Uptake, p53 Pathway Activation, and Cytotoxic Responses for Co(II) and Ni(II) in Human Lung Cells: Implications for Carcinogenicity

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Cobalt (Co) and nickel (Ni) ions display similar chemical properties and act as hypoxia mimics in cells. However, only soluble Co(II) but not soluble Ni(II) is carcinogenic by inhalation. To explore potential reasons for these differences, we examined responses of human lung cells to both metals. We found that Co(II) showed almost 8 times higher accumulation than Ni(II) in H460 cells but caused a less efficient activation of the transcriptional factor p53 as measured by its accumulation, Ser15 phosphorylation, and target gene expression. Unlike Ni(II), Co(II) was ineffective in downregulating the p53 inhibitor MDM4 (HDMX). Co(II)-treated cells continued DNA replication at internal doses that caused massive apoptosis by Ni(II). Apoptosis and the overall cell death by Co(II) were delayed and weaker than by Ni(II). Inhibition of caspases but not programmed necrosis pathways suppressed Co(II)-induced cell death. Knockdown of p53 produced 50%–60% decreases in activation of caspases 3/7 and expression of 2 highly upregulated proapoptotic genes PUMA and NOXA by Co(II). Overall, p53-mediated apoptosis accounted for 55% cell death by Co(II), p53-independent apoptosis for 20%, and p53/caspase-independent mechanisms for 25%. Similar to H460, normal human lung fibroblasts and primary human bronchial epithelial cells had several times higher accumulation of Co(II) than Ni(II) and showed a delayed and weaker caspase activation by Co(II). Thus, carcinogenicity of soluble Co(II) could be related to high survival of metal-loaded cells, which permits accumulation of genetic and epigenetic abnormalities. High cytotoxicity of soluble Ni(II) causes early elimination of damaged cells and is expected to be cancer suppressive.

Key Words: cobalt; nickel; toxicity; cancer; TP53; apoptosis.

Cobalt (Co) is a toxic metal with significant public health concerns associated with its occupational exposures and release by commonly used cobalt-chromium hip prostheses (Delaunay et al., 2010; Nordberg, 1994; Simonsen et al., 2012). Metallic and other forms of Co are used in numerous industrial applications such as electroplating, in manufacturing of high-temperature alloys, diamond polishing tools, carbides, and hard metals, and as catalysts and pigments. Exposure to Co can result in a series of adverse health effects that include impaired thyroid function, cardiomyopathy, polycythemia, and neurotoxicity. In the inhalation exposures, the main target of Co toxicity is the respiratory system. Occupationally exposed workers are at risk of developing interstitial lung fibrosis, alveolitis, and asthma. These and other toxic effects have also been found in the respiratory tract of rodents after chronic inhalation of Co(II) (Bucher et al., 1990). The 2-year inhalation study by the National Toxicology Program (NTP) found strong evidence for carcinogenicity of soluble Co(II) (cobalt sulfate) in the lungs of mice and rats (Bucher et al., 1999). Occupational exposures to certain Co-containing materials have also been associated with elevated risks of lung cancers (Simonsen et al., 2012). The mechanisms of lung carcinogenicity by Co(II) are currently poorly understood. Co(II) is clastogenic in mammalian cells but only weakly mutagenic in the Ames reversal assay (De Boeck et al., 2003). Ni(II)-containing compounds are classified as carcinogenic to humans (Goodman et al., 2011; Muñoz and Costa, 2012; Salnikow and Zhitkovich, 2008). Inhalation exposures to particulate Ni(II) forms have been found to cause respiratory tumors in humans and rodents. Although the intracellular release of Ni(II) ions is generally believed to be responsible for carcinogenic activity of the particulate forms, soluble Ni(II) (nickel sulfate) was inactive in the 2-year inhalation study by the NTP (Dunnick et al., 1995) and epidemiological findings for soluble Ni(II) have largely been negative (Goodman et al., 2011). Ni(II) and Co(II) are adjacent members of the first series of transition metals and they have similar ionic radii and other chemical properties. Both metals act as chemical hypoxia mimics by activating the hypoxia-responsive transcriptional factor HIF-1 (Salnikow et al., 2000, 2004; Wang and Semenza, 1995) and can catalyze Fenton-type reactions with hydrogen peroxide when complexed with appropriate ligands (Huang et al., 1995;
Van den Broeke et al., 1998), Ni(II) and Co(II) display strong binding affinity for soft-type ligands such as sulfur and nitrogen (Nieboer and Richardson, 1980). This ligand preference is responsible for the ability of Co(II) and Ni(II) to displace Zn(II) from zinc-finger proteins such as the DNA repair factor XPA (Bal et al., 2003; Kopera et al., 2004). Thus, based on many chemical and biological similarities, one could expect that soluble Co(II) and Ni(II) forms should have similar carcinogenic properties, which was clearly not the case in the NTP inhalation studies (Bucher et al., 1999; Dunnick et al., 1995).

In order to understand potential reasons for different respiratory carcinogenicity of soluble Co(II) and Ni(II), we conducted a detailed study of uptake, stress responses, and survival of human lung cells treated with these metals. We found that human lung cells displayed several times higher accumulation of Co(II) but showed a delayed and weaker activation of replication checkpoint, the stress-responsive transcription factor p53, and apoptosis. In contrast, Ni(II) induced a rapid and more robust apoptosis associated with a much earlier activation of p53. High tolerance of Co(II) by human lung cells, permitting DNA replication and survival despite massive internal doses of this redox-active metal, can promote accumulation of genetic damage and helps explain its lung carcinogenicity.

MATERIALS AND METHODS

Chemicals. CoCl$_2$·6H$_2$O (BioReagent grade), NiCl$_2$·6H$_2$O (BioReagent grade), and nitric acid (>99.999% purity) were obtained from Sigma-Aldrich. QVD-OpH was purchased from Innogenex. E64d, cyclosporin A, necrostatin-1, and ALLN were from Santa Cruz Biotechnology. All other reagents were purchased from Sigma-Aldrich.

Cells and treatments. H460 human lung epithelial cells were purchased from the American Type Culture Collection (ATCC) and maintained in RPMI-1640 medium (Gibco 11875-093) supplemented with 10% fetal bovine serum and penicillin/streptomycin. IMR90 human lung fibroblasts were obtained from ATCC and propagated in Dulbecco’s modified Eagle medium (Gibco 12430-062) containing 15% serum and antibiotics. Primary human bronchial epithelial (HBE) cells were obtained from Lonza and maintained in serum-free basal medium (Lonza CC-3170) supplemented with growth factors and standard antibiotics. H460 and HBE cells were grown in 95% air/5% CO$_2$, whereas IMR90 fibroblasts were kept in 5% CO$_2$, and 5% CO$_2$. Exposure to metals was done for 48 h, with a 24-h exposure between media changes.

Small hairpin RNA. The pSUPER-RETOV vector was used for a stable downregulation of p53 expression in H460 cells. Empty and scrambled small hairpin RNA (shRNA)-coding vectors were obtained from Oligoengine. The p53-targeting sequence was GAGTCCGAGTGGATAATCTAT. Retrovirus packaging, infections, and puromycin selection have been described previously (Reynolds et al., 2004). Cells infected with control and targeting vectors were grown in a medium containing 1.5 μg/ml puromycin.

Western blotting. Soluble cellular proteins were extracted as described previously (Reynolds and Zhithkovich, 2007). Proteins were separated by SDS-polyacrylamide gel electrophoresis and electrotransferred to ImmunoBlot PVDF membrane (Bio-Rad). Primary antibodies for PARP (9542), pS15-p53 (9284), pS46-p53 (2521), cleaved caspase-3 (9661), and cleaved caspase-7 (8438), BCL2 (2870), BCL-Xs (2764), MCL1 (5453), and pS70-BCL2 (2827) were from Cell Signaling Technology; for p21 (S118) from BD Biosciences; for MDM4 (A300-287A) from Bethyl Laboratories; for p53 (DO-1) from Santa Cruz Biotechnology; and for γ-tubulin (66577) from Sigma.

Cytotoxicity. Cell death was assessed by measurements of leakage of cellular lactate dehydrogenase (LDH) into media. LDH activity was determined by a commercial kit (Cayman Chemical). Cells were seeded into 96-well plates (10,000–20,000 cells/well) and treated with metals the next day. Due to high LDH levels in serum, cells were incubated in serum-free media for the determination of leaked cellular LDH. At the collection times, plates were centrifuged at 400 x g for 5 min, and media were aspirated for the determination of leaked LDH according to the manufacturer’s instructions. With the exception of the initial experiments in Figure 2, the amount of LDH leaked into the media was expressed as a percentage of total (leaked and cellular) LDH activity in each sample. Cellular LDH activity was determined after removal of growth medium and lysis of cells in fresh RPMI-1640 medium containing 0.1% Triton X-100 (10 min, room temperature). Cytotoxic effects of metals were also examined by the determinations of metabolic activity of cell populations using the CellTitre-Glo luminescent cell viability assay (Promega), which quantifies cellular ATP levels. These measurements integrate the size of cell populations and metabolic activity of individual cells, making it sensitive to all forms of cytotoxicity, such as growth arrest, cell death of any kind, and impairment of cellular metabolic activity. H460 cells were seeded into black 96-well optical bottom cell culture plates (2000 cells/well) and grown overnight prior to the addition of metals. The assay reagents were added to the plates, and luminescence measurements were recorded after 10-min incubation at room temperature. Caspase 3/7 activity was measured by a luminescence-based kit from Promega. Cells were seeded into 96-well plates (2000 cells/well) and treated with metals the next day. Caspase activity assay was performed as recommended by the manufacturer.

Clonogenic survival. A long-term survival of metal-treated cells was assessed by clonogenic assay, which measures the ability of individual cells to retain long-term proliferative activity and produce colonies. For determination of a role of p53 in clonogenic viability, sh-SCR and sh-p53 H460 cells were seeded onto 60-mm dishes (400 cells/dish) and treated with metals for 24 h on the next day. At 7–8 days postexposure, cells were fixed with methanol and stained with a commercial Giemsa solution (Sigma-Aldrich). Colonies containing 30 or more cells were scored. In clonogenic experiments comparing Co(II) and Ni(II) toxicity, H460 cells were treated with metals for 24 h in mass cultures as described in uptake studies. Floating and attached cells were combined and 400 cells/dish were seeded for the determination of clonogenic survival.

Quantitation of attached cells. The number of attached cells was determined using the CyQUANT reagent from Invitrogen. At the indicated times, media were removed and cells were incubated with a DNA-staining solution for 30 min at 37°C. Fluorescence readings were recorded with 485 nm excitation and 530 nm emission settings.

Quantitative reverse transcriptase PCR. Total RNA from H460 cells was isolated using a TRIzol reagent (Invitrogen). Purity of RNA samples was assessed by a Nanodrop spectrophotometer. Complementary DNA was synthesized by the RT First Strand kit as instructed by the manufacturer (SABiosciences). PCR reactions were performed using primers from SABiosciences and the ABI 7900HT Real-Time PCR system (Applied Biosystems). The housekeeping genes B2M, HPRT1, RPLPO, and ACTB were used for normalization purposes. Differences in gene expression were calculated by the 2-ΔΔCT method.

Cellular uptake of metals. Cellular amounts of Co and Ni were determined by graphite furnace atomic absorption spectroscopy (GF-AAS) using nitric acid extracts. Cells were seeded into 6-well plates (H460: 3 x 10$^5$ cells/well; IMR90 and HBE: 2 x 10$^6$ cells/well) and treated with metals the next day. After removal of metal-containing media, attached cells were washed twice with warm PBS and collected by trypsination in the presence of EDTA (Gibco 15400-054 Trypsin-EDTA solution). After 2 washes with ice-cold PBS, cells were resuspended in 50 μl of deionized water followed by mixing with the equal volume of 10% nitric acid. Samples were then frozen at –80°C.
thawed at 37°C, heated for 60 min at 50°C, and placed on ice for 30 min. Metal-containing supernatants were collected after centrifugation at 10,000 x g for 10 min at 4°C. The extracts were diluted to 2% nitric acid for metal analysis by GF-AAS (AAnalyst600 Atomic Absorption Spectrometer, Perkin-Elmer). Cellular precipitates formed after metal extraction were washed twice with ice-cold 5% nitric acid, centrifuged at 10,000 x g for 5 min at 4°C, and dissolved in 100 µl of 0.5M NaOH at 37°C for 30 min. The protein amount in the solubilized cellular material was used for normalization of metal uptake. In analytical recovery experiments, 0.5 and 2.5 nmol Co and Ni were added to 0.5 million H460 cells suspended in 50 µl water immediately followed by the standard procedure for metal extraction by nitric acid. Control samples containing no cells were processed in parallel. To assess the effectiveness of the nitric acid extraction of metals in uptake experiments, H460 cells were treated with 200µM Co(II) and 400µM Ni(II) for 6h, processed by the standard protocol, and then metals were measured in both nitric acid extracts and nitric acid-insoluble material solubilized in 0.5M NaOH. The effectiveness of extraction was expressed as a percentage of total cellular metals that were present in the nitric acid–soluble fractions.

Fluorescence-activated cell sorting. After removal of metal-containing media, cells were labeled with 10µM 5-ethyl-1-2-deoxyuridine (EdU) for 30 min. Cells were washed with PBS, collected by trypsinization, and fixed overnight in 80% ethanol at 4°C. PBS-washed cells were permeabilized with 0.5% Triton X-100 in PBS for 30 min at room temperature. After a wash with 2 ml PBS, cells were resuspended in 150 µl EdU Click-IT reaction cocktail from Invitrogen (Click-IT EdU-Alexa Fluor 488 Flow Cytometry Assay) and incubated for 30 min at room temperature in the dark. Cells were washed once with 2 ml PBS, resuspended in 500 µl PBS containing 4 µg/ml propidium iodide, and incubated for 30 min at room temperature protected from light. Cells were washed once with 2 ml PBS and resuspended in 0.5 ml PBS for flow cytometry analysis (FACSCalibur, BD Biosciences). Data were analyzed by CellQuest Pro software.

Statistics. p values were calculated using 2-tailed, unpaired t test.

RESULTS

Metal Uptake and p53 Activation by Co(II) and Ni(II) in H460 Cells

Lung is the main target of toxic effects by Co(II) and Ni(II) in occupational settings (Goodman et al., 2011; Nordberg, 1994). The NTP inhalation studies found that soluble Co(II) and insoluble Ni(II) produced lung tumors originated from epithelial cells (Bucher et al., 1999; Dunnick et al., 1995). Therefore, we chose H460 human lung cells as our primary cell model. These cells retained epithelial morphology and showed a normal activation of the stress-responsive transcription factor p53 by Ni(II) (Wong et al., 2013) and other lung toxicants and carcinogens (Reynolds and Zhitkovich, 2007; Wong et al., 2012; Zhang et al., 2006). Co and Ni are high-dose carcinogens producing respiratory tumors at doses that cause death of significant amounts of lung cells (Bucher et al., 1999; Dunnick et al., 1995; Goodman et al., 2011). It has been calculated that millimolar intracellular concentrations of these metals are necessary for cell transformation (Goodman et al., 2011; Muñoz and Costa, 2012). In our experiments, we tested a range of Ni and Co concentrations that included weakly (< 25% decrease in long-term cell viability) and strongly (> 50% decrease in long-term viability) cytotoxic doses. Strongly cytotoxic doses provide sufficiently robust responses that are necessary for reliable assessments of partial effects of genetic or pharmacological manipulations.

To accurately assess a relative toxicity of Ni(II) and Co(II), it was important to determine cellular accumulation of both metals under our exposure conditions. We adapted a nitric acid–based extraction procedure that was developed for measurements of cellular chromium (Messer et al., 2006). GF-AAS analyses of extracts from “spiked” H460 cells found higher than 90% analytical recovery for both metals (Fig. 1A). The effectiveness of the nitric acid protocol in extraction of both Co and Ni from metal-exposed H460 cells exceeded 95% (Fig. 1B). Uptake measurements in H460 cells treated with metals for 6h showed a much higher accumulation of Co in comparison to Ni (Fig. 1C). On average, cells accumulated 7.7 times more Co. To compare the abilities of Co(II) and Ni(II) in activation of the transcriptional factor p53, we examined its protein and Ser15 phosphorylation levels. Activation of several stress-responsive kinases can lead to phosphorylation of Ser15, which usually correlates with increased protein stability of p53 (Kruse and Gu, 2009; Toledo and Wahl, 2006). We found that Co(II) caused only marginal increases in p53 protein and its Ser15 phosphorylation at the highest dose, whereas Ni(II) produced a very clear upregulation of both p53 readouts (Fig. 1D). At maximal doses that were tested, p53 levels were increased 4.4-fold by Ni, but only 1.4-fold by Co(II), despite its 4.4 times higher cellular accumulation. Next, we investigated how H460 cells respond to metals after 24-h-long exposures. A much greater uptake of Co, averaging 8.6-fold from the slopes of dose-response curves, was detected again (Fig. 1E). Compared with 6-h treatments, cellular levels of Ni and Co were, respectively, 1.6 and 1.8 times higher, demonstrating a significant additional influx of both metals. Activation of p53 by Co(II) after 24-h exposures was clearly evident, with particularly strong increases in its protein and Ser15 phosphorylation levels at 400µM dose (Fig. 1F). Induction of the p53 target, the CDK inhibitor p21 (CDKN1A), showed similar Co(II) dose responses by Western blots and quantitative reverse transcriptase PCR (qRT-PCR; Figs. 1F and 1G). A complete loss of p21 messenger RNA (mRNA) upregulation by Co(II) in cells with a stable p53 knockdown by shRNA confirmed the p53 dependence of p21 induction (Fig. 1G). Expression of another target gene MDM2 was induced by Co(II) in control but not in p53-depleted cells, provided further evidence for p53 activation (Fig. 1G). When compared on the basis of their intracellular doses, Co(II) was still a weaker activator of the p53 pathway than Ni(II). For example, despite their 3.3 times higher metal accumulation, 200µM Co(II)-treated cells had lower p53 and p21 levels than 300µM Ni(II) samples (Fig. 1F). Striking differences between 2 metals were also found in modulation of the p53 transactivation inhibitor MDM4 (HDMX). Ni(II)-exposed cells showed severely depressed levels of MDM4 protein with less than 10% remaining amount for the 300µM treatment, whereas even the highest concentration of 400µM Co(II)
delivering 10 times more intracellular metal had no effect on MDM4 protein (Fig. 1F). Upregulation of p21 and MDM2 mRNA levels by 400 μM Ni(II) was more than 2 times lower compared with 400 μM Co(II) (Fig. 1H vs Fig. 1G), which is consistent with stronger p53 stabilization and Ser15 phosphorylation in Western blots for this dose of Co(II) (Fig. 1F).

However, considering almost 8 times higher cellular doses of Co(II) (Fig. 1E), uptake-normalized induction of MDM2 and p21 by Ni(II) was about 4 times stronger. Similar to Co(II), upregulation of MDM2 and p21 gene expression by Ni(II) was suppressed by p53 depletion (Fig. 1H). Overall, our results showed that in comparison to Ni(II), Co(II) displayed a much higher accumulation in H460 human lung cells, but its activation of the stress-responsive transcription factor p53 was delayed and weaker when normalized for uptake.

### Delayed Cytotoxicity by Co(II)

A very strong upregulation of the transcriptional factor p53 by 24-h treatment with 400 μM Co(II) prompted us to investigate the presence of apoptotic responses. Surprisingly, we found no increase in activity of the 2 main executioner caspasases, caspase-3 and caspase-7, in H460 cells treated with 400 μM and higher Co(II) concentrations (Fig. 2A). In contrast, Ni(II)-treated cells displayed a very strong upregulation of caspase activity. The addition of EDTA prior to assay reagents did not significantly change caspase activity readings, arguing against the possibility that negative findings for Co(II) were caused by its interference with the assay. Caspase activity results were corroborated by the absence of a cleaved (active) form of caspase-7 in Co(II)-exposed cells (Fig. 2B). Ni-treated cells contained large amounts of cleaved caspase-7, demonstrating a much greater ability of this metal to engage apoptosis. The absence of caspase activation does not exclude a possibility of other modes of cell death. Any form of cell death would decrease the number of attached cells, which can serve as a test for cell death responses. We found that 24-h treatments with Co(II) caused a dose-dependent suppression of cell proliferation, as evidenced by progressively smaller increases in the number of attached cells (Fig. 2C). However, even the highest dose of 600 μM Co(II) did not decrease the number of cells below the pretreatment level. In contrast, losses of cells were detected at 24 h posttreatment, pointing to delayed cell death responses. Assessment of LDH leakage from Co(II)-exposed cells showed no cytotoxicity immediately after 24-h treatments but provided a clear evidence of cell death at 24 h posttreatment (Fig. 2D). A downward trend for LDH readings in samples analyzed at the end of 24-h exposures raised a concern that the presence of Co(II) in media could have interfered with the LDH assay. To test this possibility, we measured activity of the kit-supplied LDH standard in the presence of 400 μM Co(II) and found no inhibitory effect (Fig. 2E). Therefore, a modest decrease in LDH activity in samples collected immediately at
the end of 24-h exposures most likely reflected lower background readings due to smaller numbers of cells (Fig. 2C). To explore potential reasons for the lack of apoptotic responses at the end of Co(II) exposures, we investigated expression of 3 main antiapoptotic proteins: BCL2, BCL-XL, and MCL1. Levels of MCL1 are particularly susceptible to regulatory changes, including via its stability controlled by the aminoterminal PEST domain (Domina et al., 2004). PEST domain is defined as a protein region that is rich in proline (P), glutamic acid (E), serine (S), and threonine (T). We found no changes in expression of BCL2, BCL-XL, and MCL1 in either Ni(II)-or Co(II)-treated cells (Fig. 2F). Antiapoptotic phosphorylation of BCL2 at Ser70 was also unaltered by both metals. Although Ni(II)-induced apoptosis in H460 cells involves a mitochondrial pathway (Wong et al., 2013), a clearly different apoptotic proficiency of Ni(II) and Co(II) does not appear to be related to the status of mitochondria-protecting antiapoptotic proteins. Both Ni(II)- and Co(II)-treated cells lacked proapoptotic phosphorylation of p53 at Ser46 (Fig. 2F), which is targeted by a DNA damage–responsive kinase HIPK2 (D’Orazi et al., 2002). The inability of hypoxia-mimicking Co(II) and Ni(II) ions to promote this modification could be related to the absence of DNA damage by these nongenotoxic metals or results from their induction of HIF-1α, which causes destabilization of HIPK2 (Nardinocchi et al., 2011).

To further understand tolerance of Co(II) by H460 cells, we examined inhibition of cell cycle progression, a response that has a lower threshold than cell death and is mechanistically distinct from cell death signaling. We found that 200μM Co(II), a dose delivering 3.3 times more cellular metal than highly apoptotic 500μM Ni(II), produced only modest decreases in the percentage of S-phase cells and the rate of DNA replication (Figs. 2G and 2H). Incubation of H460 cells with 400μM Co(II), which induced a very strong upregulation of p53 and p21 (Fig. 1F), caused a practically complete shutdown of DNA replication. Overall, our results showed that Ni(II) was a much more rapid and potent activator of apoptosis than Co(II). H460 cells loaded with high doses of Co(II) were able to continue replication and display delayed cell death responses.

Cytotoxicity of Co(II) Versus Ni(II)

Different cell viability measurements showed a delayed activation of cell death by Co(II), whereas Ni(II) caused a strong apoptotic response already at the end of 24-h exposure (Fig. 2). Viability of Ni(II)-treated H460 cells continued to decrease at 24 h postexposure and then showed a sharp recovery at 48 h postexposure (Fig. 3A). Co(II) caused a more modest decrease in cell viability at 24 h postexposure and a clear recovery trend after additional 24-h recovery (Fig. 3B). To compare...
cytotoxicity of Co(II) and Ni(II), we examined cell viability at 72h postexposure, a time interval incorporating both (1) acute and delayed phases of death and (2) 48-h period of growth recovery for both metals. We found that Co(II) was slightly more cytotoxic than Ni(II) with a 1.1-fold difference in slopes (Fig. 3C). Finally, we measured a long-term viability of Co(II)- and Ni(II)-treated H460 cells by scoring their cloning abilities. Clonogenic viability assesses the ability of cells to retain long-term proliferative activity, as scored by the presence of colonies resulting from proliferation of individual cells. Cells were treated with metals in mass cultures for 24 h, 400 cells from combined populations of attached and floating cells were seeded per 60-mm dish and then grown for 7 days to form visible colonies.

FIG. 3. Comparison of Co(II) and Ni(II) cytotoxicity in H460 cells. In all experiments, cells were treated with metals for 24h in complete growth media. Viability was assessed by the determination of the metabolic activity of cell populations using the CellTiter-Glo luminescent assay, which measured total cellular ATP levels in a sample. Changes in cell viability at different times after exposure to Ni(II) (A) and Co(II) (B). Data are means ± SD for 6 independent determinations. C, Cell viability at 72 h postexposure (means ± SD for 6 independent determinations). D, Clonogenic survival of H460 cells (means ± SD for 3 independent determinations). Clonogenic viability assesses the ability of cells to retain long-term proliferative activity, as scored by the presence of colonies resulting from proliferation of individual cells. Cells were treated with metals in mass cultures for 24 h, 400 cells from combined populations of attached and floating cells were seeded per 60-mm dish and then grown for 7 days to form visible colonies.

Role of p53 in Cytotoxicity of Co(II)

Activation of p53 has been found to play a critical role in Ni(II)-induced apoptosis in H460 cells (Wong et al., 2013), and there was clear evidence of p53 upregulation by Co(II) (Fig. 1F). Therefore, we next investigated whether p53 is important for manifestation of cytotoxic effects by Co(II) using H460 cells with p53 knockdown by shRNA. Consistent with the findings in parental H460 (Figs. 2A and 2B), both cells with control (sh-SCR) and targeting (sh-p53) shRNA contained no detectable cleaved (active) forms of caspase-3 and caspase-7 at the end of 24-h treatments with Co(II) (Fig. 4A). The same result was obtained in repeat Western blots of these samples along with positive controls (not shown). The highest dose of 500μM Co(II) generated a minor cleavage of PARP, which was eliminated by p53 knockdown. At 24h posttreatment, Co(II)-treated cells expressing control shRNA contained significant amounts of active caspase-3 and caspase-7 and showed a nearly complete PARP cleavage by 500μM Co(II) (Fig. 4B). Cells with p53 knockdown had approximately 2 times lower amounts of active caspases 3/7 and cleaved PARP. The partial dependence of caspase activation on p53 was not a result of its incomplete knockdown because p53 protein and phospho-Ser15 levels were almost undetectable (Figs. 4A and 4B). A complete loss of caspase-3 cleavage by Ni(II) in sh-p53 cells provided a functional validation of their severe p53 deficiency (Fig. 4C). The presence of p53 had no effect on cell growth inhibition by Co(II) during 24-h exposure time (Fig. 4D, top panel), which is consistent with the delayed upregulation of the p53 pathway (Fig. 1). However, knockdown of p53 diminished suppression of cell proliferation and decreased the loss of cells (cell death) during 24-h recovery in Co(II)-free media (Fig. 4D, bottom panel). In agreement with Western blotting results for cleaved caspases, p53 depletion diminished caspase 3/7 activity in Co(II)-treated cells by approximately 2-fold (Fig. 4E). Finally, we determined that p53 deficiency increased cell death resistance to Co(II), as evidenced by lower levels of LDH leakage and a higher clonogenic survival (Figs. 4F and 4G). Overall, our results showed that upregulation of p53 was responsible for 50%–60% of caspase 3/7 activation and cell death by Co(II). Caspase-3 activation by Ni(II) was completely eliminated by p53 knockdown. At 24h posttreatment, Co(II)-treated cells expressing control shRNA contained significant amounts of active caspase-3 and caspase-7 and showed a nearly complete PARP cleavage by 500μM Co(II) (Fig. 4B). Cells with p53 knockdown had approximately 2 times lower amounts of active caspases 3/7 and cleaved PARP. The partial dependence of caspase activation on p53 was not a result of its incomplete knockdown because p53 protein and phospho-Ser15 levels were almost undetectable (Figs. 4A and 4B). A complete loss of caspase-3 cleavage by Ni(II) in sh-p53 cells provided a functional validation of their severe p53 deficiency (Fig. 4C). The presence of p53 had no effect on cell growth inhibition by Co(II) during 24-h exposure time (Fig. 4D, top panel), which is consistent with the delayed upregulation of the p53 pathway (Fig. 1). However, knockdown of p53 diminished suppression of cell proliferation and decreased the loss of cells (cell death) during 24-h recovery in Co(II)-free media (Fig. 4D, bottom panel). In agreement with Western blotting results for cleaved caspases, p53 depletion diminished caspase 3/7 activity in Co(II)-treated cells by approximately 2-fold (Fig. 4E). Finally, we determined that p53 deficiency increased cell death resistance to Co(II), as evidenced by lower levels of LDH leakage and a higher clonogenic survival (Figs. 4F and 4G). Overall, our results showed that upregulation of p53 was responsible for 50%–60% of caspase 3/7 activation and cell death by Co(II). Caspase-3 activation by Ni(II) was completely dependent on p53.

To further understand mechanisms of Co(II)-induced apoptosis, we measured mRNA levels for 12 major pro- and antiapoptotic proteins. Stress-induced apoptosis can be executed via mitochondrial (intrinsic) or death receptor–mediated (extrinsic) mechanism. The key event in the intrinsic form of apoptosis is release of proapoptotic factors from mitochondria by oligomers of BAX and BAK that form pores in the outer mitochondrial membrane. Oligomerization of BAX/BAK is promoted by BH3-only domain proteins (PUMA, NOXA, BID, and BIM) and suppressed by BH4-containing proteins such as BCL2 and MCL1 (Green and Kroemer, 2009). Mitochondria-dependent apoptosis engages the initiator caspase-9. We found that irrespective of their p53 status, Co(II)-treated cells showed no
significant changes in gene expression for caspase-9, as well as its adaptor protein APAF1 and inhibitor BIRC5 (survivin, IAP4) (Fig. 5A). Messenger RNA levels for antiapoptotic MCL1 and BCL2 were also similar in both control and p53-depleted cells (Fig. 5B). Examination of gene expression for BH3-only activators of BAX/BAK oligomerization found that Co(II) caused a strong upregulation of NOXA (PMAIP1) and PUMA (BBC3) mRNA and this response was suppressed in sh-p53 cells by approximately 60% (Fig. 5C). Co(II)-treated cells also showed a modest, p53-independent increase in BIM expression but a slight decrease in BID. There was no effect of Co(II) on mRNA amounts of apoptotic BAX although its expression was moderately lower in p53 knockdown cells. Overall, our qRT-PCR analyses found that upregulation of NOXA and PUMA was the main proapoptotic gene expression response in Co(II)-treated cells. Induction of these genes was partially dependent on p53.

Mechanisms of Cell Death by Co(II)

Leakage of cellular LDH, which we observed in Co(II)-treated cells (Fig. 2D), can reflect the presence of late apoptotic cells or induction of necrosis. Co(II) is a prooxidant metal (Kasprzak et al., 2003), and oxidative stress is known to promote a necrotic death via opening of the mitochondrial permeability transition pore and subsequent collapse of the H⁺ gradient and loss of ATP production. Importantly, p53 protein induces this form of necrosis by binding to cyclophilin D in the inner mitochondrial membrane (Vaseva et al., 2012). Thus, it was plausible that the observed above inhibition of LDH leakage in sh-p53 H460 cells (Fig. 4F) was caused by a suppression of p53-dependent necrosis. Using different pharmacological agents, we next explored a role of necrotic and caspase-dependent and -independent processes in Co(II) cytotoxicity. The extent of cell death was monitored by LDH leakage at 24 h postexposure, which corresponds to the peak of Co(II)-induced cytotoxicity (Figs. 3 and 4). In the initial screening, inhibitors were added only for 24-h recovery time to avoid potential effects on Co(II) uptake. We found that addition of cyclosporine A, which blocks physical interactions between p53 and cyclophilin D and oxidant-induced necrosis (Vaseva et al., 2012), had no effect on LDH release from Co(II)-treated H460 cells (Fig. 6A). LDH leakage was also unchanged in the presence of necrostatin-1, which inhibits a death receptor–associated form of necrosis involving RIP1 (RIPK1) kinase (Degterev et al., 2008). Consistent with the observed earlier activation of executioner caspases, the pancaspase inhibitor QVD-Oph suppressed LDH leakage by more than 60%. Inhibition of other cysteine proteases, such as cytosolic calpains by ALLN and lysosomal cathepsins by E64d, had no effect on the amount of extracellular LDH. Western blotting results demonstrated a complete blockage of
PARP cleavage and the production of active caspases 3/7 by QVD-Oph (Fig. 6B). The presence of ALLN and E64d did not change the levels of these apoptotic markers, excluding the role of cathepsins and calpains in the initiation of caspase activation. Next, we examined whether the incompleteness of LDH release and the inhibitory effects of QVD-Oph were caused by its presence only during 24-h recovery, which could have allowed some early apoptotic events. We found that the degree of LDH inhibition was not affected by the addition of QVD-Oph during and after Co(II) exposure versus only during postexposure time (Fig. 6C, left panel). Control experiments showed that cytotoxicity-inhibitory effects of QVD-Oph were not associated with suppressive effects on p53 accumulation (Fig. 6C, right panel). Finally, we examined a relative role of p53- and caspase-dependent processes in cell death by Co(II) (Fig. 6D). From the slopes of dose-response curves, we calculated that p53 depletion inhibited LDH leakage by about 55% and QVD-Oph by approximately 75% in both control and sh-p53 cells. Overall, our results showed that cell death by Co(II) involved multiple mechanisms and was partially dependent on p53.

Co(II) and Ni(II) Treatments of Normal Human Lung Cells

To confirm our findings in H460 cells, we examined uptake of Ni(II) and Co(II) in IMR90 normal human lung fibroblasts and primary HBE cells. We found that both types of normal lung cells had several times higher accumulation of Co(II) than Ni(II) (Figs. 7A and 7B). Ni(II)-treated HBE cells showed a dose-dependent increase in caspase 3/7 activity at the end of 24-h exposures (Fig. 7C). In contrast, despite its much higher cellular doses, Co(II) produced no caspase activation. In agreement with findings in H460, Co(II)-treated HBE cells displayed a moderate activation of caspases at 24 h postexposure (Fig. 7D). Thus, studies with 2 types of normal human lung cells confirmed a much higher uptake of Co(II) but a delayed and weaker caspase activation compared with Ni(II).

**DISCUSSION**

Co(II)-Induced Cell Death

Our studies here showed that despite their chemical similarity, Co(II) had a much higher cellular accumulation than Ni(II), but it was a slower and weaker activator of p53 and death in H460 cells. In contrast to Ni(II), even several times higher cellular doses of Co(II) produced no changes in the levels of the p53 inhibitor MDM4, indicating activation of a more limited stress signaling network. Delayed death in Co(II)-treated cells was triggered via p53-dependent and -independent mechanisms (Fig. 8A). Approximately 55% of Co(II)-induced cell death was attributable to the p53-dependent, caspase-mediated apoptosis. This conclusion was drawn from the inhibition of caspase activity, decreases in active caspase-3 and caspase-7, and suppression of LDH release by p53 knockdown. Consistent with the above estimate, Co(II)-induced expression of 2 most highly upregulated proapoptotic genes NOXA and PUMA was reduced by 50%–60% in
HIGH TOLERANCE OF CO(II) BY HUMAN LUNG CELLS

FIG. 6. Mechanisms of death in Co(II)-treated H460 cells. Data are means ± SD for at least 3 independent determinations. Where not seen, error bars were smaller than data symbols. A, Leaked LDH from cells treated with 500 μM Co(II) for 24 h in complete media and then incubated for 24 h in metal-free media without serum in the presence of inhibitors: CSA—cyclosporin A (4 μM), NS1—necrostatin-1 (10 μM), QVD (QVD-Oph, 20 μM), ALLN (5 μM), and E64d (10 μM). **p < 0.01 relative to Co(II)-treated cells without inhibitors. B, Formation of active caspases 3/7 and PARP cleavage in cells treated with Co(II) and inhibitors as in panel A. C, Inhibition of LDH leakage by 50 μM QVD-Oph present during and after Co(II) treatment (QVD-all time) and added only for 24-h recovery (QVD-recovery). Right panel: p53 protein levels in H460 cells treated with QVD-Oph during 24 h Co(II) exposure and 24 h postexposure. D, LDH leakage from cells with control (sh-SCR) and p53-targeting (sh-p53) shRNA in the presence or absence of 50 μM QVD-Oph (QVD, present during and after exposure). LDH leakage was measured after 24-h recovery in Co(II)-free media.

p53-depleted cells. Unlike Ni(II) (Wong et al., 2013), loss of p53 did not cause derepression of antiapoptotic MCL1 by Co(II), which together with only a partial effect on PUMA induction and caspase-3 activation was probably responsible for the overall less potent prosurvival effect of p53 deficiency for Co(II). A caspase-mediated but p53-independent apoptosis accounted for about 20% of Co(II)-induced cell death, and the remaining 25% death was both p53 and caspase independent. In contrast to engagement of multiple cell death mechanisms by Co(II), our recent studies showed that Ni(II)-induced death in H460 cells was entirely caspase dependent, and p53 depletion completely abrogated caspase-3 activation (Wong et al., 2013). Although Co(II) is a redox-active metal and oxidative stress is known to activate necrosis by p53 binding to mitochondrial cyclophilin D (Vaseva et al., 2012), we did not detect any survival benefit by inhibiting this and alternative RIP1-dependent (Vandenabeele et al., 2010) necrotic pathways. Our findings on a similar residual cytotoxicity in control and p53-depleted cells in the presence of the pancaspase inhibitor QVD-Oph further argue against any pronecrotic effect of p53 in Co(II)-treated cells. Because p53 deficiency did not completely suppress NOXA and PUMA induction, it is possible that the continuing mitochondrial release of cytotoxic proteins such as endonuclease G and AIF promoting apoptosis directly (Green and Kroemer, 2009) was responsible for the caspase-resistant cell death by Co(II). Alternatively, caspase-inhibited severely injured cells could become nonviable due to a loss of essential respiration factors caused by a persistent permeabilization of the outer mitochondrial membrane (Lartigue et al., 2009). Human lung cells accumulated 4–9 times more Co(II) than Ni(II), as determined by direct measurements of cellular metals in our studies. The indirectly measured rates of Co(II) influx by the human divalent metal transporter DMT1 expressed in Xenopus oocytes were 4.9–7.0 times higher than for Ni(II) (Illing et al., 2012). Similar estimates for transport preference of Co(II) over Ni(II) points to the importance of DMT1 in metal uptake by lung cells. Metal selectivity for Co(II) and Ni(II) entry via other transporters such as Ca(2+) channels is currently unknown (Goodman et al., 2011; Simonsen et al., 2012).

Our findings on a high tolerance of Co(II) by human lung cells are consistent with a lack of respiratory toxicity in individuals displaying injury to other tissues caused by elevated levels of cobalt in systemic circulation (Gilbert et al., 2013;
a larger assortment of ligand arrangements, and its common square planar structure permits a peptide-specific binding and protein cleavage, which is not observed for Co(II) (Bal et al., 2000).

Implications for Ni(II) and Co(II) Carcinogenicity

The intracellular release of Ni(II) ions is generally believed to be responsible for cancer causation by insoluble Ni compounds, although epidemiological data for carcinogenicity of soluble Ni(II) are largely negative (Goodman et al., 2011; Muñoz and Costa, 2012; Salnikow and Zhitkovich, 2008). Soluble Ni(II) (nickel sulfate) was also noncarcinogenic by inhalation in the 2-year rodent carcinogenicity assay (Dunnick et al., 1995). In contrast, inhalation exposures to the same doses of soluble Co(II) (cobalt sulfate) caused lung tumors in both mice and rats (Bucher et al., 1999). Co(II) is clastogenic in vivo and in cultured cells, and this genotoxic activity has been attributed to oxidative damage (De Boeck et al., 2003). Co(II) is generally more redox active than Ni(II), and cells treated with Co(II) showed higher levels of oxidative stress (Kasprzak et al., 2003; Van den Broeke et al., 1998). A high tolerance of intracellular Co(II) by lung cells, including resistance of Co(II) to apoptosis, allows their propagation despite the presence of oxidative damage, which should promote accumulation of critical amounts of genetic rearrangements and stimulate transformation of surviving cells (Fig. 8B). A lower threshold and more rapid activation of apoptosis by soluble Ni(II) helps eliminate damaged cells prior to their acquiring of procarcinogenic genetic or epigenetic changes. The carcinogenic ability of insoluble Ni compounds is frequently attributed to the intracellular delivery of massive amounts of Ni(II) ions upon dissolution of micron-sized particles. Our data on high cytotoxicity of Ni(II) ions would argue that in addition to their gradual release from slowly dissolving particles, a major fraction of Ni(II) leaking from endosomes into cytosol and nuclear compartments could be bound by ligands, blunting cytotoxicity of Ni(II) ions but retaining or enhancing their carcinogenic properties. Ni(II) binding to histidine and other small molecules is known to enhance its oxidative chemistry (Kasprzak et al., 2003), but this complexation should diminish reactivity of Ni with proteins, limiting the overall cellular injury and permitting survival.

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