Delayed Apoptosis Post-Cadmium Injury in Renal Proximal Tubule Epithelial Cells

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Abstract
Background: Accumulation of the widespread environmental toxin cadmium (Cd) in the kidney results initially in proximal tubule dysfunction. Exposure to Cd has been previously shown to induce apoptosis in LLC-PK (Lily Laboratory Culture, Porcine Kidney) cells, which are a model of proximal tubule epithelium. Hypothesis: We postulated that modulation of the components of the apoptotic pathway triggered by Cd is amenable to therapeutic intervention. Methods: We subjected confluent LLC-PK cells grown on two-compartment filters and on plastic to Cd (1–50 µM). Apoptosis and changes in components of the apoptotic pathway were measured by immunocytochemical and immunoblot analysis during the period of exposure and following Cd withdrawal. Results: Insignificant apoptosis was seen during exposure to Cd and immediately after removal of this metal. Two waves of apoptosis were noted 6 and 48 h after the Cd was removed from the apical compartment. The apoptosis 48 h post-Cd exposure was accompanied by a decrease in cellular ATP levels and transepithelial resistance and preceded by an increase in p38 phosphorylation. Inhibition of p38 mitogen-activated protein kinase activity decreased the delayed apoptotic peak, without affecting the rate of recovery of the integrity of the renal epithelium. IGF-1 neither altered the delayed apoptosis nor facilitated the rate of recovery of the integrity of the renal epithelium. Conclusion: We demonstrate that following exposure to Cd, renal epithelial cells undergo significant apoptosis, which appears to involve p38 and is not amenable to IGF therapy.

Introduction
Cadmium (Cd) is a widespread toxin common in the food chain and in almost every environment [16, 34, 39]. Accumulation of Cd in the kidney results initially in proximal tubule dysfunction, followed by glomerular disease leading to chronic renal damage [39]. Exposure to Cd has been previously shown to induce apoptosis in LLC-PK (Lily Laboratory Culture, Porcine Kidney) cells [18–22, 31, 43], which are a model of proximal tubule epithelium [12, 14]. Although the precise mechanism of Cd-induced apoptosis in renal cells has yet to be elucidated, renal proximal tubular cell damage following a bolus dose of Cd...
is associated with inhibition of mitochondrial respiration [42].

The proximal tubule epithelium of the nephron is also particularly susceptible to damage by other nephrotoxins and by ischemia-reperfusion injury, but displays a remarkable capacity to overcome the damage inflicted and to regenerate [24, 26, 40]. Depending on the intensity of the insult, renal epithelial cells may undergo apoptosis or necrosis [25]. Commonly, ischemia-reperfusion injury results in cellular ATP depletion [25]. A reduction in ATP levels in affected cells to less than 25% of normal values triggers necrosis. In contrast, reduction of ATP levels to 25–50% of normal values causes apoptosis [25].

Necrosis is irreversible and difficult to prevent. By contrast, the apoptotic pathway can potentially be modulated and/or neutralized to maintain cell viability [37]. The understanding of the mechanisms involved in apoptosis, regeneration and repair after acute renal failure remains rudimentary. However, mitogen-activated protein kinase (MAPK) activities, including those of JNK, p38 and ERK, are markedly enhanced after ischemia in vivo and chemical anoxia in vitro, suggesting that they play a role in cellular responses to hypoxic damage [6, 7, 9, 27, 28, 32, 35]. In addition, a number of growth factors and cytokines have been implicated in renal repair after ischemia [8, 27, 30].

Activation of the c-jun N-terminal kinase (JNK) and p38 families of MAPK is an early event in Cd-induced apoptosis in non-renal cell types [2, 3, 10, 11]. Others have suggested that whereas extracellular signal-regulated kinase (ERK) may help to maintain genome integrity and survival, p38 MAPK contributes to mitotic arrest and genome instability [1]. Importantly, others have suggested that insulin growth factor-1 (IGF-1) may play a role in modulating Cd nephrotoxicity [20].

To better understand the events that regulate initiation of apoptosis and regeneration, we subjected confluent LLC-PK1 cells to Cd-induced injury, and examined them during the recovery phase following Cd withdrawal. Under these conditions, there was insignificant apoptosis during exposure to Cd and immediately after removal of this metal. However, two waves of apoptosis occurred during the recovery phase. We postulated that modulation of the components of the apoptotic pathway triggered by Cd may be amenable to therapeutic intervention. We observed that inhibition of p38 activity decreased the delayed apoptotic peak, without affecting the rate of recovery of the integrity of the renal epithelium. In contrast, IGF-1 neither altered the delayed apoptosis nor facilitated repair of the monolayer.

Methods

Cell Culture
LLC-PK1 cells (American Type Culture Collection (CT-101), Bethesda, Md., USA) were cultured in antibiotic-free Dulbecco’s modified Eagle’s Medium (DMEM) supplemented with 8% fetal bovine serum (FBS) and passaged twice a week. For experiments, the cells were plated at a density of 10^5 cells/cm^2 (confluent) on either plastic or on cell culture inserts (0.4 or 1.0 μM; Becton-Dickinson) and cultured for 5 days, with daily growth medium changes. This ensured the establishment of a well-developed, polarized epithelial monolayer [12–15]. On day 6, the cells were washed twice with PBS and incubated in FBS-free DMEM for 12 h prior to the addition of CdCl2 (0–50 μM). Following a 12- or 24-hour incubation in the presence of Cd, the epithelial monolayer was washed to 4–6 times with PBS to remove all extracellular Cd. The cells were allowed to recover by culture in normal growth medium for intervals indicated in individual experiments. To investigate the effects of IGF-1, LLC-PK1 cells were cultured in growth medium containing human IGF-1 (100 ng/ml; final; R&D Systems). To investigate the effects of p38 inhibitor (Calbiochem) or PD98059 (Sigma), cells were treated for 4 h with the inhibitor immediately prior to the peak rise in caspase-3 activity (24 h post-Cd). This short treatment time with the inhibitors ensured lack of cell death due to prolonged drug exposure.

Cellular ATP Quantification

Cellular ATP was measured by light emission using a luciferin-luciferase assay kit (Sigma). Cells cultured in 12-well plates were treated with CdCl2 or with vehicle. Luminescence was measured on a Microtiter Plate Lumimeter (Dynex Technologies). ATP levels were normalized to protein content, determined separately using Bradford assays (Bio-Rad).

Apoptosis

Apoptosis was assessed by examining nuclear morphology/chromatin condensation, and by measuring active caspase-3 levels in immunoblots. To analyze nuclear morphology, cells were washed twice with PBS and their DNA was stained with Hoechst 33258 (1 μg/ml; Sigma) for 10 min at 37°C. Staining medium also included propidium iodide (1 μg/ml; Sigma) or Sytox Green (1 μg/ml; Molecular Probes, Portland, Oreg., USA). Cells were washed with PBS, fixed with 4% paraformaldehyde, and rinsed with PBS. The frequency of apoptosis or necrosis in ten random fields (300 cells/field, 400 x magnification) was determined using an epifluorescence microscope (DMIRBE, Leica, Germany) attached to a cooled CCD camera (Ocea II, Hamamatsu, Japan) driven by Openlab Digital Imaging Software (Improvision, UK).

Atomic Absorption Spectroscopy

LLC-PK1 cells treated with CdCl2 or with vehicle were collected, centrifuged for 10 min at room temperature and then re-suspended in 1 ml of PBS. One tenth of the cells were used for protein determination. 1 ml of nitric acid was added to the remaining sample and left overnight at room temperature to digest. The digested samples were then incubated for 3 h at 110°C and then diluted with double-distilled water to a final volume of 2 ml. Metal content of the samples was quantified using a Varian Spectra A30 Atomic Absorption Spectrometer against certified standards.
**DNA Synthesis**

[3H]d-thymidine incorporation into DNA was measured as previously described [5].

**Mitotic Index**

Hoechst 33258 stains all nuclei and allowed us to visualize mitotic nuclei. The mitotic index was calculated by expressing the number of mitotic nuclei as a percentage of the total nuclei in ten random fields (300 cells/field, 400 × magnification).

**Immunoblot Analysis**

LLC-PK1 cells treated with CdCl₂ or with vehicle were harvested and total cellular protein isolated as previously described [4, 5]. Protein concentrations were determined by the Bradford method (Bio-Rad). Lysates containing 50–100 μg protein/lane were resolved by denaturing polyacrylamide gel electrophoresis (SDS-PAGE) and transferred to PVDF membranes (Roche). Membranes were probed overnight at 4°C with one of the following antibodies purchased from New England Biolabs: Active caspase-3 (Cat. #9661), phospho-AKT (#9271S), AKT (#9272), phospho-p38 (#9211S), p38 (#9212), phospho-p42/44 (#9101S) and p42/44 (#9102). Following incubation with appropriate horseradish peroxidase-labeled secondary antibodies, proteins were visualized by enhanced chemiluminescence (ECL Plus, Amersham Pharmacia). All immunoblots were also probed for β-tubulin to normalize for protein loading. No significant differences in loading were noticed between samples in a given immunoblot using this method. Protein levels were estimated by densitometric analysis of the immunoblots (AlphaImager 1220 V5.5 Software, AlphaInnotech, Calif., USA). For blots with phospho-specific antibodies, normalization was also conducted with respect to total levels of the relevant protein.

**Measurement of Transepithelial Resistance (TER)**

LLC-PK1 cells were plated at confluence on polyethylene terephthalate (PET) membrane inserts (1.0-μm pore size) in 24-well HTS Multiwell Insert System plates (Becton-Dickinson). An EVOM Ohmmeter (World Precision Instrument, Fla., USA) was used to measure TER. Since the EVOM is sensitive to changes in media volume, we ensured that 500 μl of medium was consistently added to each well (upper chamber) and 10 ml was present in the common bottom plate. TER measurements were compared to (1) an untreated monolayer growth medium and (2) the initial TER of each monolayer prior to incubation with CdCl₂.

**Data Analysis**

SPSS10 software was used for statistical analysis (Student’s t and non-parametric Mann-Whitney). All data are expressed as mean ± SEM. Significance was established at p ≤ 0.5.

**Results**

**Effects of Acute Cd Administration on LLC-PK Cells**

To begin to address the mechanisms of injury and regeneration of renal epithelial cell monolayer following Cd injury, we first established the extent and nature of damage to mature LLC-PK1 monolayers following exposure to that metal. Cells were plated at confluence, cultured for 5 days in normal growth medium, followed by 12 h of culture in serum-free medium to obtain a synchronized, quiescent cell population similar to proximal tubule epithelia. The cells were subsequently treated with various CdCl₂ concentrations in serum-free medium, to avoid Cd sequestration by serum components. Immediately following Cd treatment, insignificant apoptosis was seen across a broad dose range (1–20 μM; fig. 1a). With higher Cd concentrations or longer treatments, substantial cell death ensued, largely due to necrosis (fig. 1a and data not shown).

To verify that the observed cell loss was due to Cd-induced intracellular damage, we measured cellular Cd levels by atomic absorption spectroscopy. The results of these analyses demonstrated that the cellular Cd uptake after 12 h was proportional to extracellular Cd concentration (control: 0.01 μg Cd/μg protein; 10 μM CdCl₂: 0.7 μg Cd/μg protein).

Cd is a powerful inhibitor of mitochondrial function, thus we examined whether the Cd-induced cell death observed was correlated to changes in cellular ATP content. As shown in figure 1b, incubation of the cells with increasing CdCl₂ concentrations induced a dose-dependent decrease in cellular ATP. Notably, ATP decreases greater than 40% were associated with increased necrosis, whereas the fraction of apoptotic cells was less sensitive to substantial ATP depletion.

**Response during Recovery from Cd-Induced Exposure**

To examine the recovery mechanisms subsequent to Cd-induced injury in LLC-PK1 cells, we determined the fraction of apoptotic cells at timed intervals after Cd removal. We postulated that the induction of apoptosis by Cd was a specific response that mediated removal of damaged cells to allow undamaged cells to recapitulate the epithelial monolayer. We observed a significant increase in apoptosis 3 h after CdCl₂ removal, which was maintained over the first 24 h post-Cd (fig. 2a). The apoptotic response of the cells appeared to be biphasic, with the second wave reaching maximum levels as high as 12-fold above those in untreated cultures at 48 h of recovery (fig. 2a).

Cells enter apoptosis asynchronously. Moreover, DNA fragmentation is not evident in all apoptotic cells. Active caspase-3 was measured by immunoblot to complement the morphological assay. We observed a steady increase in active caspase-3 levels as a function of time, reaching maximum values at 24–48 h of recovery (fig. 2a), correlating with the maximum apoptotic response observed at 48 h (fig. 2a).
Fig. 1. Cell death and cellular ATP content as a function of CdCl₂ concentration. A) LLC-PK cells plated at confluence and grown for 5 days were serum starved for 12 h and then incubated with CdCl₂ for 12 h. Cells were stained with Hoechst 33258 (1 μg/ml; Sigma) and/or propidium iodide (1 μg/ml; Sigma) or Sytox Green (1 μg/ml; Molecular Probes) for 10 min at 37 °C. Each point represents the mean of 4 experiments of triplicate specimens, from which the frequency of apoptosis or necrosis (propidium iodide or Sytox Green-positive cells) in ten random fields (300 cells/field, 400 × magnification) was determined using an epifluorescence microscope (DMIRBE, Leica) attached to a cooled CCD camera (Orca II, Hamamatsu) driven by Openlab Digital Imaging Software (Improvision). The error bars are the SEM. Square = Serum starved; ■ = CdCl₂. B) Cellular ATP was measured by light emission using a luciferin-luciferase assay kit (Sigma). Luminescence was measured on a Microtiter Plate Luminometer (Dynex Technologies). ATP levels were normalized to protein content, determined separately using the Bradford assay (Bio-Rad). Each point represents the mean of 4 experiments of triplicate specimens and the SEM.

Fig. 2. Cell death, cellular ATP content and TER as a function of time following CdCl₂ exposure. A) LLC-PK cells plated at confluence and grown for 5 days were serum starved for 12 h, incubated with 10 μM CdCl₂ for 12 h and then washed (0 h) and fed with normal medium augmented with 8% FBS. The number of apoptotic cells (■, solid line) was counted in 10 random fields (300 cells/field) in triplicate. Relative abundance (-fold increase) of active caspase-3 (●, broken line) was quantified by densitometric analysis of the immunoblots (AlphaImager 1220 V5.5 Software, AlphaInnotech). Loading was normalized to β-tubulin. B) Cellular ATP was determined by the luciferin/luciferase assay and C) TER using EVOM ohmmeter (World Precision Instrument) in LLC-PK cells plated at confluence, grown for 5 days after 12 h serum starvation and 12 h incubation with 10 μM CdCl₂ (0 h) or after 12 h of serum starvation cell cultures (●) and then after the cells were washed and fed with normal medium supplemented (3–120 h) with 8% FBS. Each point represents the mean of 4 experiments done in triplicate and SEM.
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Fig. 3. Mitotic index and [3H]-thymidine incorporation as a function of time following CdCl₂ exposure. A The number of mitotic bodies was counted in 10 random fields (300 cells/field) and B [3H]-thymidine incorporation measured in triplicate in LLC-PK cells plated at confluence, grown for 5 days after 12 h serum starvation and 12 h incubation with 10 μM CdCl₂ (0 h) (■, broken line) or after 24 h of serum starvation cell cultures (0 h) (●, solid line) and then after the cells were washed and fed with normal medium supplemented (3–120 h) with 8% FBS. Each point represents the mean of 4 experiments done in triplicate and SEM.

The delayed apoptosis during the recovery phase suggest that Cd may be taken up, and stored intracellularly to interfere with various functions, including mitochondrial energy production. Hence, we determined if the relationship between decreased cellular ATP levels and onset of apoptosis was maintained during the recovery phase. Consistent with this concept, there was a time-dependent drop in cellular ATP levels to a nadir of 60% of untreated cells at 48 h of recovery, coinciding with the time of maximum observed apoptosis (fig. 2b), and slowly recovered to normal levels over the next 3 days (fig. 2b).

We also measured the TER of the cell monolayer, which is a functional indicator of epithelial barrier function. Consistent with the observed changes in ATP levels and apoptosis, TER values decreased to 60% of control by 48 h of recovery. TER values remained low until 72 h, and slowly reached normal levels at 120 h (fig. 2c). Thus, delayed Cd toxicity in vitro during the recovery phase results in cell loss, which in turn results in loss of barrier integrity, similar to renal Cd toxicity in vivo.

Proliferative Responses of Cd-Treated LLC-PK1 Cells during the Recovery Phase

According to the current understanding of renal epithelial regeneration, a nephrotoxic insult that results in cell death and loss of epithelial barrier integrity also induces mitogenic responses in the remaining cell population. Thus, viable cells will divide and migrate to cover denuded areas and re-establish renal function. To investigate whether Cd-induced damage would trigger cell division in the LLC-PK1 monolayers, we determined changes in DNA synthesis and mitotic index in untreated and in Cd-damaged cells. Addition of serum to control cells following a 24-hour incubation in serum-free medium resulted in a 1.5-fold increase in [3H]dThd incorporation into DNA, which was maintained for 96 h. This increase likely stems from stimulation of quiescent cells to re-enter the cell cycle after serum starvation. The decrease in dThd incorporation noted 120 h following initial serum addition likely reflects the activation of contact-inhibition mechanisms (fig. 3a). Addition of serum to cells cultured in serum-free medium for 24 h, and in the presence of CdCl₂ for the last 12 h, resulted in two waves of DNA synthesis within 120 h after serum addition. Specifically, a significantly greater, 2.5-fold increase in [3H]dThd incorporation into DNA was observed by 3 h, a time course similar to that found in untreated cells (fig. 3a). This greater increase likely reflects a greater proportion of quiescent cells advancing to the S-phase of the cell cycle as a result of Cd-induced damage and/or neighboring cell loss. The maximum increase in DNA synthesis between 3 and 6 h after serum addition was followed by a peak in the mitotic fraction 3- to 4-fold greater than in control cultures at 24 h (fig. 3b), thus mimicking the renal epithelial regenerative response in vivo. Thus, it appears that, in the recovery stage following Cd-induced injury, a fraction of the cell population is recruited back into the cell cycle, whereas another portion of the cells undergoes apoptosis. These two responses follow distinct time courses, with a
Regulation MAPK in Cd-Induced LLC-PK1 Cell Apoptosis

The involvement of several MAP kinases in Cd injury and apoptosis in non-renal cells has been demonstrated. Further, activation of the MAPK cascade is an essential element of mitogenic responses to serum. Specifically, p38, JNK and ERK play an important role in these responses [2, 10]. We assessed changes in the levels of phosphorylated kinases, which are a measure of their activation. Culturing cells for 24 h in the absence of serum did not abolish ERK42/44 expression, but levels of phosphorylated ERK in these cells were negligible in the absence or presence of Cd (fig. 4a). Addition of serum resulted in rapid transient activation of ERK p42/p44 within the first 60 min (data not shown), followed by a smaller, but sustained increase in phosphorylated ERKp42/p44 levels in cells cultured in the absence of CdCl2, as late as 96 h, similar to ERK p42/44 abundance in other proliferating cells [pers. unpubl. data]. The reduction in mitotic index and peak in mitotic activity at 24 h of recovery, and maximum apoptosis at 48 h.

Fig. 4. Relative abundance of total and phosphorylated ERK and p38 as a function of time following CdCl2 exposure. A Representative immunoblots showing the relative abundance of total and phosphorylated ERK and B total and phosphorylated p38 in LLC-PK cells plated at confluence and grown for 5 days were serum starved for 12 h, incubated with 10 μM CdCl2 in serum-free medium for 12 h and then washed and fed with normal medium augmented with 8% FBS. Culturing cells for 24 h in the absence of serum did not abolish ERK42/44, but levels of phosphorylated ERK in these cells were negligible in absence or presence of Cd. Similar to total ERK p42/p44, total levels of p38 did not appreciably change in Cd-treated or -untreated cultures upon serum addition. Phosphorylated p38 levels in untreated cultures did not change early after serum stimulation, but appeared to increase between 36 and 96 h, and returned to barely detectable levels by 120 h. The immunoblots are representative of 4 separate experiments.

DNA synthesis observed at 120 h in untreated cultures was accompanied by a reduction in phosphorylated ERK abundance. Those cells recovering from CdCl2 treatment also showed an early increase in phosphorylated ERK p42/p44. In contrast to the untreated cells, these substantial increases in the levels of active ERKp42/p44 upon serum addition were maintained throughout the time course of analysis, reaching a maximum of about 8-fold above those observed prior to serum stimulation (fig. 4a).

Similar to total ERK p42/p44, total levels of p38 did not appreciably change in Cd-treated or -untreated cultures upon serum addition (fig. 4b). Phosphorylated p38 levels in untreated cultures did not change early after serum stimulation, but appeared to increase between 36 and 96 h, and returned to barely detectable levels by 120 h, a time during which DNA synthesis and mitosis were minimal (fig. 4b). Phosphorylated p38 was undetectable in Cd-treated cultures early after serum addition, but increased by 6 h of recovery, to a maximum at 36 h, just preceding the peak in cellular apoptosis (fig. 2a, 4b). Thus, Cd-induced injury differentially modulates the responses of two stress- and mitogen-modulated MAP kinases, like-
Fig. 5. Effect of p38 and ERK inhibition on apoptosis and TER after 10 μM CdCl₂. LLC-PK cells plated at confluence and grown for 5 days were serum starved for 12 h, incubated with 10 μM CdCl₂ for 12 h and then washed and fed with normal medium augmented with 8% FBS. At 20 h after the CdCl₂ was removed, either p38 inhibitor (p38I) or PD98059 was added to the media for 4 h and the number of apoptotic cells counted (A, C) and the relative abundance of active caspase-3 measured (B, D). In addition (E), we measured the TER of the monolayer in the presence or absence of p38I (λ, 24 h serum starved, p38I: μ, 12 h CdCl₂, p38I) or PD98059 (σ, serum starved, PD98059: Δ, 12 h CdCl₂, p38I) in cells starved for 12 h and incubated with 10 μM CdCl₂ for 12 h (θ) or serum starved for 24 h (υ) prior to washing and refeeding with serum supplemented medium. Each point represents the mean of 4 experiments done in triplicate and SEM. * p < 0.05.

ly reflecting a differential role for these two proteins in the recovery phase following Cd²⁺-induced injury.

In this model, no alterations in total or phosphorylated JNK were seen in response to Cd (data not shown).

Functional Role of p38 and ERK p42/p44 in Cd²⁺-Induced Delayed Apoptosis

Given that the most pronounced increments in phosphorylated p38 and in ERKp42/p44 occurred just prior to the second apoptotic wave during recovery, we examined the involvement of these kinases in the apoptotic response. LLC-PK1 cells to CdCl₂ were treated as before, but included ERK or p38 inhibitors were included from 20 to 24 h post-Cd. We reasoned that, as both apoptosis and active caspase-3 levels were substantial at that time, the use of MAPK inhibitors would provide information about their involvement in the apoptotic response. Control experiments using a 4-hour incubation with p38I concentrations demonstrated to abolish p38 activity did not significantly alter the apoptotic cell fraction or active caspase-3 levels in our cultures (fig. 5a, b). This time of incubation was maintained in all subsequent experiments, as prolonged p38I treatment can be cytotoxic by itself [10, 11]. The presence of p38I during recovery in Cd²⁺-treated cultures resulted in significant decreases in apoptosis and active caspase-3 levels (fig. 5a, b).

We next investigated the effect on the apoptotic cascade of abrogating ERK p42/p44 activation by interfering...
IGF-1 and Cd²⁺-Induced Apoptosis

We determined whether IGF-1 afforded protection against the biphasic apoptotic response observed in CdCl₂-injured cells. The results of these experiments showed that IGF-1 was able to prevent the earlier apoptotic wave at 6 h of recovery, but did not affect the second, more pronounced apoptotic phase at 48 h of recovery (fig. 6a), demonstrating that distinct mechanisms likely mediate the two waves of apoptosis. IGF-1 did not significantly alter the time course of recovery of TER function (fig. 6b). To confirm that IGF-1 signaling pathways were activated in the presence of this growth factor, we measured levels of phosphorylated PKB/AKT, a known IGF-1 target. We observed an increase in phosphorylated PKB/AKT levels in cells treated with IGF-1 with or without prior incubation with CdCl₂, demonstrating appropriate signaling through the IGF-1 receptor (fig. 6c). In addition, the presence of IGF-1 did not interfere with increments in either activated caspase-3 or phosphorylated p38 levels associated with apoptosis, consistent with the proposed involvement of these proteins in Cd²⁺-induced apoptosis in LLC-PK1 cells (fig. 6c).

Discussion

The objective of this study was to examine the mechanisms underlying Cd-induced apoptosis in the course of recovery from an acute insult. With the advent of better-characterized cell lines and culture techniques, in vitro models are becoming more useful in toxicological research [14, 41]. Pointedly, most of the previous work on
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Cadmium-induced apoptosis in vitro has focused on the acute event [20, 43]. To the best of our knowledge, this is the first series of experiments to explore the recovery of an epithelial cell monolayer from an acute insult with Cd over a 5-day period. By extending the period of examination to 5 days post-exposure to Cd, we demonstrated that Cd induces a prodigious amount of apoptosis via a p38-dependent mechanism 48 h post-exposure, which may be associated with depletion in cellular ATP, induced by intracellular Cd disruption of mitochondrial respiration.

Whether Cd induces necrosis or apoptosis depends on the extracellular concentration and appears to reflect the degree of cellular ATP depletion induced by exposure to the metal. In vivo, renal proximal tubule epithelial cell damage following a bolus dose of Cd-metallothionein involves disruption of mitochondrial respiration, secondary to the accumulation of Cd [42]. Others have determined that ATP depletion to about 15–25% of control represents a threshold that determines whether proximal tubule epithelial cells die by necrosis or apoptosis [25]. Consistent with these studies, high concentrations of Cd induced acutely severe drops in cellular ATP content, resulting in necrosis, whereas lower doses had lesser effects on cellular ATP and produced minimal necrosis and/or apoptosis.

The late prodigious apoptosis seen 48 h post-Cd is also associated with cellular ATP content. As previously reported [20], in vitro LLC-PK cells undergo apoptosis soon after removal of Cd. Our data indicate that the immediate apoptosis is minor at best and that a more significant event takes place later in the recovery phase. In other experimental models of acute renal failure in vivo, apoptosis of renal tubule epithelial cells has been shown to occur in two distinct phases [23]. The first phase of apoptosis occurs between 12 and 48 h after the acute ischemic or nephrotoxic insult. The second phase of apoptosis occurs many days later, during the recovery phase. Tubular cell apoptosis occurring shortly after the acute insult probably contributes to tubular cell loss and the tubular dysfunction associated with injury. In contrast, the apoptosis associated with the recovery phase has been postulated to contribute to the remodeling of injured tubules. In the present study, the ongoing apoptosis seen upon withdrawal of Cd correlates with a continuing decrease in cellular ATP content. Thus, it is likely that a significant portion of Cd toxicity may be secondary to the lingering effects of intracellular Cd on mitochondria well after the withdrawal of the metal from the extracellular environment, but reflecting the first phase of injury following the acute nephrotoxic insult. It is likely that if we had continued to follow our culture system past the 5 days, a second remodeling phase of apoptosis would have been apparent.

Recovery from a renal insult that causes cell loss is associated with epithelial cell proliferation and tubular remodelling. Consistent with this concept, we observed waves of DNA synthesis and mitotic activity in the recovering monolayers. Thus, it appears that, in the recovery stage following Cd-induced injury, a fraction of the cell population is recruited back into the cell cycle, whereas another portion of the cells undergoes apoptosis. These two responses follow distinct time courses, with a peak in mitotic activity at 24 h of recovery, and maximum apoptosis at 48 h.

The activation of p38 by Cd has been previously noted in non-renal cells [2, 3, 10, 11, 17]. At least two of these studies find that p38 activation is an early and specific regulatory event for the Cd-provoked apoptosis [1, 10]. None of these investigators report a delayed activation of p38. Pointedly, the concentrations of Cd used by these studies to induce p38 activity acutely were 4- to 20-fold greater than we used in our work, although the length of incubation was less [1–3, 10, 11, 17]. However, it should be noted that LLC-PK cells have been previously shown to be more sensitive than other cells to Cd [31, 43]. Lower cytotoxic doses (<80 µM) were reported to also activate JNK and ERK, but not p38 [2]. Although we note an increase in the abundance of phosphorylated ERK, neither native nor phosphorylated JNK was altered in our model.

A number of investigators have postulated that minimizing or inhibiting p38 activity or apoptosis may accelerate repair [10]. Whereas the later phase of apoptosis seen 48 h after the Cd pulse was associated with an increase in active p38, the early minimal rise in regulated cell death was not. Although judicious use of p38 inhibitor clearly blocks the secondary apoptotic wave, we see little improvement in the rate at which the epithelial monolayer re-establishes integrity. This may reflect the short period of inhibition or that other factors are involved in the regulating integrity of the epithelial monolayer. TER, which we used to measure epithelial integrity, is in fact a measure of the integrity of tight junctions. Tight junction recovery is a regulated process dependent on tyrosine phosphorylation [29]. Although Cd causes ATP depletion and changes in the cytoskeleton, the pattern of changes in the cytoskeleton and junctional proteins is different from that seen in chemical anoxia or ischemia [33, 36, 38, 43], suggesting that this metal may have other more significant effects.
The inability of IGF-1 to inhibit the late apoptotic peak while blocking the earlier cell death is intriguing. Although much has been made of the anti-apoptotic properties of IGF-1 acting through PI3-K and AKT in ischemic injury, renal growth factors do not ameliorate apoptotic cell death induced by ATP depletion [25]. Importantly, IGF-1 inhibited the p38-independent apoptosis seen in the first 6 h, when the ATP content was still >75% of control. Clearly, the mechanism by which the cells undergo apoptosis in the second wave is either different from the first or simply not amenable to modification by AKT. We are in the process of delineating the upstream regulation and downstream targets of p38 activation during the second wave to better understand the regulated cell death.

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