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Differential cellular metabolite alterations in HaCaT cells caused by exposure to the aryl hydrocarbon receptor-binding polycyclic aromatic hydrocarbons chrysene, benzo[a]pyrene and dibenzo[a,l]pyrene

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A B S T R A C T

Polycyclic aromatic hydrocarbons (PAHs) are ubiquitous in the human environment. Since they are present in crude oil fractions used for the production of rubber and plastics, consumers may come into direct dermal contacts with these compounds (e.g., via tool handles) on a daily basis. Some individual PAHs are identified as genotoxic mutagens thereby prompting particular toxicological and environmental concern. Among this group, benzo[a]pyrene (BAP) constitutes a model carcinogen which is also used as reference compound for risk assessment purposes. It acts as a strong agonist of the aryl hydrocarbon receptor (AHR) and becomes metabolically activated toward mutagenic and carcinogenic intermediates by cytochrome P450-dependent monoxygenases (CYPs). While BAP has been exhaustively characterized with regard to its toxicological properties, there is much less information available for other PAHs. We treated an AHR-proficient immortal human keratinocyte cell line (i.e., HaCaT) with three selected PAHs: BAP, chrysene (CRY) and dibenzo[a,l]pyrene (DALP). Compound-mediated alterations of endogenous metabolites were investigated by an LC–MS/MS-based targeted approach. To examine AHR-dependent changes of the measured metabolites, AHR-deficient HaCaT knockdown cells (AHR-KD) were used for comparison. Our results reveal that 24 metabolites are sufficient to separate the PAH-exposed cells from untreated controls by application of a multivariate model. Alterations in the metabolomics profiles caused by each PAH show influences on the energy and lipid metabolism of the cells indicating reduced tricarboxylic acid (TCA) cycle activity and β-oxidation. Up-regulation of sphingomyelin levels after exposure to BAP and DALP point to pro-apoptotic processes caused by these two potent PAHs. Our results suggest that in vitro metabolomics can serve as tool to develop bioassays for application in hazard assessment.

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1. Introduction

In daily life, consumers are exposed to a variety of potentially toxic compounds that may be released from consumer products [1,2]. For instance, polycyclic aromatic hydrocarbons (PAHs) are ubiquitous in the human environment. Generated from incomplete combustion of organic matter like coal, grilled meat or cigarettes they are present in air, food and soil [3–5]. Furthermore, PAHs occur in crude oil fractions and carbon black used for the production of rubber, certain plastics and elastomers [6,7]. Dermal contact with consumer products containing petroleum and subsequent dermal uptake of diesel oil and other lubricants were identified in human volunteer trial experiments [8]. Therefore, direct skin contact (e.g., via tool handles or lubricants and fuels) in combination with possible migration is supposed to be an important route of exposure
for various PAHs and percutaneous absorption of PAHs has been shown before [9–11]. Moreover, a strong correlation between dermal exposure to PAHs and the occurrence of skin cancer has been reported in the past [12].

Carcinogenic PAHs require activation through biotransformation to form reactive intermediates and to target biological macromolecules such as DNA [5]. Cytochrome P450-dependent monooxygenases (CYPs), in particular CYP1A1, 1A2 and 1B1, are primarily involved in the metabolism of PAHs [13,14]. Enzymatic generation of sterically hindered bay or fjord region diol-epoxides and subsequent DNA adduct generation is considered the dominant mechanism in chemical carcinogenesis resulting in mutational miscoding [15]. Many bay region PAHs such as benzo[a]pyrene (BAP) are activated through CYP1A1 [16]. By contrast, CYP1B1 seems to play a more important role in the metabolism of fjord region PAHs such as dibenzo[a,l]pyrene (DALP; IUPAC notation: dibenzo[def,fp]chrysene) [17]. Besides activation through CYP enzymes, two other activation pathways are important in PAH metabolism: the formation of radical cations by one electron oxidation via peroxides [18], and the formation of o-quinones by dihydrodiol dehydrogenases [19]. It is assumed that all three pathways—to different extents—might contribute to PAH-mediated carcinogenesis [20]. Most PAHs act as activators of the aryl hydrocarbon receptor (AHR) [21]. AHR activation then leads to gene expression of metabolizing enzymes such as CYPs, which—in turn—contribute to the formation of activated PAH metabolites and subsequent DNA damage. Besides its role in the enzymatic biotransformation of xenobiotics, the AHR may also contribute more directly to alterations in cell signaling and proliferation [22]. The complexity of PAH toxicity is further illustrated by the fact that the level of DNA adduct formation does not necessarily correlate with the extent of tumor formation in skin [23,24].

BAP was classified as human carcinogen by IARC [25]. By contrast, chrysene (CRY) has less carcinogenic activity in comparison to BAP. It also contains a bay region that can be converted into reactive diol-epoxide moieties though [26], which then may induce cancer in mice [27,28]. DALP is one of the strongest carcinogenic PAHs containing a fjord region [29]. The described toxicological mechanisms triggered by different PAHs might lead to interactions and non-additive effects if such compounds occur in mixtures [30]. Hence risk assessment strategies for exposures to PAH mixtures need to be developed.

The three selected PAHs applied in our study are present in the environment and may therefore come into contact with human skin. All three PAHs possess different chemical structures and carcinogenic potencies, but all of them are metabolized by CYP enzymes. We used an immortal human keratinocyte cell line (HaCaT) as an in vitro model to study the effects that occur upon exposure to these PAHs. HaCaT cells have been shown to be a suitable metabolically competent model system to mimic human skin cells [31]. We examined the effects of the three selected PAHs using a targeted metabolomics approach by analyzing 188 endogenous metabolites. Thereby we analyzed the end-products of cellular metabolism via comparative analysis of endogenous metabolites and examined AHR-dependent changes induced in HaCaT cells. The overall aim of our studies is to gain more molecular information on the adverse signaling pathways associated with dermal exposures to PAHs.

## 2. Materials and methods

### 2.1. Chemicals

Cell culture media and supplements were purchased from PAN Biotech (Aidenbach, Germany). All other chemicals were purchased from Sigma Aldrich with the highest purity available (Munich, Germany). Substances were routinely dissolved as stock solutions in dimethylsulfoxide (DMSO).

### 2.2. Cell lines

Human adult wild-type (HaCaT WT) and AHR knockdown (HaCaT AHR-KD) keratinocyte cell lines were kindly provided by Prof. Ellen Fritsche (IFU – Leibniz Research Institute for Environmental Medicine, Germany) and are described elsewhere [32]. Both cell lines were maintained in DMEM (Dulbecco’s Modified Eagle Medium supplemented with 10% fetal calf serum (FCS, v/v), 2 mM L-glutamine, penicillin (100 U/mL), streptomycin (0.1 mg/mL) at 37 °C in humidified atmosphere with 5% CO₂. After 24 h of initial incubation the cells were exposed to the test substances.

### 2.3. MTT assay

In brief, cells (2 × 10⁵ cells/mL) were seeded in 24-well plates and exposed to various concentrations (0–20 μM) of the chemicals (BAP, CRY, DALP). The final concentration of DMSO did not exceed 0.1%(v/v). After 24 h and 48 h of exposure, MTT reagent (0.5 mg/mL) was added and incubated for 2 h at 37 °C, 5% CO₂. Afterwards, formazan crystals were dissolved by adding 400 μL DMSO into each well. The absorption of the samples was measured in triplicates on a microplate TECAN reader at 595 nm.

### 2.4. Real-time PCR

For gene expression studies cells (2 × 10⁵ cells/mL) were seeded in 12-well plates and incubated in DMEM. Cells were exposed to the test substances for 24 h or 48 h, respectively. Following substance treatment the total RNA was extracted using Trizol (Invitrogen, Carlsbad, CA, USA). RNA quantity and quality was determined using a Nanodrop-1000 (Thermo Fisher Scientific, Asheville, NC, USA). Subsequently, cDNA was synthesized of 1 μg of RNA using High Capacity cDNA Reverse Transcription Kit (Applied Biosystems, Carlsbad, CA, USA). Levels of individual transcripts were then quantified using SYBR®Green Fast (Applied Biosystems). Quantitative real-time polymerase chain reaction (qRT-PCR) was performed according to the manufacturer’s instructions on an ABI-7500 Fast Real Time PCR system (Applied Biosystems). Gene selective primers for CYP1A1 (forward 5′-CCAGAATCCACCTTCCCAGCCT-3′, reverse 5′-GAGGGCGCAAGAAATCCCTG-GGC-3′), CYP1B1 (forward 5′-TGGATTTGAGAAGCTACC-3′, reverse 5′-CCAGG-ACCTGATCATAATTC-3′) and HPRT (forward 5′-GCTCTGAGGACGTCGG-3′, reverse 5′-GCCAAAAGGAGTGAATG-3′) were used. Relative gene expression was calculated using the ΔΔCt method and normalized to expression levels of HPRT. Vehicle control (0.1% DMSO) sample expressions were set to a fold change value of 1 and used as comparison for treated cells.

### 2.5. Western blots

Equal amounts of proteins (30 μg) obtained from RIPA buffer lysed cells were applied to SDS-PAGE, transferred onto nitrocellulose membranes and immune-blotted according to the manufacturer’s instructions. Antibodies against CYP1A1 (sc-20772) and CYP1B1 (sc-32882) were used (Santa Cruz Biotechnology, Santa Cruz, CA, USA). Primary antibody probed blots were visualized with appropriate horseradish peroxidase-coupled secondary antibodies (Santa Cruz Biotechnology) using enhanced chemo-luminescence (34078; Thermo Scientific, Waltham, MA, USA) for detection.
2.6. Cell culture for metabolomics approach

Cells (1 × 10^5 cells/mL) were seeded in 10 cm cell culture dishes and incubated in DMEM. Cells were exposed to the test substances (4 replicates for each treatment) for 48 h. After substance treatment, cells were quickly washed twice with ice-cold ammonium formate buffer solution (155 mM, pH 7.4). Cells were detached from the culture dish by scraping into 1 mL of buffer solution, transferred to 1.5 mL safe-lock tubes (Eppendorf, Hamburg, Germany) and centrifuged (500 × g, 5 min, 4 °C). After centrifugation the supernatant was removed and the cell pellet weight was recorded before freezing them in liquid nitrogen. The frozen samples were stored until analysis at −80 °C A volume of 0.5 mL UHPLC grade ice-cold methanol (J.T. Baker, Deventer, Holland) was added for cell pellet extraction followed by 3 freeze and thaw cycles: cells were frozen for 5 min in liquid nitrogen and afterwards thawed for 5 min at room temperature followed by 1 min of ultrasonication in a water bath. After centrifugation (12,000 x g, 30 min, 4 °C), supernatants were immediately analyzed using the Biocrates AbsoluteIDQ p180 kit.

2.7. Targeted metabolomics approach

The AbsoluteIDQ platform (Kit p180, Biocrates, Innsbruck, Austria) was employed for targeted metabolite profiling as described by the manufacturer. This platform detects a total of 188 metabolites, including 42 amino acids (AAs) and biogenic amines (BAs), 40 acylcarnitines (Cx:y), hydroxycarboxylicarnitines (Cx:y-OH) and dicarboxyphosphatidylcholines (PC:Cx:y) (ACs), the sum of hexoses, 15 sphingomyelins (SM:Cx:y) and sphingomyelin derivatives [SM(OH)Cx:y], as well as 14 lysophosphatidylcholines (LPC:Cx:y) and 76 phosphatidylcholines (PC:Cx:y). LPCs and PCs were further differentiated with respect to the presence of ester (“a”) or ether (“e”) bonds in the glycerol moiety (“a” = acyl, “aa” = diacyl, and “ee” = acyl-alkyl). The lipid side chain composition is abbreviated with “Cx:y”; “x” denotes the number of carbons in the side chain and “y” the number of double bonds. Detailed information on the metabolites analyzed along with the protocol of metabolite quantification is presented elsewhere [33]. The cellular extracts were prepared following the manufacturer’s protocol. Briefly, samples were prepared as follows: (a) pipetting of a standard mixture labeled with stable isotopes onto the filter paper inserts of the 96-well plate of the kit and drying under a stream of nitrogen using an EVA LS evaporator (VLM, Bielefeld, Germany), (b) adding of 20 μL cell extract onto the filter inserts and drying under a stream of nitrogen, (c) derivatization of AAs using phenylisothiocyanate, (d) extraction of metabolites and internal standards with 5 mL ammonium acetate in methanol, (e) centrifugation through a membrane filter followed by dilution with solvent for mass spectrometry. One part of the eluate was diluted with 40% methanol in water (1/10 (v/v)) for LC–MS/MS analysis, the other part with Biocrates kit running solvent (1/50 (v/v)) for FIA-MS/MS. The final extracts were then analyzed using a QTRAP 5500 (AB Sciex, Darmstadt, Germany) triple quadrupole mass spectrometer coupled to an Agilent 1200 Series HPLC (Agilent, Waldbronn, Germany) using ESI MS/MS. For the mass spectrometry analysis the software package Analyst 1.6 was used (AB Sciex, Darmstadt, Germany). LC–MS/MS analysis was performed in the positive mode which enables the quantification of 42 metabolites, covering 21 AAs and 21 BAs. A ZorbaX Eclipse XDB-C18 column (3 × 100 mm, 3.5 μm; Agilent, Waldbronn, Germany) was used for metabolite separation by LC. Subsequently, the FIA-MS/MS procedure was used for the quantification of 146 additional metabolites, covering 40 ACs including free carnitine, 38 PCAA’s, 38 PCa’E’s, 14 LPCs, 15 SMs, and the sum of hexoses in both positive and negative ion mode. Multiple reaction monitoring (MRM) was used for quantification and compound identification using the MetIDQ™ software package which is an integral part of the AbsoluteIDQ kit. Concentrations of all metabolites were calculated in micromolar units. The method is in concordance with FDA guidelines, warranting reproducibility within a given error range [34]. Metabolite concentrations were standardized to cell pellet weight. For statistical analyses, only metabolites were chosen which were present in each sample, thus restricting the profile to a total of 166 metabolites.

2.8. Statistical analysis

Processing of metabolomics data was carried out using the statistical package SPSS* (version 12.0.2G) as described elsewhere [35]. In brief, each sample was standardized to the mean of the control (set to 100%) and normalized using z-score values. Then, a MANOVA (multivariate analysis of variance) including Bonferroni correction was used for compound selection and reduction of the dataset to metabolites that are significantly changed in at least one exposure group compared to controls. Metabolites that showed a significant group difference (p < 0.05) were selected for multivariate analysis. Principal component analysis (PCA) was performed to achieve dimension reduction of the datasets and the resulting factors were used for a post-hoc Fisher’s Discriminant analysis as described elsewhere [36] for group separation. The performance of the discriminant models was subsequently verified by applying the “leave-one-out” cross-validation formalism [37].

3. Results

Prior to the metabolomics analysis, dose-dependent cytotoxicity was investigated in HaCaT WT and AHR-KD cells upon exposure to each of the three PAHs (Fig. 1).

To indirectly assess AHR activation by these compounds, induction of the downstream genes CYP1A1 and CYP1B1 was examined at different concentrations and at both the mRNA (Fig. 2) and the protein level (Fig. 3).

After selection of appropriate concentrations and time-points corresponding cellular metabolites were analyzed using a targeted metabolomics platform.

3.1. Cytotoxicity

Cell viability was only weakly affected in both cell lines after treatment with up to 20 μM BAP for 48 h (Fig. 1A). In HaCaT WT cells viability only decreased to about 80% at the highest concentration applied. In contrast, CRY caused strong cytotoxic effects in both cell lines (calculated EC50 values of 3.8 μM for WT and 2.0 μM for AHR-KD cells; Fig. 1B). Treatment with DALP also led to enhanced cytotoxicity in WT cells with a calculated EC50 value of 0.035 μM (Fig. 1C). On the other side, the viability of AHR-KD cells remained unaffected at DALP concentrations of up to 20 μM.

3.2. CYP induction on mRNA level

To investigate the activation of AHR by BAP, CRY or DALP, induction of the downstream genes CYP1A1 and CYP1B1 was measured at the mRNA level. Both CYP enzymes are involved in the metabolism of the three PAHs [13]. As expected, BAP caused a very strong and concentration-dependent induction of CYP1A1 in WT but only a slight elevation of transcription levels in AHR-KD cells after 24 h of incubation (Fig. 2A). For CYP1B1 induction a clear chemical-related effect could be observed as well albeit to a much lower extent. Compared to BAP, CRY was also found inducing both genes in WT cells but less strongly. There was again no significant induction in AHR-KD cells (Fig. 2B). Conversely, DALP only weakly increased
the transcription level of CYP1A1 and those of CYP1B1 were even less affected (Fig. 2C). Although this finding is matched by another study where DALP showed similar results in the skin of mice [24], it nevertheless seems kind of surprising and also pointed to the possibility that the time-points picked might have been too late to assess this end-point adequately. Therefore, we also tested earlier time-points (2 and 8 h) and confirmed that no significant induction of CYP1A1 or CYP1B1 occurred upon treatment of HaCaT cells with DALP concentrations of up to 1 μM (data not shown).

3.3. CYP induction on protein level

Western blots confirmed the qPCR data on the protein level. Both CYP1A1 and CYP1B1 showed a strong concentration-dependent induction by BAP, a moderate induction by CRY and no induction by DALP in HaCaT WT cells (Fig. 3). For AHR-KD cells faint signals were detected for BAP only.

Based on the results obtained in cytotoxicity tests, qPCR and western blots, concentrations of 1 μM (BAP and CRY) and 0.01 μM (DALP) were selected to be applied in the subsequent metabolomics study. HaCaT cells (WT and AHR-KD) were exposed to each chemical for 48 h. This experimental setting uses a sub-toxic dose for each PAH sufficient to activate the AHR in HaCaT WT cells. Exposure for 48 h was regarded to be suitable for the detection of metabolomic alterations. So, this time-point represents a scenario after maximal AHR activation and CYP mRNA induction at which downstream CYP protein levels were still found elevated.

3.4. Metabolomics approach: identification of metabolomics biomarkers

In total, 166 metabolites were detected at all exposure scenarios and analyzed using the AbsoluteIDQ platform. For HaCaT WT and AHR-KD cells 126 and 45 metabolites, respectively, were selected for a multivariate model since they showed significant group differences (p ≤ 0.05). PCA was performed to achieve dimension reduction of the datasets and the resulting factors were used for a post-hoc Fisher’s discriminant analysis resulting in group separation as described elsewhere [36]. For HaCaT WT cells, this discriminant model resulted in 6 factors accounting for 100% of the observed variance in the system. Evaluation of the PCA factors that contributed most to discriminant functions F(1×1) and F(2×2) revealed a high contribution (90% explanation of the overall variance observed) of PCA factors 1–3 to the separation. Interestingly, mainly PCae’s and PCaa’s were loading high on factor 1, acylcar- nitines on factor 2 and a mix of metabolite classes including the amino acid glutamine (Gln) and α-aminoacidic acid (alpha-AAA) were loading high on factor 3. For further data reduction the following thresholds were applied to the loading factors: for factor 1 ≥ 0.970, for factor 2 ≥ 0.950 and for factor 3 ≥ 0.330, respectively. This methodology revealed 24 metabolites (see Table 1) that could be used to achieve complete separation of all exposure groups and controls in the model applied (Fig. 4).

The performance of the discriminant models was subsequently verified by applying the “leave-one-out” cross-validation formalism [37,38] resulting in 100% accuracy. In the case of HaCaT AHR-KD cells, Fisher’s discriminant analysis of the selected 45 metabolites led to a separation of the exposure groups by including 7 factors, but cross-validation revealed only 81.3% of accuracy. One case of each exposure group was falsely assigned: one BAP-exposed sample was predicted as control, one CRY-treated sample as DALP-treated sample and one DALP-treated sample as CRY-treated sample. Accordingly, no reliable separation of exposure groups in HaCaT AHR-KD cells could be achieved. In HaCaT WT cells, the 24 identified metabolites that were used in the multivariate model to distinguish between different treatment groups comprise 6 different metabolite classes (BAs, AAs, ACS, PCaa’s, PCae’s and SMs, see Tbl. 1): the two polar metabolites alpha-AAA (BAs) and the amino acid Gln, two PCaa’s of medium chain length (i.e., PCaa_C32:2 and PCaa_C36:2), several PCae’s of medium and long chain length as well as SM_C16:1. Scatter dot plots (Fig. 4) show examples of metabolites of each class providing insight in the metabolomics patterns altered by the three PAHs that were essential for group separation. All PAHs show a down-regulation of Gln. Furthermore, metabolomics patterns of BAP and DALP show similarities in a strong down-regulation of alpha-AAA and a strong up-regulation of
Fig. 2. Effect of BAP, CRY and DALP on the AHR target genes CYP1A1 and CYP1B1 in HaCaT WT and AHR-KD cells. Cells were exposed for 24 h and 48 h to (A) BAP (0.1 μM, 1.0 μM, 3.5 μM), (B) CRY (0.1 μM, 1.0 μM, 3.5 μM), and (C) DALP (0.01 μM, 0.02 μM, 0.03 μM), respectively. Controls contained 0.1% DMSO (vehicle) only. Relative gene expression of CYP1A1 and CYP1B1 was subsequently quantified with SYBR Green using real-time PCR. Target gene expression was normalized to the transcription level of an endogenous gene (HPRT). The data represent the mean of two independent experiments (±SD).

3.5. Metabolomics approach: alterations of metabolomics profiles by each PAH

We used the described multivariate model to examine which metabolites contribute to the separation of the three PAHs. In an additional statistical analysis we evaluated alterations, as caused by each PAH in comparison to non-treated controls, of the complete metabolomics pattern consisting of a total of 166 metabolites. In Venn diagrams and volcano plots the metabolomics changes are visualized (Fig. 5).

The plots depict the effects on the measured metabolites caused by each PAH in HaCaT WT cells. Venn diagrams (Fig. 5A) summarizing all significant changes show the differential alterations caused by the three compounds. Most of the changes were noticed as up-regulations of the levels of metabolites mainly caused by the two potent carcinogens BAP and DALP. These two compounds share a third of all up-regulated metabolites. Still 26 metabolites are uniquely elevated upon BAP exposure. BAP is also sharing certain up-regulated metabolites with CRY, while DALP and CRY do not have any metabolite changes in common. Down-regulation of metabolites was mainly caused by DALP with 30 uniquely altered metabolites compared to none by BAP and CRY. Only Glu has been consistently found down-regulated in all treatment groups (BAP p = 0.001, CRY p = 0.008, DALP p = 0.002). Therefore we verified relative Glu levels by an enzymatic assay. As seen in the
targeted metabolomics approach, Gln was downregulated approximately two-fold in HaCaT WT cells treated with BAP and DALP (Supplementary Fig. 4). However, this orthogonal method could not confirm the effect of CRY on Gln levels. In AHR-KD cells, only BAP provoked a decrease of Gln concentration in accordance with the LC–MS/MS measurements. Metabolomics profile changes caused by BAP (Fig. 5B) and DALP (Fig. 5D) show similarities in up-regulation of ACs (e.g., C18:2 BAP \( p = 0.01 \); DALP \( p \leq 0.003 \)), SMs (e.g., SM_C26:0; BAP \( p = 0.025 \); DALP \( p = 0.009 \)) and LPCs (e.g., LPC_C14:0; BAP \( p = 0.008 \); DALP \( p = 0.007 \)), as well as down-regulation of AAs (e.g., Lys; BAP \( p = 0.034 \); DALP \( p \leq 0.0005 \)) and one BA (i.e., alpha-AAA; BAP \( p = 0.004 \); DALP \( p \leq 0.0005 \)) (Supplementary Tbl. 1). Treatment with BAP specifically induces a significant up-regulation of 9 SMs and SM(OH)\(_2\), for example SM(OH)\(_2\)_C14:1 \( p = 0.006 \) and SM_C24:0 \( p = 0.009 \), which are not altered in DALP- or CRY-treated cells (Supplementary Tbl. 1). In the case of DALP 37 ACs are significantly up-regulated. Twenty eight among them are also found up-regulated in cells treated with BAP. Conversely, treatment of WT cells with CRY did not affect its AC levels. The treatment of cells with DALP is further down-regulating 11 PCa\(_e\)'s that remain unaffected upon exposure to BAP and CRY. Twenty PCa\(_a\)'s were found increased in cells after BAP exposure, 12 of which are also up-regulated after CRY exposure. Treatment with DALP caused a strong down-regulation of 15 AAs and 3 BAs (i.e., alpha-AAA, sarcosine,
metabolites using SMs OH concentrations taurine). By contrast, treatment of cells with CRY and BAP resulted in the down-regulation of only 1 (Gln) and 3 (Ala, Gln and Lys) AAs, respectively. In CRY-exposed cells, many PCaa’s and PCae’s were up-regulated similarly to BAP-exposed cells. On the other side, SM, AC, LPC and BA concentrations remained completely unaffected in HaCaT WT cells upon treatment with CRY when compared to control cells. The results obtained for AHR-KD cells are presented in Supplementary Fig. 1 and Supplementary Tbl. 2.

### 4. Discussion

Carcinogenic PAHs occur in crude oil fractions used for the production of rubber, certain plastics and elastomers. Since keratinocytes are potential target cells for PAH exposure we studied the influence of BAP, CRY and DALP on 166 endogenous metabolites of HaCaT cells. HaCaT cells represent a non-tumorigenic permanent epithelial cell line from adult human skin [39] which
is known to be metabolically competent [40]. Hence, exposure of these cells against PAHs may lead to PAH biotransformation and the activation of certain toxicological signaling cascades involved in the expression of an adverse outcome.

4.1. PAH-mediated cytotoxicity and influence on CYP1A1 and CYP1B1 expression

Even though BAP is metabolized in HaCaT WT cells to its ultimate carcinogen (−)-anti-BAP-7,8-diol-9,10-epoxide (BPDE) [40] and has been shown to cause metabolism-independent pro-apoptotic effects [41], viability tests revealed only little cytotoxicity of BAP in HaCaT WT or AHR-KD cells after 48 h at the concentrations applied. Compared to the other two PAHs investigated, BAP caused a strong induction of CYP1A1 and CYP1B1 on both mRNA and protein level (Figs. 1 and 2). This is in accordance with its high potency to bind and activate the AHR [42]. However, besides CYP induction, AHR activation may also promote defensive mechanisms such as, e.g., the synthesis of glutathione S-transferases (GST) [5], which then—in turn—may detoxify BAP-derived electrophiles such as BPDE [43]. This mechanism might contribute to the high tolerance of HaCaT cells against BAP. On the other side, the tetracyclic PAH CRY revealed with some cytotoxicity in HaCaT cells irrespective of its AHR expression status and most likely due to the induction of oxidative stress [44]. Conversely, DALP showed very strong cytotoxicity only in HaCaT WT cells (EC50: approx. 0.035 μM) without inducing CYP1A1 or CYP1B1. Since the strong cytotoxicity of DALP was observed in HaCaT WT without significant induction of CYP enzymes, but completely disappeared in HaCaT AHR-KD cells, AHR-dependent toxicity of DALP in this particular cell line seems rather mediated through other mechanisms than diol-epoxide-DNA adduct formation. Notwithstanding, formation of DALP-11,12-diol-13,14-epoxides (DALPDE) has been

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**Fig. 5.** Differential alterations of metabolomics profiles caused by BAP, CRY and DALP in HaCaT WT cells. Venn diagrams of up- and down-regulated levels of metabolites (A). Volcano plots of metabolites influenced by each single PAH compared to control: BAP (B), CRY (C) and DALP (D). Fold changes were calculated as the ratio between the median of the two groups compared. The significance expressed as p-values is calculated by a two-sided, unpaired t-test. (B–D) The x-axes represent the fold changes as log2 values of the ratio between the median of the treated cells compared to untreated controls. The y-axes represent the −log10 values of the determined p-values via t-test. Axes on the right side represent the p-values (p < 0.05 is regarded as significant). On the top axis, fold changes are presented. Abbreviations used: AAs, amino acids; ACs, acylcarnitines; BAs, biogenic amines; PCs, phosphatidylcholines; LPCs, lysophosphatidylcholines; SMs, sphingomyelins.
already shown to contribute to DALP-mediated cytotoxicity in certain cell models such as mouse embryo fibroblasts [45].

4.2. Effects of BAP, CRY and DALP on metabolites of HaCaT WT and AHR-KD cells

For the first time a metabolomics analysis was conducted in HaCaT cells and the differential influences of the three selected PAHs were recorded and considered as individual signatures of toxicity. Applying a multivariate model, the 166 metabolites analyzed allowed no separation of the different exposure groups in HaCaT AHR-KD cells while HaCaT WT cells revealed strongly altered and differentiable metabolomics profiles (Figs. 4 and 5). This result points to differences of AHR signaling dependent on the particular PAH used. Since CYP1A1 and CYP1B1 expression was found significantly increased in WT cells treated with BAP and CRY (Figs. 2 and 3), metabolites formed through enzyme-mediated catalysis most likely contributed to the overall metabolomics alterations observed with these two PAHs. On the other hand, DALP-treated WT cells showed a lack of considerable CYP induction. In addition, metabolite levels remained virtually unaffected in AHR knockdown cells treated with DALP (Supplementary Fig. 1). Aiming to explain these results for DALP, we considered three possible reasons: Firstly, AHR-KD cells might become insensitive against PAH effects and differentiate differently due to the lack of AHR. Contrasting this assumption, however, on the metabolite levels only minor changes between AHR-KD cells and WT cells could be detected in untreated controls (Supplementary Fig. 2). Secondly, albeit extremely low, basal CYP1B1 levels in WT cells could be sufficient for the formation of genotoxic metabolites of DALP. We detected 4-fold lower basal CYP mRNA levels in AHR-KD cells compared to WT cells (Supplementary Fig. 3). Also Topinka et al. found a negligible CYP induction at DALP concentrations that nevertheless caused high levels of DNA adducts [46]. A 10,000 fold reduction of basal CYP1B1 levels has been shown after knockout of AHR in MCF-7 cells [47]. As CYP1B1 is mainly responsible for DALP’s metabolic activation [48], a decrease in the cellular level of this enzyme might explain the insensitivity of AHR-KD cells against the potential toxicity of this compound. Thirdly, the AHR-dependent toxicity of DALP could also be caused by non-genotoxic AHR signaling alone. Although not investigated yet DALP might function as a selective AHR modulator (SAHRM), thereby triggering alternative AHR responses without xenobiotic response element (XRE) activation [49].

4.3. Alterations in metabolite levels of amino acids and biogenic amines

The levels of the amino acid Gln were found strongly lowered upon exposure of HaCaT WT cells to BAP and DALP and weakly upon CRY exposure (Figs. 4 and 5). Conversely, in AHR-KD cells Gln levels were only decreased compared to controls upon treatment with BAP, but not in cells treated with CRY or DALP (Supplementary Figs. 1, 4). Significantly depleted levels of Gln have been reported in the literature in lung-derived A549 cells treated with different PAH quinones [50]. Since BAP quinones can easily be generated via both CYP-mediated monoxygenation followed by non-enzymatic oxidation and peroxidase-mediated one-electron oxidation [5], in the absence of a functional AHR the contribution of the latter pathway might become more relevant in the generation of PAH quinones.

The strong depletion in the levels of Gln points to a PAH-mediated impairment of both glycolysis and the tricarboxylic acid (TCA) cycle whose intermediates provide anabolic precursors for the biosynthesis of fatty acids, nucleic acids, and proteins. This is further supported by significantly reduced levels of alpha-AAA, an intermediate in lysine biosynthesis, which has also been found depleted in HaCaT cells after exposure to BAP or DALP (Fig. 5, Supplementary Tbl. 1). Altogether, the levels of 15 AAs were decreased after DALP exposure (Supplementary Tbl. 1). Three of them (Ala, Gln and Lys) were also lowered in HaCaT cells after exposure to BAP. In particular, Gln and Gly are necessary in the biosynthesis of glutathione (GSH), a tripeptide which acts as protective agent against reactive PAH metabolites such as DALPDE and thus can prevent DNA adduct formation [43]. Therefore, depletion of these AAs could be also due to enhanced GSH synthesis.

4.4. Alterations in the lipid metabolite profile caused by the three different PAHs

Upon conversion of long-chain acyl-CoAs into ACs, these products are actively transported across the inner mitochondrial membrane in exchange for free carnitine. Then, through catalysis of carnitine acyltransferase II, acyl-CoA is being recovered within the mitochondrial matrix and directed towards β-oxidation [51]. Recently, Westmann et al. showed that fatty acid β-oxidation in chicken embryo liver was significantly reduced in response to the treatment with a mixture of 16 different PAHs [52]. In transgenic mice activation of AHR inhibited hepatic fatty acid β-oxidation along with acyl-CoA oxidase 1, the rate limiting enzyme in peroxisomal β-oxidation [53]. Down-regulation of acyl-CoA synthetase was shown after BAP exposure of marine diatoms, thus providing evidence that PAHs may interfere with the regulation of lipid metabolism [54]. In our experiments significant increases of AC levels have been detected in HaCaT WT cells after exposure to BAP and DALP (Fig. 5, Supplementary Tbl. 1). A more than 3-fold increase of carnitine palmitoyltransferase 1 (CPT1), which is localized in the outer mitochondrial membrane, and is responsible for the formation of ACs by catalyzing the transfer of the acyl group of a long-chain acyl-CoA to carnitine, was reported in the liver of rats upon treatment with 2,3,7,8-tetrachlorodibenzo-p-dioxin (TCDD) [55,56]. This might explain why ACs start to accumulate in cells that have been challenged by the carcinogens BAP and DALP. TCDD is known as a strong AHR inducer, thus the detected changes may be linked to AHR activation. While the exact molecular mechanisms of this interference between these carcinogenic PAHs and AC degradation remain obscure, lowered rates of β-oxidation and subsequent TCA cycle activity can explain why AA profiles were found strongly decreased under the very same conditions (Fig. 5, Supplementary Tbl. 1).

The phospholipid groups of PCs and SMs are important components of cell membranes and involved in regulation of cell function, membrane protein trafficking and inflammation [57]. PAHs are highly lipophilic compounds that can be directly incorporated into the cell membrane thereby disturbing the biological membrane function [58]. Perturbation of membrane’s integrity and the ultimate loss of functionality have been shown in diatoms upon BAP exposure [54] and might also be causative for the strong PC levels changes observed in our study (Supplementary Tbl. 1). Additionally, inducing canonical AHR signaling can be assumed to play a role in the phospholipid pattern changes of HaCaT cells exposed to BAP or CRY, but not to DALP. A recent study predicted genes like PPARδ and Lpin involved in lipid biosynthesis to be direct AHR targets [59]. An activation of PPARδ was also found in a recent proteomics study using Hepa1c1c7 cells exposed to BAP [60]. Yet, this interference of AHR signaling with lipid metabolism is still poorly understood. For instance, the PCaa_C36:2 levels of cells were only increased after BAP and CRY exposure and therefore identified as a potential biomarker for canonical AHR activation after PAH exposure. PCaa_C36:2 was also detected as a possible biomarker to distinguish smokers from non-smokers in the KORA (Cooperative Health Research in the Region of Augsburg, Germany) study [61]. As cigarette smoke contains a mixture of carcinogenic PAHs including...
BAP [62], our data provide further evidence that this particular lipid can serve as biomarker indicating the exposure of cells against AHR activating PAHs. By contrast, DALP, that did not induce AHR downstream signaling in HaCaT cells (i.e., induction of CYP enzymes), indeed caused a measurable but insignificant decrease in the level of PCaC36:2 (Table 1).

The increases of the levels of certain SMs (i.e., SM(OH)C24:1, SM,C18:0, SM,C26:0, SM,C26:1) found in HaCaT WT cells after treatment with BAP or DALP can be associated with the onset of apoptosis. Sphingolipids represent a complex group of biomolecules that function as transducers in eukaryotic cells, modulating cell growth, differentiation and cell death [63]. SMs can be hydrolyzed by sphenoglycanine (SMase) to ceramides. After hydrolysis of SMs, ceramides function as second messengers activating various cytokines such as TNF-α that are involved in the signaling of apoptosis [64]. Therefore, the increases of SM levels found in cells after exposure to the strong carcinogens BAP and DALP are likely a result of the pro-apoptotic effects of these compounds. Exposure of HaCaT WT cells against BAP at concentrations known to activate the AHR revealed a strong enhancement of CD95 (Fas)–mediated apoptosis [41], which can similarly be triggered by ceramides [65].

5. Conclusion

Applying a multivariate model we have identified altogether 24 biomarkers in HaCaT WT cells that revealed sufficient to discriminate between exposure groups after treatment with three different PAHs. In previous PAH toxicity studies, long known cellular endpoints such as CYP1A1 induction and DNA adduct formation have been repeatedly proposed to be used in the risk assessment of these environmental pollutants. The data presented complement this set of rather historical biomarkers by novel metabolites originating from lipid metabolism and turnover. The alterations in the metabolomics profiles measured suggest influences of PAHs on the energy and lipid metabolism in the cells investigated.

This study is a proof of principle work using single compound exposure only. As PAH exposure might lead to interactions if such compounds occur in mixtures, in future experiments binary mixtures as well as multi PAH mixtures should be studied and the resulting metabolomic pattern changes compared. We would suggest experiments using PAH concentrations with identical potency in combination with experiments investigating co-exposure scenarios, where one of the investigated chemicals is at low dose in order to identify additive or synergistic effects. Further mechanistic insights should be gained by experiments that relate metabolic changes to pathological ways. By specific inhibition of steps in relevant signaling pathways (e.g., AHR, p53 or Nrf2) resulting metabolomic marker sets could be related to those pathway steps. This way metabolomic biomarkers indicative for inhibition or activation of relevant signaling pathways could be identified [35]. All this information would finally help to develop in vitro risk assessment strategies for PAH mixtures.

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Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.toxrep.2016.09.003.

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