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Acetaminophen and NAPQI are toxic to auditory cells via oxidative and endoplasmic reticulum stress-dependent pathways

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ABSTRACT

Pain relievers containing N-acetyl-para-aminophenol, also called APAP, acetaminophen or paracetamol, in combination with opioid narcotics are top-selling pharmaceuticals in the U.S. Individuals who abuse these drugs for as little as sixty days can develop tinnitus and progressive bilateral sensorineural hearing loss. Recently published studies indicate that APAP and its metabolic product N-acetyl-p-benzoquinoneimine (NAPQI) are the primary ototoxic agents in this type of pain relievers. However, the mechanisms underlying the deleterious effects of these drugs on auditory cells remain to be fully characterized. In this study, we report cellular, genomic, and proteomic experiments revealing that cytotoxicity by APAP and NAPQI involves two different pathways in Immortomouse/C212-derived HEI-OC1 cells, implicating ROS overproduction, alterations in ER morphology, redistribution of intra-cisternal chaperones, activation of the eIF2α-CHOP pathway, as well as changes in ER stress and protein folding response markers. Thus, both oxidative and ER stress are part of the cellular and molecular mechanisms that contribute to the cytotoxic effects of APAP and NAPQI in these cells. We suggest that these in vitro findings should be taken into consideration when designing pharmacological strategies aimed at preventing the toxic effects of these drugs on the auditory system.

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1. Introduction

APAP is a pro-drug metabolized by the cytochrome P450 2E1 isozyme (CYP2E1) to an electrophilic metabolite called N-acetyl-p-benzoquinoneimine (NAPQI). APAP hepatotoxicity has been known and investigated for over 50 years, and experiments performed in the 1970s suggested it was actually associated with covalent binding of NAPQI to tissue macromolecules (Bromer and Black, 2003) (and references therein). Studies on the metabolic processing of these molecules revealed that only a small fraction (5%–10%) of a therapeutic dose of APAP is converted by CYP2E1 into NAPQI in humans, whereas the other 90% is excreted as conjugates of glucuronide and sulfate (Zimmerman, 1998). In addition, inside the cells NAPQI is usually inactivated by glutathione, though a plethora of studies have suggested that acute overdose may overwhelm glutathione stores and result in oxidative toxic effects to the liver (Hersh et al., 2007). Other studies, however, have provided plausible evidence that APAP, but not NAPQI, can kill hepatocytes through oxidative stress (Harman et al., 1991). Moreover, recent results suggested that glutathione depletion itself would be insufficient to provoke hepatocyte apoptosis suggesting that an APAP-induced intraluminal redox imbalance within the endoplasmic reticulum as the actual pro-apoptotic event (Nagy et al., 2007, 2009).

In contrast to liver injury, APAP ototoxicity has been only recently recognized, eliciting significant attention by both the scientific and popular press (Blakley and Schilling, 2008; Curhan et al., 2010, 2012; Friedman et al., 2000; Ho et al., 2007; Oh et al., 2000;
Differences were evident at 48 h (APAP than NAPQI exposed to APAP and NAPQI at both time points. Although APAP effect was stronger this drug daily, an amount significant higher than the known hepatotoxic doses (Yorgason et al., 2010). Thus, it is likely that ototoxicity resulting from acute APAP overdoses was masked because individuals were dying from liver failure before developing hearing loss. In contrast, protection against APAP hepatotoxicity in long-term abusers, which could potentially be provided either by hydrocodone or an adaptive response of the liver to increased doses of APAP (Bromer and Black, 2003), would avoid premature death by liver failure and give enough time for ototoxicity to develop. After the initial studies reporting hearing loss caused by the abuse of analgesics containing APAP and hydrocodone, most research on the mechanisms underlying ototoxicity has focused on hydrocodone and/or its metabolite hydromorphone. However, studies in our laboratory have provided convincing evidence that APAP, but not hydrocodone, is the primary ototoxic agent, though the mechanisms of its deleterious effects remains unknown (Yorgason et al., 2010).

In a previous study we demonstrated, using neonatal mouse cochlear cultures and cells derived from Immortomouséd organ of Corti (HEI-OC1), that outer hair cells (OHCs) were the major target of APAP toxicity in the organ of Corti, and that HEI-OC1 cells responded to APAP just like actual OHCs (Yorgason et al., 2010). Thus, the current study has been designed to elucidate the mechanisms of APAP and NAPQI cytotoxicity in auditory cells, using the HEI-OC1 cell line as an in vitro model. Accordingly, we have performed experiments aimed at defining changes in gene expression and protein levels in HEI-OC1 cells elicited by exposure to APAP or NAPQI. The results of these studies unravel that the toxic effects of APAP and NAPQI in HEI-OC1 cells follow two different mechanism involving the induction of both oxidative and endoplasmic reticulum (ER) stress. In addition, we provide markers for this phenomenon, which can be used in the future to design strategies to monitor the occurrence and severity of this toxic response. Altogether, these studies extend our current understanding of the mechanisms responsible for the cytotoxic effects caused by the abuse of this commonly used analgesic and, although the use of HEI-OC1 cells instead of actual cochlear hair cells limits the extrapolation of our results to the organ of Corti, we believe that these findings should be taken into consideration when designing pharmacological strategies aimed at preventing the toxic effects of these drugs on the auditory system.

2. Materials and methods

2.1. Dose-response studies

Immortomouse-derived HEI-OC1 cells were grown in plastic cell culture dishes at permissive conditions, 33 °C, 5% CO2 in DMEM supplemented with 10% FBS, as previously described (Kalinec et al., 2003). HEI-OC1 cells were first trypsinized, counted and concentration adjusted to 5.0 × 10^3 cells/mL using Cellometer Auto T4 (Nexelcom Bioscience, Lawrence, MA). Next, cells were transferred to 48-well flat bottom plates (250 μL each well), incubated again at 33 °C for 4 h, and then some of them exposed to differing concentrations of APAP (0.3, 0.6, 1.5, 3.75, 7.5 mg/mL) and NAPQI (30, 60, 150, 375, 750 μg/mL) (Sigma–Aldrich, Saint Louis, MO) whereas others were not exposed to any drug (Control). Direct cell count was performed at different time points (0, 24, and 48 h) using Celligo, an adherent cell cytometer (Cyntellect Inc., San Diego, CA).

2.2. MTT Assay

HEI-OC1 cells growing at permissive conditions were collected and concentration adjusted to 5.0 × 10^3 cells/mL using Cellometer Auto T4 as described before. Next, cells were transferred to 96-well flat bottom plates (100 μL each well), incubated at 33 °C overnight.
and then exposed to 10 μM S600125 (JNK inhibitor, Sigma–Aldrich), 5, 50, and 100 μM AEBSF (4-(2-Aminoethyl) benzene-sulfonfonyl fluoride, inhibitor of the ATF6-mediated pathway, Sigma–Aldrich), or no drugs (Control) for an additional 2 h period. After this second incubation, cells were exposed to 1.5 mg/mL APAP, or 150 μg/mL NAPQI, and incubated for either 24 h or 48 h at 33 °C. At these time points, the TACS® MTT Cell Proliferation Assay (Treivigen, Gaithersburg, MD) was used following manufacturer’s protocol to investigate cell viability. Absorbance was measured with Spectra Max 5 Plate Reader with Soft Max Pro 5.2 Software (Molecular Devices LLC, Sunnyvale, CA) and average OD in Control cells was taken as 100% of viability.

2.3. ROS determinations

To detect endogenous reactive oxygen species (ROS) production, 1 × 10⁵ HEI-OC1 cells/mL, both without treatment (Control) and treated for 1 min, 5 min, 10 min, 15 min, 30 min, 1 h, 6 h, 12 h, 24 h, and 48 h with APAP and NAPQI and then washed, were incubated with 5 μM 2,7-dichlorodihydrofluorescein diacetate acetyl ester (DCFH-DA, Sigma–Aldrich) for 15 min at 37 °C in the dark. In the presence of intracellular ROS, nonfluorescent membrane-permeable DCFH-DA is first deacetylated by cellular esterases and rapidly oxidized to impermeable and highly fluorescent 2,7-dichlorodihydrofluorescein (DCF). ROS production was thus monitored by measuring DCF emission at 530 nm with excitation at 485 nm using the Spectra Max 5 nm with excitation at 485 nm using the Spectra Max 5 Plate Reader with Soft Max Pro 5.2 Software.

2.4. Western blotting

Cells were lysed at 4 °C in a 50 mM Tris buffer solution (pH 7.4) containing 1% Nonidet P-40, 2 mM EDTA, 100 mM NaCl, 1 mM vanadate, 10 μM PMSF, 2 μM 10 mg/mL leupeptin, and 2 μM 10 mg/mL aprotinin. Samples were mixed with loading buffer (non-reducing lane marker (Thermo Scientific/Pierce, Rockford, IL) plus 10μM DTT), heated at 95 °C for 5 min, analyzed by SDS/PAGE gels (30 μg of protein per lane), transferred to nitrocellulose membranes, and incubated with primary antibodies at 1:100 to 1:500 dilutions and HRP-linked secondary antibodies at 1:1500 to monitor by measuring DCF emission at 530 nm with excitation at 485 nm using the Spectra Max 5 Plate Reader with Soft Max Pro 5.2 Software.

2.8. Transmission electron microscopy of HEI-OC1 cells

HEI-OC1 cells were fixed in 4% paraformaldehyde for 1 h at room temperature, washed with PBS 3 times for 5 min each, incubated 10 min with 0.001% Triton X-100 in PBS, and then in blocking solution (10% fish gelatin, Norland Inc., Cranbury, NJ) for 30 min. All the primary antibodies (listed in Section 2.4) were used at 1:100 to 1:500 dilution in PBS in overnight incubations at 4 °C. Rhodamine phalloidin from Molecular Probes-Invitrogen (Eugene, OR, USA), and DAPI from Sigma were used to stain actin and cell nuclei, respectively, and Alexa 488 (anti-goat and anti-rabbit) and Alexa 546 (anti-rabbit) from Molecular Probes–Invitrogen were used as secondary antibodies at 1:200 dilution. Samples were observed with a TCS-SP5 Broadband Spectra laser confocal microscope with a 63x (NA = 1.2) objective (Leica Microsystems, Deerfield, IL, USA). Images were cropped, resized, and brightness and contrast over the whole image adjusted where necessary, using Photoshop (Adobe Software).

2.9. Transmission electron microscopy of HEI-OC1 cells

Cells were washed once with 100 mM sodium cacodylate buffer (Electron Microscopy Sciences, EMS, Hatfield PA), and then fixed in a solution of 2.5% glutaraldehyde in 100 mM sodium cacodylate buffer (EMS; pH 7.2). Fixed cells were centrifuged for 5 min and the pellet immersed in fresh fixative (2.5% glutaraldehyde in 100 mM sodium cacodylate buffer). After 2 h, the fixative was removed and pellet was left in 100 mM sodium cacodylate buffer overnight. The following day the buffer was discarded and the pellet washed with double-distilled H2O (ddH2O). The pellet was then incubated at room temperature in a 1% solution of osmium tetroxide (EMS) for 35 min, washed with ddH2O and dehydrated using 5 different concentrations (30, 50, 70, 95, and 100%) of ethanol (Pharmco-AAPER, Brookfield, CT) in an ascending order. The dehydrated pellet was then immersed in propylene oxide (EMS) and later left in a mixture of epoxy resin (EMBed-812, EMS) and propylene oxide (1:1) overnight. The next day, the 1:1 mixture of epoxy resin and propylene oxide was replaced with a 3:1 mixture. After 6 h, the 3:1 mixture of epoxy resin and propylene oxide was exchanged with only epoxy resin, and the samples were left in a rotating incubator overnight. Afterward, fresh epoxy resin plus an accelerator (BDMA, EMS) were added to the tubes containing the cells, and they were
incubated for 1 h at room temperature. Finally, the cells along with the mixture of epoxy resin and accelerator were poured into plastic embedding molds and incubated at 60 °C overnight. When blocks containing the cells were hardened, 70 nm sections were obtained using an EM UC7 ultra-microtome (Leica Microsystems, Buffalo Grove, IL) with a diamond knife (Diatome Ultra 45°, Fort Washington, PA). Sections were picked up with grids (EMS) and stained with 3% uranyl acetate for 4 min (SPI Supplies, West Chester, PA) and lead citrate for 30 s (EMS). Samples were examined with a Tecnai G2 20 Twin transmission electron microscope (FEI, Hillsboro, OR) at 80 kv.

2.9. ATF4 knockdown

shRNA Lentiviral Particles, shRNA-ATF4 (sc35112VV) and shRNA Negative Control (sc-108080) were obtained from Santa Cruz Biotechnology (Santa Cruz, CA). HEI-OC1 cells were cultured in 12-well plates for 24 h prior to viral infection. Next day, with the cells approximately 50% confluent, the media was removed from place wells and replaced with 1 mL of complete medium with Polybrene® (sc-134220) at a final concentration of 5 μg/mL. Cells were infected by adding the shRNA Lentiviral Particles, shRNA-ATF4 and shRNA Negative Control to the culture and incubating for 24 h. Next, the culture medium was removed and replaced with 1 mL of complete medium (without Polybrene®) and the cells were again incubated overnight. Stable clones expressing the shRNAs were selected via Puromycin dihydrochloride (sc-108071) selection at 3 μg/mL. Finally, cells were exposed to APAP and NAPQI for 24 h or 48 h, and then collected and processed for WB, PCR, MTT and confocal microscopy.

2.10. Statistical analysis

Statistical analysis of the data, including One-way and Two-way ANOVA, was performed using JMP 9 software (SAS Institute, Cary, NC) and p < 0.05 as the criterion for statistical significance.

3. Results

3.1. Oxidative stress contributes to the cytotoxic effects of APAP, but not NAPQI, on HEI-OC1 cells

To examine the cytotoxic effects of APAP and NAPQI, we treated HEI-OC1 cells with six different doses of these drugs and investigated dose-dependence by software-performed continuous counting of number of cells in culture plates. All NAPQI doses used in our studies were set at 10% of those of APAP, since it is well established that no more than 5—10% of APAP is metabolized into NAPQI (Bromer and Black, 2003; Mikus and Weiss, 2005). As shown in Fig. 1A, both APAP and NAPQI significantly decreased the number of cells in all the experimental conditions investigated. Exposure to APAP resulted in more pronounced effects than those observed with NAPQI, revealing that the toxic effects of this drug on cells from auditory lineage occur, at least in part, independently of its metabolite. The toxicity curves guided subsequent studies, which, accordingly, were performed using 1.5 mg/mL APAP and 0.15 mg/mL NAPQI doses that consistently reduced the number of HEI-OC1 cells. Using these doses, we obtained similar results using a second, independent method for measuring cell viability, namely the MTT assay. As shown in Fig. 1B, both APAP and NAPQI significantly decreased viability of HEI-OC1 cells at 24 and 48 h respect to Control condition, but while APAP’s toxic effect was stronger than NAPQI’s effect at 24 h, no differences were observed at 48 h.

It has been suggested that the toxic effects of aminoglycoside antibiotics on auditory cells would be mediated by overproduction of ROS that cannot be neutralized by the natural anti-oxidant mechanisms (Forge and Schacht, 2000). Moreover, some of the work involving aminoglycosides describe ROS increasing within minutes to hours of exposure of chick hair cells to gentamicin in vitro (Hirose et al., 1997). Thus, we investigated whether the cytotoxic effects of APAP and NAPQI could also be associated with oxidative stress at time points ranging from 1 min to 48 h. As shown in Fig. 2, APAP significantly decreased ROS levels at short exposure times and then increased them, reaching Control values at the 6 h time-point and up to 4-fold higher for longer treatments. ROS levels in NAPQI-treated cells, in contrast, were always lower than in Control cells. These results suggest that APAP cytotoxic effects, but not those from NAPQI, would be mediated by oxidative injury.
3.2. Contribution of endoplasmic reticulum stress to the cytotoxic effects of APAP and NAPQI on HEI-OC1 cells

Subsequently, we tested whether HEI-OC1 cells treated with APAP or NAPQI would undergo endoplasmic reticulum stress (ERS). For this purpose, we initially performed confocal microscopy and Western blotting using antibodies against GRP78 (glucose regulated protein of mw 78, a.k.a. BiP) and CHOP (CCAAT/enhancer binding protein homologous transcription factor, also called GADD153), surrogate marker proteins for this phenomenon. GRP78 function as a major ER-enriched chaperone for facilitating protein folding, and can bind to misfolded proteins and unassembled complexes. Its expression is induced by ERS, but once the stress is removed GRP78 is post-transcriptionally modified into a biologically inactive form. CHOP, on the other hand, is induced during ERS to integrate and amplify the death pathway (Schonthal, 2009; Woehlbier and Hetz, 2011).

We investigated GRP78 and CHOP expression at 12, 24 and 48 h, since our previous experiments indicate extreme values of ROS at these time points. We found that Control (untreated) cells showed

Fig. 3. The cytotoxic effects of APAP and NAPQI on HEI-OC1 cells involves ERS. Confocal microscopy using antibodies against ERS markers GRP78 (green, A–I) and CHOP (green, J–R) at 12, 24 and 48 h. In all panels red color corresponds to rhodamine phalloidin and blue to DAPI. In Control and APAP-treated cells GRP78 immunolabeling at 12 and 24 h was mostly perinuclear and homogeneous, although at 12 h reactivity was more extended in APAP-treated cells (B, arrowheads); in NAPQI-treated cells immunolabeling was coarse and cluster-like, with a perinuclear distribution at 24 h (F, arrows), but also present in cellular projections at 12 h (C, arrowheads). At 48 h it was still perinuclear in Control cells (G, arrows), but it was also detected in filopodia and other cellular projections in APAP- and NAPQI-treated cells (H–I, arrowheads). CHOP immunoreactivity was weaker than GRP78. At 12 h a diffuse labeling was observed in the perinuclear region of Control cells (J, arrow), whereas in APAP-treated cells the nucleus of some cells were clearly immunoreactive (K, arrowhead) but not others (K, arrow); a faint labeling was observed in the nucleus and cytoplasm of in NAPQI-treated cells (L, arrow and arrowhead). At 24 h immunolabeling was more evident in all the conditions, with well defined spots of immunoreactivity in the perinuclear region (arrows) and cell periphery (arrowheads) (panels M–O). At 48 h cells' labeling was more similar to that at 12 h than at 24 h, with a faint cytoplasmic labeling in Control cells (P, arrow), nuclear reactivity in some APAP-treated cells (Q, arrow), and NAPQI-treated cells showing CHOP immunoreactivity both in the perinuclear region (arrow) and cell periphery (arrowhead) (panel R).
GRP78 immunolabeling mostly at the perinuclear region of the cell (Fig. 3 A–D). Interestingly, we found that the general pattern of GRP78 labeling was more similar at 12 and 48 h (Fig. 3 A–C and G–I) than at 24 h (Fig. 3 D–F), with immunoreactivity moving beyond the perinuclear region to the periphery at 12 and 48 h. Distinct cytoplasmic clusters highly reminiscent of ER fragmentation and vacuolization characterized NAPQI immunoreactivity and, as in the other conditions, labeling was mostly perinuclear at 24 h and moved to the cell periphery at 12 and 48 h (Fig. 3 C,F,I). By WB analysis, we found that both APAP and NAPQI actually induced an increase in GRP78 expression, but APAP-induced peak occurred at 48 h whereas NAPQI-induced peak took place 24 h after exposure (Fig. 4 A).

CHOP expression was also affected by APAP and NAPQI. CHOP immunolabeling was not strong either in Control or treated HEI-OC1 cells, but this could be associated to the particular antibody used in our study (Fig. 3 J–R). At 24 h CHOP reactivity was mostly perinuclear in Control cells, whereas it was found in all the cytoplasm in APAP or NAPQI-treated cells (Fig. 3 M–O). At 12 and 48 h CHOP immunoreactivity was lower than at 24 h, weak in the cytoplasm of Control cells (Fig. 3 J and P, arrows), present in the nucleus of only some APAP-treated cells (Fig. 3 K and Q, arrowheads), and the nucleus and cytoplasm of NAPQI-treated cells (Fig. 3 L and R, arrows and arrowheads). WB studies showed a significant increase in CHOP expression in cells treated with APAP, NAPQI and tunicamycin (positive Control) at 12 h (Fig. 4B). However, although real, this increase could be magnified by the very low expression in Control cells. APAP did not induce any effect in HEI-OC1 cells treated for 24 h, whereas a significant decrease respect to Control values was observed at 48 h. NAPQI, on the other hand, significantly decreased CHOP expression at 24 and 48 h. Curiously, CHOP expression in tunicamycin-treated cells also decreased to about 50% of Control values at 48 h (Fig. 4B). Finally, transmission electron microscopy images showed evident ER vacuolization coupled to a dramatic reduction in membrane-bound ribosomal particles in APAP and, particularly, NAPQI-treated cells but not in Control cells (Fig. 4C).

These results demonstrate that both APAP and NAPQI induce the expression of ERS markers, but that only the latter appears to form the intracellular structures that characterize this phenomenon.

### 3.3. HEI-OC1 cells treated with APAP and NAPQI express genomics and proteomic markers of both oxidative and ER stress

Next, we investigated changes in gene expression induced in HEI-OC1 cells by exposure to APAP and NAPQI for 24 and 48 h, the time-points showing qualitatively different responses in our previous experiments. For this purpose, we used pathway specific RT² Profiler PCR arrays which measure the levels of 308 different genes associated with cell death, oxidative stress, and key signaling pathways involved in cell growth, survival, and death. Using this approach, we find that 232 transcripts were significantly (more than 2-fold) up or down regulated by exposure to APAP and NAPQI (see Supplemental Table 1). By generating Venn diagrams, we defined that among these 232 genes, 32 transcripts were regulated by APAP alone, 163 by NAPQI alone, and 37 by both compounds. For example, by focusing on the ontological association of these genes, we observed that APAP up regulated caspases 2 and 3 whereas NAPQI did not change the expression of these genes but instead up regulated caspases 1, 4, 6, 7, 8, 9, 12 and 14. Similarly, expression of some glutathione-s-transferases like Gsta1, Gstp2 and Gstt1 was affected only by APAP, whereas others like Gsta1 and Gstm1 or glutathione reductase (Grs) were targeted only by NAPQI. These results provide insights on the intracellular molecules associated with the toxic effects of both APAP and NAPQI on HEI-OC1 cells.

We applied semantic-based algorithms (IPA™ software, Ingenuity Systems/Qiagen) to gain insight as how the genes that changed their expression levels in response to drug treatment were linked into single functional networks. This approach indicated that approximately 50% of the genes tested were related to ERS and
protein degradation, a result congruent with our cell biological experiments that further support a role for these interrelated pathways in the response to APAP and NAPQI. In addition, more than 80% of genes whose expression was either increased or decreased more than 2-fold by APAP and NAPQI (199 out of 232) belong to five different functional gene networks, with some of them being ascribed to more than one group. These groups include genes associated with xenobiotic metabolism, oxidative radical metabolism, ER stress, protein ubiquitination, and cell death (Fig. 5). Notably, whereas it can be expected that genes whose expression is necessary for drug metabolism, oxidative stress and cell death are differentially regulated in response to toxic drug exposure, our finding that in HEI-OC1 cells approximately 50% of the genes tested were related to ERS and protein degradation led us to hypothesize that these highly interrelated pathways would play an important role in the response. However, we were aware that this result could potentially be biased by the particular selection of the set of gene arrays chosen for this study. In addition, we thought it was important to validate the gene expression results at the proteins level. Thus, to extend our results, we decided to perform experiments using quantitative proteomics techniques, which would address both the gene selection bias and provide gene-to-protein expression validations. For this purpose, we investigated the changes in protein levels induced by 24 and 48 h exposure to APAP and NAPQI with iTRAQ techniques. These experiments identified more than 900 proteins, with 234 proteins that were significantly up or down regulated by treatment with these drugs (Supplemental Table 2). iTRAQ studies performed at Keck Biotechnology Resources Laboratory at Yale University; complete results are available at their website). IPA™-assisted ontological analyses of this data revealed that approximately 50% of these proteins groups identified by our gene expression studies (Fig. 5). When combined, both the gene and protein expression datasets indicate that APAP- and NAPQI-induced death of HEI-OC1 cells would be mediated, at least in part, by ERS and the functionally related alterations in protein folding and degradation. These genes and their products, therefore, are bonafide surrogate markers that can be used in future cell biological experiments to monitor the toxic effects of these drugs.

3.4. The cytotoxic effect of APAP and NAPQI on HEI-OC1 cells is dependent of eIF2α and CHOP but independent of IRE1, ATF6, and ATF4-mediated signaling

We performed experiments to investigate whether the ERS signaling cascade mediated by PKR-like endoplasmic reticulum kinases (PERK) (Lin et al., 2009; Woehlbier and Hetz, 2011) could be the actual culprit for APAP/NAPQI-induced cell death. When activated by ERS, PERK induces phosphorylation of eukaryotic initiation factor 2 alpha (eIF2α), which works together with eIF2β to initiate protein biosynthesis. PERK-induced phosphorylation of eIF2α inhibits its interaction with eIF2β, thereby reducing protein synthesis and rescuing ERS. Inhibition of eIF2α also impairs the function of other well-characterized inducers of ERS, such as activating transcription factor 4 (ATF4) and CHOP, which function downstream eIF2β by regulating transcriptional responses (Wiseman and Kelly, 2011) (see Fig. 10). Guided by this knowledge, we searched for changes in eIF2α phosphorylation in response to APAP and NAPQI. We found that levels of phosphorylated eIF2α (p-eIF2α) increase in HEI-OC1 cells treated with APAP for 24 h, and decrease in those exposed to NAPQI at 24 and 48 h, suggesting a quick activating effect of APAP and a consistent inhibitory effect of
NAPQI on eIF2α phosphorylation (Fig. 6). Therefore, these results support a role for p-eIF2α in mediating APAP and NAPQI cytotoxic effects, leading us to study the role of other ER regulators working in conjunction with p-eIF2α.

We genetically inactivated ATF4 using specific shRNA for this transcription factor (Fig. 7A) and treated cells with APAP and NAPQI. To our surprise, we found that ATF4 knockdown was not able to rescue the cytotoxic effects of APAP and NAPQI (Fig. 7B).

In addition to PERK, ERS may result in the activation of two additional signaling cascades, one mediated by IRE1 (inositol-requiring enzyme 1) and the other by ATF6 (activating transcription factor 6) (Schonthal, 2009; Woehlhiber and Hetz, 2011) (see Fig. 10). Activation of IRE1 results in the unconventional splicing of the mRNA of its downstream target, the X-box binding protein 1 (XBP1), with the spliced form of XBP1 working as a transcription factor that induces expression of ER chaperones to ameliorate ERS (Lin et al., 2009, 2007; Woehlhiber and Hetz, 2011). Thus, we monitored the splicing of XBP1 mRNA in HEI-OC1 cells by RT-PCR from cells treated with APAP and NAPQI. Vehicle- and tunicamycin-treated cells were used as negative and positive Control, respectively. We found that cells exposed to APAP and NAPQI, in contrast to those exposed to tunicamycin, showed no clear signals of XBP1 splicing, suggesting that IRE1-mediated signaling was not readily activated by these drugs (Fig. 8A). We also evaluated the integrity of signaling pathways activated by IRE1 independently of XBP1, such as c-Jun N-terminal kinases (JNK) signaling, which is amenable for inhibition by the small drug SP600125. JNK inhibition by SP600125, however, did not affect HEI-OC1 cells’ survival (Fig. 8B). Combined, these results support the notion that IRE1 is unlikely involved in the toxic response to APAP and NAPQI in HEI-OC1 cells.

Finally, we tested the participation of ATF6 in the analgesic-induced toxic response. ERS induces translocation of this protein from the ER membrane to the Golgi, where it is enzymatically processed into a transcriptional regulator that activates the expression of ER chaperones and Endoplasmic Reticulum Associated Degradation genes (ERAD-related genes) (Woehlhiber and Hetz, 2011). The processing of ATF6 can be specifically inhibited by AEBSF (4-(2-aminoethyl) benzenesulfonyl fluoride), which antagonizes the activation of ATF6 within the Golgi, precluding its inducing effect on ERS. Thus, we speculated that, if ATF6-mediated signaling cascades were involved in the cytotoxic effect of APAP or NAPQI, inhibition of ATF6 with AEBSF would increase viability of HEI-OC1 cells treated with these drugs. We tested AEBSF at three different doses, 5 μM, 50 μM and 100 μM, but viability of APAP or NAPQI treated cells was not increased by any of them (Fig. 9). In agreement with these results, WB experiments indicated that CHOP expression was not changed either (at 24 h (n = 4):

Fig. 6. APAP and NAPQI changed levels of eIF2α phosphorylation. A – WB studies indicated that APAP maintains or increases eIF2α phosphorylation, whereas NAPQI nearly abolished it. B – Densitometric quantification of WB bands showed a significant APAP-induced increase in eIF2α phosphorylation at 24 h, and a significant NAPQI-induced decrease both at 24 and 48 h (At 24 h (* p = 0.003*). At 48 h (n = 4): Control – 100.0 ± 0.197, APAP – 1.583 ± 0.197, P < 0.0018; NAPQI – 0.327 ± 0.197, P < 0.016*. At 48 h (n = 4): Control – 1000 ± 0.197, APAP – 0.870 ± 0.197, N.S., NAPQI – 0.06 ± 0.197, P ≤ 0.0033).

Fig. 7. ATF4 knockdown did not affect APAP or NAPQI cytotoxicity. A – ATF4 shRNA transfected HEI-OC1 cells express ~20% of the ATF4 expressed either by Control or cells transfected with scrambled shRNA (negative Control). B – ATF4 shRNA did not increase viability of HEI-OC1 cells treated with APAP or NAPQI (At 24 h (n = 3): Control – 100.0 ± 4.9%; ATF4 shRNA Control – 93.4 ± 4.9%, P = N.S., re Control; APAP – 53.7 ± 4.9%; ATF4 shRNA APAP – 55.1 ± 4.9%, P = N.S. re APAP; NAPQI – 44.0 ± 4.5%; ATF4 shRNA NAPQI – 47.3 ± 4.9%, P = N.S. re NAPQI. At 48 h (n = 3): Control – 100.0 ± 5.1%; ATF4 shRNA Control – 81.8 ± 5.1%, P < 0.01* re Control; APAP – 33.4 ± 5.1%; ATF4 shRNA APAP – 45.6 ± 5.1%, P = N.S. re APAP; NAPQI – 38.3 ± 5.1%; ATF4 shRNA NAPQI – 40.4 ± 5.1%, P = N.S. re NAPQI).
Control = 1.000 ± 0.224; APAP = 1.624 ± 0.224, P=N.S. re Control; APAP + AEBSF (50 μM) = 1.260 ± 0.593, P=N.S. re APAP; NAPQI = 0.951 ± 0.224, P=N.S. re Control; NAPQI + AEBSF (50 μM) = 0.902 ± 0.593, P=N.S. re NAPQI. At 48 h (n = 4) : Control = 1.000 ± 0.224; APAP = 1.021 ± 0.242, P=N.S. re Control; APAP + AEBSF (50 μM) = 0.741 ± 0.593, P=N.S. re APAP; NAPQI = 0.796 ± 0.242, P=N.S. re Control; NAPQI + AEBSF (50 μM) = 0.741 ± 0.593, P=N.S. re NAPQI.

4. Discussion

Analogic preparations containing APAP and opioids, such as hydrocodone, have recently elicited significant attention because, if abused, they can rapidly induce profound sensorineural hearing loss (Blakley and Schilling, 2008; Curhan et al., 2010, 2012; Friedman et al., 2000; Ho et al., 2007; Oh et al., 2000; Rigby and Parnes, 2008; Yorgason et al., 2010). The cellular and molecular mechanisms underlying the ototoxic effects of this drug combination are currently unknown. In an attempt to fill the gap in this important field of research, we have previously reported studies on the role of the endoplasmic reticulum (ER) stress response (ERS) in mediating the ototoxic effects of APAP and NAPQI. ERS is activated by various stimuli, including drugs, and is characterized by the accumulation of unfolded and misfolded proteins in the ER, which leads to the induction of specific transcription factors and the activation of downstream pathways, including PERK, IRE1, and ATF6. 

Fig. 8. IRE1-mediated ERS pathway is not involved in APAP or NAPQI cytotoxicity. A – HEI-OC1 cells exposed to APAP and NAPQI, in contrast to those exposed to tunicamycin, showed no clear signals of XBP1 splicing. B – JNK inhibition by SP600125 did not affect the viability of HEI-OC1 cells treated with APAP or NAPQI. (At 24 h (n = 3) : Control = 100.0 ± 2.9%; S660125 = 103.2 ± 2.9%, P=N.S. re Control; APAP = 42.3 ± 2.9%; APAP + S660125 = 37.7 ± 2.9%, P=N.S. re APAP; NAPQI = 44.4 ± 2.9%; NAPQI + S660125 = 45.1 ± 2.9%, P=N.S. re NAPQI. At 48 h (n = 3) : Control = 100.0 ± 2.9%; S660125 = 102.5 ± 2.9%, P=N.S. re Control; APAP = 32.0 ± 2.9%, P = 0.0001* re Control; APAP + S660125 = 27.3 ± 2.9%, P=N.S. re APAP; NAPQI = 31.4 ± 2.9%, P ≤ 0.0001* re Control; NAPQI + S660125 = 31.2 ± 2.9%, P=N.S. re NAPQI).

Fig. 9. ATF6-mediated ERS pathway is not involved in APAP or NAPQI cytotoxicity. Treatment of HEI-OC1 cells with AEBSF, inhibitor of the ATF6-mediated pathway, did not prevent APAP or NAPQI cytotoxicity. (At 24 h (n = 3) : Control = 100.0 ± 3.0%; APAP = 51.4 ± 3.0%, P < 0.0001* re Control; APAP + AEBSF (5 μM) = 31.4 ± 3.0%, P < 0.0001* re APAP; APAP + AEBSF (50 μM) = 39.5 ± 3.0%, P < 0.0001* re APAP; APAP + AEBSF (100 μM) = 43.0 ± 3.0%, P = 0.0265* re APAP; NAPQI = 67.3 ± 3.0%, P < 0.0001* re Control; NAPQI + AEBSF (5 μM) = 61.7 ± 3.0%, P=N.S. re NAPQI; NAPQI + AEBSF (50 μM) = 44.8 ± 3.0%, P < 0.0001* re NAPQI; NAPQI + AEBSF (100 μM) = 57.0 ± 3.0%, P < 0.0008* re NAPQI. At 48 h (n = 3) : Control = 100.0 ± 3.0%; APAP = 29.2 ± 3.0%, P < 0.0001* re Control; APAP + AEBSF (5 μM) = 29.2 ± 3.0%, P=N.S. re APAP; APAP + AEBSF (50 μM) = 26.7 ± 3.0%, P=N.S. re APAP; APAP + AEBSF (100 μM) = 26.9 ± 3.0%, P=N.S. re APAP; NAPQI = 48.8 ± 3.0%, P < 0.0001* re Control; NAPQI + AEBSF (50 μM) = 48.3 ± 3.0%, P=N.S. re NAPQI; NAPQI + AEBSF (100 μM) = 45.6 ± 3.0%, P=N.S. re NAPQI.)

Fig. 10. Highly simplified diagram of the signaling pathways activated by ERS in mammalian cells. There are three major pathways mediated by the ER-resident transmembrane proteins ATF6, IRE1 and PERK. ERS increases GRP78 expression in the ER, and induces activation of one or more of the signaling pathways. After activation, ATF6 translocates to the Golgi, where it is cleaved by the Sit-1 and Sit-2 transmembrane proteins ATF6, IRE1 and PERK. ERS increases GRP78 expression in the ER, and induces activation of one or more of the signaling pathways. After activation, ATF6 translocates to the Golgi, where it is cleaved by the Sit-1 and Sit-2 transmembrane proteins. ERS increases GRP78 expression in the ER, and induces activation of one or more of the signaling pathways. After activation, ATF6 translocates to the Golgi, where it is cleaved by the Sit-1 and Sit-2 transmembrane proteins.
that clarified the pathobiological effects of APAP and hydrocodone on cochlear hair cells, providing evidence that APAP, and not hydrocodone, was the actual culprit of the ototoxic effects of the compound (Yorgason et al., 2010). We also identified OHCs as the major target of APAP toxicity in the mouse organ of Corti, and provided evidence that HEI-OC1 cells, although not actual hair cells, are a good model to investigate the cellular and molecular mechanisms underlying APAP ototoxicity (Yorgason et al., 2010). Thus, in the current study, our goal has been to further increase the current understanding of the process of analgesic-induced ototoxicity by identifying mechanisms by which APAP affects the viability of cultured HEI-OC1 cells. Our results indicate that both APAP and its metabolite NAPQI are toxic for HEI-OC1 cells, but different cellular mechanisms contribute to mediate their cytotoxicity.

4.1. APAP, but not NAPQI, causes HEI-OC1 cells death by inducing oxidative stress

Our investigation focused on evaluating the toxic effects associated with APAP and its metabolite, NAPQI, separately. We designed our studies starting from dose–response curves that describe the concentrations of both APAP and NAPQI that cause noticeable ototoxic effects in HEI-OC1 cells, so that future studies can be guided by these experiments. We used a concentration of 1.5 mg/mL (~10 mM) APAP, which is less than APAP EC50 in human hepatocyte cultures (28.2 mM), but more than APAP EC50 in mouse hepatocyte cultures (3.8 mM), as determined by cell viability studies following 24 h treatment (Jennitz et al., 2008). Since approximately 10% APAP is metabolized into NAPQI, a concentration of 150 µg/mL (~1 mM) NAPQI was used in our experiments with this compound. Using these doses, we demonstrated that treatment with APAP for short periods actually decrease ROS levels in HEI-OC1 cells, but exposure for more than 6 h induces a significant increase in oxidative stress; NAPQI, in contrast, decreases ROS levels in HEI-OC1 cells at all the time-points investigated (Fig. 2). The observed anti-oxidant effect of APAP at short periods of exposure was at odds with the abundant literature about its effects on hepatocytes, but consistent with reports that APAP acts as an anti-inflammatory and anti-oxidant compound in neurons both in vitro and in vivo. For example, APAP protects hippocampal neurons and PC12 cultures from oxidative stress by lowering cytoplasmic levels of peroxides and reducing of lipid peroxidation (Bisaglia et al., 2002), and cerebral cortical cultured neurons and brain endothelial cells exposed to the superoxide-generating compound menadione (Tripathy and Grammas, 2009a, 2009b).

In rats, APAP induces apoptosis of cortical neurons (Posadas et al., 2010), but significantly attenuates superoxide production by the neutrotoxic quinolinic acid, a metabolite implicated in the pathogenesis of neurodegenerative diseases (Maharaj et al., 2006). In a C. elegans model of Parkinson’s disease APAP also protects dopaminergic neurons from oxidative damage evoked by exposure to 6-hydroxy dopamine or excessive levels of dopamine (Lock et al., 2008). Therefore, we can speculate that HEI-OC1 cells probably respond to APAP just as neurons, which seems reasonable given their neuro-epithelial origin. The decrease in ROS levels induced by NAPQI, in contrast, is consistent with the results of Harman et al. (1991) in cultured hepatocytes. These authors presented solid evidence that killing of hepatocytes by APAP was dependent on a cellular source of ferric iron, potentiated by 1.3-bis-(2-chloroethyl)-1-nitrosourea (BCNU, an inhibitor of glutathione reductase), and very sensitive to antioxidants. By contrast, NAPQI hepatotoxicity was not prevented by chelation of ferric iron, was unaffected by BCNU, and was insensitive to antioxidants (Harman et al., 1991). Moreover, the killing of cultured hepatocytes by NAPQI was preceded by a collapse of the mitochondrial membrane potential and a depletion of ATP, suggesting that mitochondrial de-energization by a mechanism unrelated to oxidative stress was likely the mechanism of hepatocytes killing by NAPQI (Harman et al., 1991). Thus, the results by Harman et al. and our own results suggest that, though important, oxidative stress would not completely explain the hepatotoxin- and otoxic effects of both drugs, a finding that led us to search for additional mechanisms underlying this process in HEI-OC1 cells.

4.2. ER stress is a shared mechanism underlying the toxic effects of APAP and NAPQI on HEI-OC1 cells

Information on APAP neurotoxicity is scarce and, to our knowledge, no toxic effects have been described in the human nervous system. In agreement with our results, reported studies suggest that APAP concentrations up to 100 mM or doses up to 100 mg/kg prevent ROS production and cytotoxicity in neurons, but higher concentrations, far from protecting cells from ROS toxicity, might actually contribute to ROS production (Posadas et al., 2010). Also in agreement with our results indicating that APAP and NAPQI effects would be mediated by different cellular mechanisms, NAPQI, but not APAP itself, would activate mouse and human transient receptor potential ankyrin-1 (TRPA1) channel, reducing voltage-gated calcium and sodium currents in primary sensory neurons and releasing pro-inflammatory neuropeptides from sensory nerve terminals (Nassini et al., 2010).

The mechanisms of APAP and NAPQI liver toxicity, in contrast, have been extensively studied (see, for example, Jaeschke et al., 2012, 2011; Moyer et al., 2011). However, there is no complete consensus about the multiple steps and signaling pathways involved in the process. It is generally believed that APAP hepatotoxicity is initiated by formation of its reactive metabolite, NAPQI, which depletes glutathione and binds to cellular proteins, especially in mitochondria. The resulting mitochondrial oxidant stress and peroxynitrite formation, in part through amplification by c-Jun-N-terminal kinase activation, leads to mitochondrial DNA damage and opening of the mitochondrial permeability transition pore. Endonucleases from the mitochondrial inter-membrane space and lysosomes would be responsible for nuclear DNA fragmentation. The mitochondrial dysfunction and nuclear DNA damage ultimately cause oncocytic cell death with release of molecular byproducts that trigger a sterile inflammatory response (Jaeschke et al., 2012, 2011). Therefore, some authors believe that NAPQI, not APAP, is responsible of liver toxicity. Interestingly, studies in hepatoma cell lines, which possess very low levels of CYP450 enzymes and therefore cannot efficiently metabolize APAP into NAPQI, indicate that APAP exposure induces caspase-dependent apoptosis, in contrast to cell death of metabolically competent hepatocytes that occurs by oncocytic necrosis as indicated by cell and organelle swelling, massive cell contents release, and karyorrhexis and karyolysis (Bajt et al., 2004; Gujral et al., 2002; McGill et al., 2011). Thus, APAP and NAPQI cytotoxicity would be associated with different pathways: one leading to cell apoptosis and the other to necrosis, and the balance between these two mechanisms would be cell type-dependent. Although the cellular effects of APAP itself are still under discussion, there are no doubts about NAPQI cytotoxicity. Thus, NAPQI cytotoxicity has been investigated with genomic, proteomic and transcriptomic approaches. In a recent work, 176 different lymphoblastic cell lines from healthy human subjects were used in a genom-wide study (Moyer et al., 2011). A group of single nucleotide polymorphisms (SNPs) in linkage disequilibrium on chromosome 3 was highly associated with NAPQI toxicity. This group of SNPs mapped to a “gene desert” area, but chromatin immunoprecipitation assays demonstrated binding of several
transcription factor proteins including heat shock factor 1 (HSF1) and HSF2. The mechanism responsible for this association, however, was unclear, and the study provided no decisive information about the actual mechanism of NAPQI cytotoxicity (Moyer et al., 2011).

In the present study we demonstrate that, in HEI-OC1 cells, both APAP and NAPQI are cytotoxic, but their toxic effects are mediated by different mechanisms. To elucidate the molecular mechanisms underlying APAP and NAPQI cytotoxicity in HEI-OC1 cells, we complemented our cellular biology experiments with gene expression levels and proteomic profiling. Our initial design involved the use of pathway specific arrays to study the expression of genes involved in toxicity in a focused manner. This method was subsequently complemented using proteomics (iTRAQ), which provides both an unbiased validation of genomic studies as well as complementing the information about gene expression levels with information about protein production. These experiments produced information leading to the conclusion that both compounds regulate gene and protein networks that participate in the cell response to xenobiotic metabolism, oxidative radical metabolism, ER stress, protein ubiquitination, and cell death. Consequently, guided by these results and those of OMIC experiments with APAP performed by other researchers in liver cells, we designed cellular biological experiments to define whether ER stress was indeed part, together with oxidative stress, of the cellular mechanisms associated with the toxic effects of APAP and NAPQI. We examined in detail the impact of these compounds on the structure and function of the ER using well-characterized markers of ER stress, and our observations revealed that APAP and NAPQI changed the levels of intra-cisternal chaperones in a similar way to what it has been previously reported as surrogate of ER stress, ultimately causing the fragmentation of this organelle. Further experiments supported the involvement of the PERK-mediated pathway, but not the IRE1- and ATF6-mediated pathways, in the APAP/NAPQI-induced ER stress (Fig. 10). Interestingly, APAP and NAPQI showed different effects on eIF2α phosphorylation, and consequently on protein synthesis, and their effects were not associated with ATF4 activation. Thus, these studies reveal that ER stress might be a key pathophysiological mechanism responsible for the toxic effects of APAP and NAPQI on auditory cells and outlines molecules and pathways that, if inactivated, could prevent these deleterious effects.

5. Conclusions

In summary, we have used HEI-OC1 cells as an in vitro model for studying the potential ototoxic effects of APAP, a commonly used analgesic drug that causes hearing loss. More importantly, our results suggest that APAP and its metabolite NAPQI could be achieving its ototoxic effects by triggering both oxidative stress and endoplasmic reticulum stress. Since several approaches are being currently developed to modulate both of these two deleterious cellular responses, we are optimistic that our results will guide future studies aimed at designing and testing potential therapeutic intervention aimed at preventing hearing loss caused by these compounds.

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Appendix A. Supplementary data

Supplementary data related to this article can be found at http://dx.doi.org/10.1016/j.heares.2014.04.007.
causes neurogenic inflammation in the airways and other tissues in rodents. FASEB J. 24, 4904–4916.