demonstrated a possible connection of PKD activation during mitosis and its role in cell cycle (33). The effects of PKD inhibitors are in line with previous reports of anticancer effects of PKDs. CRT0866101 (CRT101) in particular has been shown to block cell proliferation by inducing G2–M arrest, downregulation of mitotic regulatory proteins and induction of apoptosis in colorectal (34), bladder (35), pancreatic (36), and prostate (37) cancer cell lines. Our laboratory has also shown that kb-NB142-70, another novel PKD inhibitor causes dramatic G2–M cell-cycle arrest in PC3 prostate cancer cells (38). Both of these two PKD inhibitors were also shown to inhibit HeLa cell proliferation (Supplementary Fig. S3). Collectively, our study has identified PKD as a crucial cell-cycle regulator that is activated during G2–M and promotes mitotic entry. Moreover, in this study, we have provided possible mechanistic insights to PKD inhibitor-mediated cell-cycle arrest, downregulation of cell-cycle regulators and inhibition of cell proliferation, which reflected a positive role of PKD in promoting G2–M transition by modulating Aurora A protein stability.

Aurora A modulates the function of several cell-cycle regulators critical for G2–M transition and mitosis, such as PLK1, CDK1/cyclin B, and CDC25C (18) and is essential for key mitotic events, such as centrosome maturation and separation (31), mitotic entry and formation of mitotic spindles (27). Aurora A must be degraded to ensure proper mitotic exit. During late anaphase, it loses its interaction with TPX2 and is targeted by anaphase-promoting complex/cyclosome (APC/C) ubiquitin ligase for proteasomal degradation (16). Aurora A is also downregulated by Fbxw7, a p53-dependent tumor suppressor (15). Mechanistically, Fbxw7 physically interacts with Aurora A and facilitates the ubiquitination-mediated proteasomal degradation in a Gsk3β-dependent manner. Aberrant expression and proteostasis of Aurora A have been linked to many cancers, thus enabling it as a promising target for therapeutic interventions in cancer (18). Our study has revealed for the first time, a functional link between PKD and Aurora A kinase in G2–M progression and mitotic entry of the cell cycle. We have shown evidence that PKD is critical for late G2 cells to enter mitosis, and that abrogation of its activity delays G2–M

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PKD interferes with Fbxw7-mediated ubiquitination of Aurora kinase A. A, Western blot analysis was performed to show that CRT101-mediated downregulation of Aurora A was proteasome-dependent. B, Half-life of Aurora A protein was measured from the immunoblot of A. C, HeLa cells were synchronized at the start of M phase by nocodazole and preincubated with MG132 for 30 minutes. DMSO (control) or CRT101 was added to the culture and the cells were further incubated for 3 hours. Aurora A was pulled down by IP and the protein was analyzed by Western blotting to assess its ubiquitination by CRT101. Downregulation of Aurora A by knockdown of PKD2 and Cdh1 (D) or PKD2 and Fbxw7 (E) was analyzed by Western blotting (top). Quantification of Aurora A protein levels was performed by densitometry analysis (bottom). F, Western blot analysis depicting that CRT101-mediated downregulation of Aurora A is inhibited by knockdown of Fbxw7. GAPDH was used as loading control. G, A model of regulation of Aurora A protein stability by PKD. The graph represents average of at least three independent experiments with error bars representing SEM (\( \ast, P < 0.5; \ast\ast, P < 0.01; \ast\ast\ast, P < 0.001; \text{ns, not significant})

PKD2 Promotes G2-M Transition via Aurora A

We have shown that loss of PKD has profound effects on cell cycle, such as, (i) rapid downregulation of mitotic regulators including cyclin B1, phospho-CDC25\(^{148}\) (Supplementary Fig. S5), (ii) induction of mitotic catastrophe characterized by defects in spindle formation, apoptosis and cell death, (iii) defects in centriole separation in late G2 and most importantly, (iv) degradation of Aurora A by Fbxw7-mediated proteasomal degradation. Cdh1 is a component of anaphase-
promoting complex that is responsible for substrate degradation during anaphase of mitosis (39). Whereas Fbxw7 (α, β, and γ) expression spans all cell-cycle stages (40), which may be a potential cause for siFbxw7, not siCdh1 to be effective against Aurora A turnover in an asynchronous culture where mitotic cell population is significantly low. Therefore, it is conceivable that a cell harboring inactive or no PKD2 eventually lose Aurora A at the protein level and displays somewhat an "Aurora A null" phenotype (41).

Another significant finding of our study is that PKD2 resides on the centrosome during G2–M and controls centriole separation. We showed that Aurora A and PKD2 colocalized in centrosome in early G2 and throughout G2–M. It is known that the maturation and separation of centrosomes are dependent on the recruitment and activation of Aurora A kinase on centrosomes (42). It is unclear whether PKD2 plays a direct role in recruiting Aurora A to the centrosomes. We did not detect direct interaction between PKD2 and Aurora A although both proteins colocalize in the centrosomes and bind to γ-tubulin. Several possibilities may explain this result: (i) the interaction was limited to the centrosomes, which is difficult to detect using whole-cell lysates, (ii) the interaction may be low affinity and transient like many protein kinases and substrates; (iii) the regulation may be indirect through other regulators. Although Fbxw7 was found to mediate the regulation of Aurora A by PKD, it remains to be determined whether PKD directly regulates Fbxw7 activity. Functionally, it is evident that PKD activity modulates the availability/stability of Aurora A at the centrosomes. Thus, inhibition of PKD results in degradation of Aurora A, which may impede the maturation and separation of centrosomes, and thereby block the entry of cells into mitosis. Klenzle and colleagues have shown that PKD positively controls Golgi complex fragmentation at G2 phase of the cell cycle via Raf–MEK1 pathway, thus, enabling cells to enter mitosis (43). This study is in line with our findings because centrosome duplication and separation is an early step that required for Golgi fragmentation in G2 (44). Thus, the alteration of Aurora A stability and the resulting defects in centrosome separation may be linked to Golgi fragmentation that occurs at G2 phase. Beyond Aurora A, the localization of PKD2 in centrosome, the major microtubule-organizing center of the cell, raises the possibility that it might modulate other centrosomal proteins to ensure tight regulation of cell cycle.

In summary, our results provide strong evidence for a novel function of PKD2 in regulating G2–M transition, mitotic entry and cell-cycle progression. PKD2 does so by protecting critical cell-cycle regulators including Aurora A kinase, an essential mitotic regulator from Fbxw7-mediated proteasomal degradation (Fig. 7G). The regulation of Aurora A by PKD2 provides mechanistic insights to the oncogenic effects of these kinases and may have important therapeutic implications in cancer.

Disclosure of Potential Conflicts of Interest
No potential conflicts of interest were disclosed.

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Administrative, technical, or material support (i.e., reporting or organizing data, constructing databases): Q.J. Wang
Study supervision: Q.J. Wang

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Protein Kinase D2 Modulates Cell Cycle By Stabilizing Aurora A Kinase at Centrosomes

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