Enzyme-Mediated Protein Haptenation of Dapsone and Sulfamethoxazole in Human Keratinocytes: I. Expression and Role of Cytochromes P450

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ABSTRACT
Cutaneous drug reactions (CDRs) are among the most common adverse drug reactions and are responsible for numerous minor to life-threatening complications. Several arylamine drugs, such as sulfamethoxazole (SMX) and dapsone (DDS), undergo bioactivation, resulting in adduction to cellular proteins. These adducted proteins may initiate the immune response that ultimately results in a CDR. Recent studies have demonstrated that normal human epidermal keratinocytes (NHEKs) can bioactivate these drugs, resulting in protein haptenation. We sought to identify the enzyme(s) responsible for this bioactivation in NHEKs. Using immunofluorescence confocal microscopy and an adduct-specific enzyme-linked immunosorbent assay (ELISA), we found that N-acetylation of the primary amine of SMX and DDS markedly reduced the level of protein haptenation in NHEKs. Detection of mRNA and/or protein confirmed the presence of CYP3A4, CYP3A5, and CYP2E1 in NHEKs. In contrast, although a faint band suggestive of CYP2C9 protein was detected in one NHEK sample, a CYP2C9 message was not detectable. We also examined the ability of chemical inhibitors of cytochromes P450 (aminobenzotriazole and 1-dichloroethylene) and cyclooxygenase (indomethacin) to reduce protein haptenation when NHEKs were incubated with SMX or DDS by either confocal microscopy or ELISA. These inhibitors did not significantly attenuate protein adduction with either SMX or DDS, indicating that cytochromes P450 and cyclooxygenase do not play important roles in the bioactivation of these xenobiotics in NHEKs and thus suggesting the importance of other enzymes in these cells.

Sulfonamides are used in the treatment of numerous infectious diseases. Their effectiveness in the treatment of Pneumocystis carinii pneumonia, especially in AIDS patients, has increased their importance as therapeutic agents in the modern antimicrobial era (Cribb et al., 1996a). Sulfamethoxazole (SMX) and the sulfone dapsone (DDS) are widely used antimicrobials for the treatment of Pneumocystis carinii pneumonia resulting in the recovery of approximately 75% of the patients suffering from this ailment (Hughes, 1987; Goldie et al., 2002). However, adverse drug reactions, especially those of a cutaneous nature, limit their use. These reactions most commonly occur 7 to 10 days after initiation of therapy and are associated with fever and skin rash. Morbilliform or maculopapular nonurticarial types of skin rash are most commonly observed with these agents. Some patients, however, progress to Stevens-Johnson syndrome or toxic epidermal necrolysis, which can have a mortality rate as high as 40 to 50%. A multiorgan syndrome, manifested as fever, rash, eosinophilia, and hepatotoxicity, has also been reported in patients receiving these drugs (Dujovne et al., 1967; Berg and Daniel, 1987; Rieder et al., 1989; Svensson et al., 2001).

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ABBREVIATIONS: SMX, sulfamethoxazole; DDS, dapsone; NHEK, normal human epidermal keratinocyte; CYP450, cytochrome(s) P450; COX, cyclooxygenase; MADDS, monoacetyl dapsone; ABT, 1-aminobenzotriazole; INDO, indomethacin; 7-BQ, 7-benzyloxyquinoline; 7-HQ, 7-hydroxyquinoline; DCE, trans-dichloroethylene; ELISA, enzyme-linked immunosorbent assay; NASMX, N-acetyl sulfamethoxazole; PCR, polymerase chain reaction; PBS, phosphate-buffered saline; ANOVA, analysis of variance; DMSO, dimethyl sulfoxide.
Several studies have demonstrated that SMX and DDS undergo biotransformation to form reactive N-arylhydroxylamine metabolites (Fig. 1), which are believed to be responsible for causing cutaneous drug reactions (Cribb et al., 1996a; Reilly and Ju, 2002; Svensson, 2003). It has been proposed that the bioactivation of these drugs at the site of manifestation (i.e., skin) may be a critical element in the initiation of these reactions (Reilly et al., 2000). Indeed, we have previously demonstrated that normal human epidermal keratinocytes (NHEKs) are able to bioactivate SMX and DDS to their respective arylhydroxylamine metabolites (sulfamethoxazole hydroxylamine and dapsone hydroxylamine) (Reilly et al., 2000) and that such bioactivation results in protein haptenation (Roychowdhury et al., 2005).

Studies have demonstrated that skin cells express a variety of drug-metabolizing enzymes, such as cytochromes P450 (CYP450), cyclooxygenase (COX), flavin-containing monooxygenases, and peroxidases, which may bioactivate numerous chemical agents (Kanekura et al., 1998; Rys-Sikora et al., 2000; Baron et al., 2001; Jannmohamed et al., 2001; Saeki et al., 2002). Hydroxylation of SMX and DDS by various CYP450s has been reported (Cribb et al., 1995; Mitra et al., 1995; Gill et al., 1999; Winter et al., 2000). Several arylamine drugs, including procainamide, are oxidized to arylhydroxylamine metabolites by COX-2 (Liu and Levy, 1998; Goebel et al., 1999).

In the present investigation, we sought to identify the enzyme(s) that catalyze the bioactivation of SMX and DDS, leading to protein haptenation in NHEKs. Our studies demonstrate that although COX-1, CYP3A4, and CYP2E1 are able to catalyze the bioactivation of SMX and DDS in vitro, inhibitors of these enzymes do not reduce adduct formation when NHEKs are exposed to either SMX or DDS. These data suggest that these enzymes do not play an important role in the bioactivation and subsequent protein adduction of SMX and DDS in NHEKs.

**Materials and Methods**

Materials. Monoacetyl dapsone (MADDS) was generously provided by Parke-Davis (now Pfizer, Ann Arbor, MI). DDS, SMX, 4-dimethylaminopyridine, 1-aminobenzotriazole (ABT), troleandomycin, disulfiram, and indomethacin (INDO) were obtained from Sigma (St. Louis, MO). 7-Benzoyloxquinoline (7-BQ) and 7-hydroxyquinoline (7-HQ) were purchased from Gentest (Bedford, MA). DDS and SMX hydroxylamine metabolites were synthesized as described previously (Vyas et al., 2005). Trans-dichloroethylene (DCE) was obtained from TCI America (Portland, OR). Rabbit anti-sera were raised against SMX- and DDS-keyhole limpet hemocyanin conjugates and specificity assessed as described previously (Reilly et al., 2000). Rat tail collagen (type I) was obtained from Sigma. Normal human epidermal keratinocytes (as first passage cells) and keratinocyte culture media were obtained from CAMBREX (Walkersville, MD). The HaCaT cell line was generated by Dr. N. Fusenig (DKFZ Heidelberg, Heidelberg, Germany) and obtained from Dr. Michael Southhall (Johnson and Johnson, Skillman, NJ). Microtiter ELISA plates (96-well) were obtained from Rainin Instruments (Woburn, MA). Rabbit anti-CYP2C9 antibody was custom made by Covance Research Products Inc. (Denver, PA). Donkey anti-rabbit IgG conjugated to horseradish peroxidase was obtained from Amersham Pharmacia Biotech Inc. (Piscataway, NJ). SuperSignal West Pico Chemiluminescent Substrate was purchased from Pierce (Rockford, IL), and a SynGene GeneGnome chemiluminescence detection system was obtained from Synoptics (Cambridge, UK). Rabbit polyclonal anti-CYP3A4 has been described previously (Beaune et al., 1985); rabbit α-3A5 primary antibody was purchased from Gentest. Goat-anti-rabbit IgG conjugated with Alexa fluor 488 and goat anti-rabbit antibody conjugated with alkaline phosphatase and YoYo-1 were purchased from Molecular Probes (Eugene, OR). Bradford assay reagent was purchased from Pierce. Immunomount was obtained from Vector Laboratories (Burlingame, CA). Trizol was purchased from Invitrogen (Carlsbad, CA). All other chemicals and reagents were purchased from Sigma or Fisher Scientific (Chicago, IL). Human liver microsomes were processed at St. Jude Children's Research Hospital (Nashville, TN), and tissue was provided by the Liver Tissue Procurement and Distribution System (National Institutes of Health Contract N01-DK-9-2310) and by the Cooperative Human Tissue Network.

**Synthesis of N-Acetyl sulfamethoxazole.** SMX (250 mg) was dissolved in 1 ml of acetic anhydride in the presence of 0.5 g of 4-dimethylaminopyridine and 10 ml of dichloromethane. The reaction was carried out for 1 h at room temperature with continuous stirring to obtain N-acetyl sulfamethoxazole (NASMX). Excess of acetic anhydride was removed by neutralizing the reaction mixture with saturated sodium bicarbonate solution and NASMX obtained by recrystallizing the product in an acetone and hexane (1:3) mixture. NASMX was characterized by NMR, and the purity was determined by high-performance liquid chromatography-UV. Product purity was 98%, whereas the observed yield was 79%.

**Cell Culture.** Adult NHEKs and immortalized HaCaT cells were cultured as detailed previously (Reilly et al., 2000). In brief, cells were propagated in 75-cm² flasks using basal media (KBM-2 supplemented with bovine pituitary extract (7.5 mg/ml), human epidermal growth factors (0.1 ng/ml), insulin (5 μg/ml), hydrocortisone, (0.5 μg/ml), epinephrine, transferrin, gentamicin (50 μg/ml), and amphotericin (50 μg/ml) at 37°C in an atmosphere containing 5% CO₂. Media were replaced every 2 to 3 days. When cell cultures reached near confluency (70–90%), cells were disaggregated using 0.025% trypsin/0.01% EDTA in HEPES followed by neutralization with 2 volumes of Trypsin neutralizing solution. Cell suspensions were then centrifuged at 220g for 5 min followed by washing in basal media and resuspension in KGM-2 (supplemented growth medium). Cells were
then either subjected to subculturing or cryopreservation for further purposes. All experiments were performed using third to fourth passage cells.

**Determination of mRNA for CYP2C9 in NHEKs.** Total RNA was isolated from fourth passage NHEKs and seventh passage HaCaT cells using Trizol (Invitrogen) following the manufacturer’s protocol. cDNA was generated using 500 ng of total RNA in a synthesis reaction using SuperScript II RT with random hexamers according to the supplier’s protocol. PCR was performed using the CYP2C9-specific primers 5'-GATCTGAAATATTTTTCTC-3' and 5'-TCTCAGGTGTGTCTGTC-3', with diluted amounts of cDNA from the keratinocytes resulting in a 280-bp product. Plasmid CYP2C9 cDNA in pcW was used as a positive control for amplification. Actin was also amplified from the keratinocyte cDNA using the primers 5'-TCATGGAATGTGACAGTCCTG-3' and 5'-CTCTAGAAGCATTTGCGGTGCACGATG-3', resulting in a 285-bp fragment. Amplifications were performed in 20-μl reactions using AmpliTaq Gold (Applied Biosystems, Foster City, CA) according to the manufacturer’s protocol with a final concentration of MgCl₂ of 2 mM. Cycling conditions consisted of an initial denaturation at 94°C for 5 min followed by 40 cycles of 94°C for 30 s, 53°C for 10 s, and 72°C for 10 s. Reactions were analyzed on 4% agarose gels stained with ethidium bromide.

**Immunoblot Analysis for CYP2C Proteins in NHEKs.** Cell lysates and yeast-expressed recombinant CYP2C proteins were fractionated by electrophoresis in SDS-10% (w/v) polyacrylamide gels and transferred to nitrocellulose membranes. Membranes were immunoblotted with a polyclonal rabbit anti-CYP2C9 antibody (1:500) raised to bacterially expressed human CYP2C9 and donkey anti-rabbit IgG conjugated to horseradish peroxidase. This anti-CYP2C9 polyclonal antibody to bacterially expressed recombinant CYP2C9 was made for National Institute of Environmental Health Sciences rabbit antibody production protocol. Yeast-expressed CYP2C proteins were used as standards on immunoblots because there is no N-terminal modification for cDNA expression in yeast. Therefore, the recombinant yeast-expressed CYP2C proteins have mobility identities to those of the human CYP proteins expressed in vivo. Specific bands were visualized with SuperSignal West Pico Chemiluminescent Substrate and a SynGene GeneGnome chemiluminescence detection system.

**Determination of mRNA for CYP3A4 and CYP3A5 in NHEKs.** Total RNA was isolated from NHEKs and HaCaT cells using Trizol Reagent (Invitrogen) and first strand cDNA was synthesized from 3 μg of total RNA according to the manufacturer’s instructions (SuperScript First Strand cDNA Synthesis kit; Invitrogen). Amplification of CYP3A4 cDNA (GenBank accession no. AF182273) from first-strand cDNA was performed using: forward, 5'-CCACAGCTTGTGGCATGACACAC-3' and reverse, 5'-GAGGTCTCCTTGTGGTCTGTCCAG-3' primers that annealed to nucleotides in exons 1 and 13. PCR was carried out in a total reaction volume of 50 μl consisting of 5× PCR buffer with 1.5 mM MgCl₂, 1 μl, 5 pmol of each primer, 0.2 mM dNTP (Life Technologies, Gaithersburg, MD), and 2.5 U of TaqDNA Polymerase (Expand High Fidelity PCR system; Boehringer Mannheim, Mannheim, Germany). PCR consisted of initial denaturation at 94°C for 3 min followed by 35 cycles of denaturation at 94°C for 30 s, annealing at 55°C for 30 s, synthesis at 72°C for 90 s, and a final extension at 72°C for 10 min. No CYP3A4 was detected, so an aliquot of the first round product was PCR amplified for a second round with the same primers and conditions to detect whether very small amounts of mRNA were present. CYP3A5 was amplified using: P1-forward, 5'-AACAGC-CCACGAAAGAAGCAGC-3' (forward); and P2-reverse, 5'-TAAGCC-CCATCCTTTTATTTCTAAGGT-3' primers as described previously (Kuehl et al., 2001), and product was detected after the first round amplification.

**Determination of Protein for CYP3A4 and CYP3A5 in NHEKs.** NHEKs, HaCaT cells, and human liver samples were re-suspended in storage buffer (100 mM potassium phosphate, pH 7.4, 1.0 mM EDTA, 20% glycerol, 1 mM dithiothreitol, 20 μM butylated hydroxytoluene, and 2 mM phenylmethylsulfonyl fluoride), lysates were generated by sonication, and protein concentrations were determined using the Bio-Rad protein assay. Lysate (30 μg) was separated on 10% SDS-polyacrylamide gels and immunoblotted with monoclonal anti-CYP3A4 K03 or rabbit α-3A5 (Gentest), followed by appropriate secondary antibodies coupled with peroxidase. The blot was developed with the ECL detection system (Amersham Biosciences).

**Determination of Catalytic Activity for CYP3A4 in NHEKs.** NHEKs (10⁶ cells) were incubated in KGM-2 overnight. After 24 h, the media were discarded and replaced with fresh media. 7-BQ (50 μM), a substrate for CYP3A4, was added to measure the catalytic activity of CYP3A4 in these cells. The plate was incubated for 30 min at 37°C after the addition of 7-BQ followed by the fluorescence measurement every 30 min for 24 h using the CytoFluo multwell fluorescent plate reader (excitation wavelength, 410 nm; emission wavelength, 538 nm). The fluorescent metabolite of 7-BQ, 7-HQ, was also added to NHEKs at varying concentrations (0, 5, 10, 25, and 50 μM) to determine the ability to detect this metabolite in the presence of NHEKs.

**Determination of CYP2E1 in NHEKs.** CYP2E1 mRNA expression was quantified in NHEKs and HaCaT cells essentially as described by Haufroid et al. (2001). Briefly, total RNA was isolated from keratinocytes using Trizol reagent (Invitrogen) and quantified using a Quant-it RiboGreen kit (Invitrogen) as per the manufacturer’s instructions. Using 2 μg of total RNA from each cell preparation, cDNA was prepared using random hexamer primers and ImProm-II reverse transcriptase following the procedures outlined in the ImProm-II Reverse Transcription System kit (Promega, Madison, WI). RT-PCR reactions to quantify CYP2E1 mRNA concentrations were carried out in a total volume of 40 μl containing 5.5 mM MgCl₂, 0.2 mM each deoxyribonucleotide, 0.3 μM of each PCR primer, 0.2 μM fluorescent probe, 5 or 25 ng of keratinocyte cDNA, 0.01 U/μl uracil N-glycosylase, and 0.025 μM AmpliTaq Gold Polymerase (TaqMan PCR Core Reagents kit; Applied Biosystems) in an Option System (MJ Research, Watertown, MA). 18S rRNA concentrations were determined using the TaqMan Ribosomal RNA Control Reagents (Applied Biosystems), which were designed to generate a 187-bp product. Each 40-μl PCR reaction contained 5.5 mM MgCl₂, 0.2 mM each deoxyribonucleotide, 0.05 μM of each PCR primer, 0.2 μM fluorescent probe, 5 or 25 ng of keratinocyte cDNA, 0.01 U/μl uracil N-glycosylase, and 0.025 μM AmpliTaq Gold Polymerase (TaqMan PCR Core Reagents kit; Applied Biosystems).

**Determination of the Cytotoxicity of Inhibitors in NHEKs.** To determine the maximal noncytotoxic concentrations of various inhibitors in NHEKs for the inhibition of the respective enzymes, the cytotoxicity of concentrations ranging from 25 μM to 25 mM were examined. Cytotoxicity was determined using an impermeable DNA binding dye (YoYo-1), as we have described previously (Vyas et al., 2005).

**ELISA Analysis of Drug/Metabolite-Protein Adducts.** Formation of covalent adducts following SMX or DDS exposure, in the presence or absence of the selective enzyme inhibitors, was determined by cultivating NHEKs (1 × 10⁶ cells) for 24 h in 50-ml centrifuge tubes containing 10 ml of complete growth medium. The concentrations of the selective inhibitors used were the maximum noncytotoxic concentrations of those inhibitors in NHEKs. Cells were then incubated with selective enzyme inhibitors for 3 h followed by SMX (1 mM ascorbic acid was added before the addition of SMX) or DDS treatment for 3 h (concentrations specified under Results). Following total 6-h incubation, tubes were centrifuged at 220 g at 4°C for 30 min to pellet the cells. Covalent adducts were determined as previously described (Vyas et al., 2005).

**Immunocytochemistry.** Drug/metabolite-protein covalent adduct formation was visualized using immunofluorescence confocal microscopy. Cells were grown on collagen-coated (0.1 mg/ml) coverg...
slips placed in Petri dishes containing 2 ml of complete growth medium. After 24 h, cultures were subjected to different selective enzyme inhibitors for 3 h (maximum noncytotoxic concentrations were used), followed by SMX or DDS treatment for 3 h (concentrations specified under Results). After the 3-h incubation, cells were washed (three times) with phosphate-buffered saline (PBS; 0.05 M sodium phosphate, 0.15 M NaCl, pH 7.4) and fixed for 20 min with 4% paraformaldehyde in PBS. After fixation, cultures were washed three times with PBS followed by blocking for 60 min with Tris-casein buffer containing 0.3% Triton X-100 and overnight incubation with the anti-DDS or anti-SMX antisera (1:50 diluted in blocking buffer) at 4°C. Coverslips were then washed with PBS, incubated for 3 h at 37°C with the fluorochrome-conjugated secondary antibody (Alexa Fluor-488-labeled goat-anti-rabbit IgG, 1:500 diluted in blocking buffer), and mounted on glass slides using Immunomount containing antifade reagent.

Fluorescence images were acquired with a Zeiss Laser Scanning Microscope (LSM 510, Zeiss Axiovert stand, Zeiss 63× and 20× objective lenses; Carl Zeiss GmbH, Jena, Germany) using excitation at 488 nm. Emission was set to a long-pass filter at 505 nm.

**Image Analysis.** For imaging with the confocal laser scanning microscope, laser attenuation, pinhole diameter, photomultiplier sensitivity, and exposure were kept constant for every set of experiments. Images were acquired from three different view fields of each slide. The obtained images were quantitatively analyzed for changes in fluorescence intensities within regions of interest (boxes drawn over cell somata) using the NIH Image J software (Bethesda, MD). Fluorescence values from minimum of three view fields consisting of 15 to 20 NHEK cells in each field from three different slides of each treatment were averaged and expressed as mean (S.D.) fluorescence intensity.

**Statistical Analysis.** Data are presented as mean (S.D.). Data were analyzed using SigmaStat (Systat Software, Inc., Point Richmond, CA). Statistical comparisons between two groups were made using either Student’s t test (parametric method) for normalized data or Friedman’s rank sum test (nonparametric method) for the data that did not pass the normality test. For the comparison between more than two groups, ANOVA and the Holm-Sidak method for multiple pair-wise comparisons were used. p < 0.05 was considered to be significant.

**Results**

**Covalent Adduct Formation with DDS and SMX and Their Acetylated Metabolites in NHEKs.** Protein haptenation of the parent arylamine drugs, DDS and SMX, was readily detected by both confocal microscopy and ELISA (Figs. 2 and 3). N-Acetylation of SMX or DDS resulted in marked attenuation of protein haptenation. As we have noted previously, the amount of adduct formed with DDS appears to be greater than that seen with SMX, although this may represent differences in antisera affinity for the respective adducts.

**Expression of Various Cytochromes P450 in Keratinocytes.** NHEKs and an immortalized keratinocyte cell line (HaCaT) were probed for the presence of various CYP450 known to catalyze the formation of the arylhydroxylamine metabolites of SMX and DDS in vitro. As shown in Fig. 4, CYP2C9 mRNA was not detected in cells from cultures of primary keratinocytes (NHEKs) and HaCaT cells using multiple dilutions of cDNA. However, immunoblot analysis suggested the possible presence of CYP2C9 (a diffuse polypeptide band with approximately the same mobility as recombinant CYP2C9) in the first sample of NHEKs, but that was not seen in the second sample of cells from the same patient or in HaCaT cells. Moreover, CYP2C8, CYP2C18, or CYP2C19 were not detected in either cell type (Fig. 5). The recombinant CYP2C protein standards were expressed in a yeast cDNA expression system as described previously (Gold-
stein et al., 1994). Interestingly, although both CYP3A4 and CYP3A5 mRNA were observed in NHEKs, only CYP3A5 mRNA was observed in HaCaT cells (Fig. 6A). Immunoblot analysis using an antibody specific for CYP3A5 yielded positive results in both NHEKs and HaCaT cells, as did an antibody that recognizes both CYP3A4 and CYP3A5 (Fig. 6B). However, the levels of CYP3A4 and CYP3A5 proteins were much lower compared with the human livers. CYP2E1 mRNA was also found to be expressed in both NHEKs and HaCaT cells (Fig. 7), whereas CYP2E1 protein was barely detectable (data not shown).

**CYP3A4 Activity in NHEKs.** After the addition of 7-BQ, no increase in fluorescence above background (NHEKs alone) was observed (data not shown). In contrast, a concentration-dependent increase in fluorescence was observed when the expected metabolite (7-HQ) was added to NHEKs (data not shown).
human livers were probed for CYP3A proteins using a monoclonal antibody. Moreover, we did not detect the presence of the most probable enzymes for bioactivation of SMX and DDS in NHEKs is unknown.

To assess the importance of arylamine oxidation, we evaluated the impact of SMX and DDS N-acetylation on protein haptentation in NHEKs. We found that the N-acetyl metabolites gave rise to a lower level of protein haptentation as measured by both ELISA and confocal microscopy (Figs. 2 and 3). It should be noted that it is possible that the apparent reduction in adduct formation could be secondary to a reduced affinity of the antisera for the adduct in the presence of an acetyl group on the drug, as opposed to an actual reduction in the amount of adduct formed. Differentiation of these two potential explanations could only be accomplished by the characterization of and development of chemical methods for the quantification of the drug-protein adducts.

Before evaluating the effect of various enzyme inhibitors on protein haptentation with these drugs, we sought to confirm the presence of the most probable enzymes for bioactivation in NHEKs. Because HaCaT cells are widely used as an alternative to primary cultures of NHEKs, together with our previous observation of protein haptentation in these cells when exposed to SMX or DDS (Roychowdhury et al., 2005), we also examined the presence of CYP450 enzymes in this cell line. CYP2C9 appears to be the most important enzyme for bioactivating SMX and DDS in the human liver (Cribb et al., 1995; Uetrecht, 1990; Mitra et al., 1995; Gill et al., 1999; Winter et al., 2000). Several arylamine drugs, including procainamide, are oxidized to arylhydroxylamine metabolites by COX-2 in vitro (Liu and Levy, 1998; Goebel et al., 1999). Which of these enzymes, if any, mediates the bioactivation of SMX and DDS in NHEKs is unknown.

Protein Haptenation of Dapsone and Sulfamethoxazole

Discussion

The biotransformation of xenobiotics to reactive metabolites is believed to be responsible for a wide range of adverse reactions. Sulfonamides such as SMX and the sulfone DDS have been reported to be bioactivated to arylhydroxylamine metabolites, which readily auto-oxidize to arylnitroso species (Fig. 1). These metabolites are believed to initiate the cascade of events that ultimately provoke a cutaneous drug reaction (Cribb et al., 1996b; Svensson, 2003). We have proposed previously that bioactivation in the skin may play an important role in these reactions (Reilly et al., 2000). Indeed, incubation of NHEKs with SMX or DDS results in protein haptentation (Roychowdhury et al., 2005).

In vitro studies have shown that the bioactivation of these parent arylamine drugs to their arylhydroxylamine metabolites may be mediated by various oxidizing enzymes such as CYP2C9, CYP2E1, CYP3A4, and MPO (Cribb et al., 1990, 1995; Uetrecht, 1990; Mitra et al., 1995; Gill et al., 1999; Winter et al., 2000). Several arylamine drugs, including procainamide, are oxidized to arylhydroxylamine metabolites by COX-2 in vitro (Liu and Levy, 1998; Goebel et al., 1999). Which of these enzymes, if any, mediates the bioactivation of SMX and DDS in NHEKs is unknown.

Covalent Adduct Formation of DDS and SMX in the Presence and Absence of Inhibitors of Various Enzymes.

The effect of various enzyme inhibitors on protein haptentation in NHEKs exposed to SMX or DDS was evaluated by both confocal microscopy and ELISA. For each inhibitor, we used the highest concentration that did not cause cytotoxicity in NHEKs. As shown in Fig. 8, a broad inhibitor of CYP450 (ABT) that inhibits CYP3A4 and CYP2C9, a selective inhibitor of CYP2E1 (DCE), and an inhibitor of COX (INDO) failed to reduce the protein haptentation of DDS in NHEK cells. Similar results were observed with SMX by ELISA (Fig. 9). Similar results were obtained when inhibitors were added simultaneously with SMX or DDS and incubated for 3 h (data not shown). We also found that trolenadomycin (a potent inhibitor of CYP3A4) and disulfiram (CYP2E1 inhibitor) did not attenuate the protein haptentation with either DDS or SMX (data not shown).
CYP2C9 message in either cell type (Fig. 4). Yengi et al. (2003) have previously reported the expression of CYP2C9, CYP2C18, and CYP2C19 transcripts in human skin samples. Because these investigators probed full thickness skin, it is not possible to determine which skin cells gave rise to these transcripts. Saeki et al. (2002) have observed the presence of CYP2C transcripts in human keratinocytes from several subjects. However, their use of nonspecific primers does not permit identification of the specific CYP2C genes that were expressed in these cells. Moreover, transcripts can be detected in the absence of detectable protein. Although our data suggest the possible presence of a low level of CYP2C9 protein in NHEKs, further studies are needed to more carefully determine the expression of this enzyme in human skin.

The results shown in Fig. 6 suggest that CYP3A5 is the protein detected in both NHEKs and HaCaT cells because it was detected by an antibody that is immunospecific for CYP3A5. Moreover, CYP3A5 mRNA was readily amplified from NHEKs and HaCaT cells, whereas a second round of PCR amplification was required to detect CYP3A4 mRNA transcript in NHEK cells (Fig. 6B). To our knowledge, this differential expression of CYP3A enzymes has not been demonstrated previously in keratinocytes. The differential expression of CYP3A4/5 proteins, especially much lower expression in NHEKs and HaCaT cells as compared with liver, might suggest the minor role of these enzymes for the bioactivation of these parent drugs in skin. Interestingly, we were unable to demonstrate the metabolism of a CYP3A4 substrate (7-BQ) in NHEKs, although its metabolite (7-HQ) was readily detected when added directly to NHEKs. This suggests a very low level of CYP3A4-mediated catalytic activity.

**Fig. 8.** Protein haptenation of DDS in NHEKs in the presence of inhibitors of CYP450s and cyclooxygenase. NHEKs were incubated for 3 h in the presence of vehicle (1% DMSO), 5 mM ABT, 5 mM DCE, or 100 µM INDO. The concentrations of inhibitors selected were the maximal concentrations that did not increase cell death in NHEKs under the incubation conditions. After preincubation with inhibitors, cells were incubated for an additional 3 h with 250 µM DDS. Covalent adducts were quantified using confocal microscopy (A) or ELISA (B), as described under Materials and Methods. A, quantification of adducts was performed as described under Materials and Methods. Control, NHEKs incubated with vehicle (1% DMSO) alone. Results were analyzed using ANOVA with Holm-Sidak method for multiple pair-wise comparisons. *, p < 0.05 compared with NHEKs incubated with vehicle alone. B, covalent adducts were determined by the ELISA as described under Materials and Methods. Data presented represent the mean (S.D.) optical density of three different experiments having three replicates in each experiment. Data were analyzed statistically using ANOVA with the Holm-Sidak test for multiple pair-wise comparisons. *, p < 0.05 compared with NHEKs incubated with vehicle alone.

**Fig. 9.** Protein haptenation of SMX in NHEKs in the presence of inhibitors of CYP450s and cyclooxygenase. NHEKs were incubated for 3 h in the presence of vehicle (1% DMSO), 5 mM ABT, 5 mM DCE, or 100 µM INDO. The concentrations of inhibitors selected were the maximal concentrations that did not increase cell death in NHEKs under the incubation conditions. After preincubation with inhibitors, cells were incubated for an additional 3 h with 250 µM SMX. Covalent adducts were quantified using ELISA as described under Materials and Methods. Data presented represent the mean (S.D.) optical density of three different experiments having three replicates in each experiment. Data were analyzed statistically using ANOVA with the Holm-Sidak test for multiple pair-wise comparisons. *, p < 0.05 compared with NHEKs incubated with vehicle alone.
in these cells. We were also able to demonstrate the presence of CYP2E1 mRNA in both NHEKs and HaCaT cell types (Fig. 7). This is consistent with the reports of other investigators who have demonstrated the presence of CYP2E1 transcripts and protein in NHEKs (Baron et al., 2001; Saeki et al., 2002).

Based on the mRNA and protein data, we concluded that CYP2C9 was not a likely mediator of the bioactivation of these drugs in NHEKs. Indeed, preliminary studies demonstrated that sulfinaphenazine, an inhibitor of CYP2C9, did not inhibit protein haptenation in NHEKs exposed to SMX or DDS (Wurster et al., 2004). To evaluate the role of CYP3A4/5 and CYP2E1 in the generation of drug-protein adducts, we evaluated the effect of inhibitors of these enzymes (at their maximum noncytotoxic concentration) on protein haptenation in NHEKs exposed to SMX or DDS. Our results indicate that a general inhibitor of CYP450, ABT (which has been shown to completely (~90%) inhibit most of the major CYP450, such as CYP1A2, 2B6, 2C9, 2C19, 2D6, and 3A4) (Balani et al., 2002), did not reduce the covalent adducts formed after exposure of NHEKs to DDS and SMX (Figs. 8 and 9). The selective inhibitor of CYP2E1 (DCE) also failed to decrease adduct formation. Other inhibitors of CYP3A4 (troleandomycin) and CYP2E1 (disulfiram) also did not reduce the protein haptenation of DDS and SMX in NHEKs (data not shown). These results suggest that CYP450 does not play a major role in the oxidative metabolism of these drugs in NHEKs. In addition, although COX-2 has been shown to mediate the oxidation of arylamines in vitro, an inhibitor of cyclooxygenase (indomethacin) did not attenuate the protein haptenation in NHEKs exposed to either DDS or SMX (Figs. 8 and 9). This is consistent with our recent observation that recombinant COX-2 does not mediate the oxidation of these drugs (Vyas et al., 2006).

Hence, our studies confirm our previous observations that NHEKs are able to bioactivate SMX and DDS, giving rise to haptenated proteins. We found that CYP450 that are important in the bioactivation of these drugs in the liver do not appear to play a significant role in their bioactivation in NHEKs. It should be noted, however, that recent data suggests the level of CYP450 expression may differ as keratinocytes differentiate (Du et al., 2006). Nevertheless, we observed significant protein haptenation in NHEKs in the absence of evidence for involvement of CYP450. These studies demonstrate that the role of various drug-metabolizing enzymes in the bioactivation of drugs may vary from tissue to tissue. Likewise, although COX-2 has been reported to oxidize arylamines, neither COX-1 nor COX-2 appears to contribute to the bioactivation of these drugs in NHEKs. In our companion paper to this current investigation, we demonstrated the role of flavin-containing monooxygenase and peroxidases in the bioactivation of these arylamine drugs in keratinocytes.

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