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Formation of trans-epoxy fatty acids correlates with formation of isoprostanes and could serve as biomarker of oxidative stress

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ABSTRACT

In mammals, epoxy-polyunsaturated fatty acids (epoxy-PUFA) are enzymatically formed from naturally occurring all-cis PUFA by cytochrome P450 monooxygenases leading to the generation of cis-epoxy-PUFA (mixture of R,S- and S,R-enantiomers). In addition, also non-enzymatic chemical peroxidation gives rise to epoxy-PUFA leading to both, cis- and trans-epoxy-PUFA (mixture of R,R- and S,S-enantiomers). Here, we investigated for the first time trans-epoxy-PUFA and the trans/cis-epoxy-PUFA ratio as potential new biomarker of lipid peroxidation. Their formation was analyzed in correlation with the formation of isoprostanes (IsoP), which are commonly used as biomarkers of oxidative stress. Five oxidative stress models were investigated including incubations of three human cell lines as well as the in vivo model Caenorhabditis elegans with tert-butyl hydroperoxide (t-BOOH) and analysis of murine kidney tissue after renal ischemia reperfusion injury (IRI). A comprehensive set of IsoP and epoxy-PUFA derived from biologically relevant PUFA (ARA, EPA and DHA) was simultaneously quantified by LC-ESI(-)-MS/MS. Following renal IRI only a moderate increase in the kidney levels of IsoP and no relevant change in the trans/cis-epoxy-PUFA ratio was observed. In all investigated cell lines (HCT-116, HepG2 and Caki-2) as well as C. elegans a dose dependent increase of both, IsoP and the trans/cis-epoxy-PUFA ratio in response to the applied *t*-BOOH was observed. The different cell lines showed a distinct time dependent pattern consistent for both classes of autoxidatively formed oxylipins. Clear and highly significant correlations of the trans/cisepoxy-PUFA ratios with the IsoP levels were found in all investigated cell lines and *C. elegans*. Based on this, we suggest the trans/cis-epoxy-PUFA ratio as potential new biomarker of oxidative stress, which warrants further investigation.

1. Introduction

Oxidative stress, i.e. the imbalance of antioxidative and oxidative mechanisms is associated with the pathophysiology of several diseases. It is characterized by an impairment of redox signaling and control in the organism, ambivalently caused and/or resulting by the disease state [1]. Various diseases are linked to oxidative stress including inflammatory, cardiovascular, respiratory and neurological diseases [2].

Elevated levels of reactive oxygen and nitrogen species formed in the course of oxidative stress oxidatively modify biomolecules including lipids, proteins, DNA, thereby altering their biological properties and function. Polyunsaturated fatty acids (PUFA) which are essential constituents in membrane phospholipids are susceptible to free radical mediated autoxidation [3]. During this lipid peroxidation radical abstraction of a bisallylic hydrogen and subsequent addition of molecular oxygen results initially in conversion of PUFA to hydroperoxy fatty acid radicals which are further converted in chain oxidation reactions giving rise to a multitude of reactive and stable secondary oxidation products which inter alia bear hydro(pero)xy-, epoxy- and cyclic moieties [4]. Several of these products arising during non-

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enzymatic autoxidation are structurally similar to products formed by enzymatic conversion. However, enzymatic conversion of PUFA by cyclooxygenase (COX) results in regio- and stereospecific formation of prostaglandins (PG) whereas non-enzymatic lipid peroxidation leads to a complex mixture of regio- and stereoisomeric prostaglandin-like products, referred to as isoprostanes (IsoP), comprising in the case of arachidonic acid (ARA, C20:4n6) 64 different F₂-IsoP isomers [5,6]. While only free, i.e. non-esterified, PUFA are enzymatically converted by COX, IsoP are formed from PUFA esterified in phospholipids and are released upon stimuli [7]. PG, like PGE2, are known to be potent mediators of inflammation and for selected IsoP also biological activity has been shown, e.g. 15-F_{2t}-IsoP acting as renal vasoconstrictor [8]. ARA derived F_2 -IsoP, especially 15- F_{2t} -IsoP (8-*iso*-PGF_{2a}), were assessed to be valuable for the evaluation of oxidative stress in vivo and the latter is commonly used as biomarker in diseases and environmental exposures implicated with oxidative stress [9-11]. However, in recent years the validity of the sole measurement of 15-F_{2t}-IsoP as biomarker of oxidative stress is regarded controversial as it besides non-enzymatic autoxidation may also arise from enzymatic conversion by COX [12,13]. To account for the contribution of both pathways to the detected 15-F_{2t}-IsoP level, determination of the 15-F_{2t}-IsoP/PGF_{2 α} ratio may be used to differentiate between non-enzymatic and enzymatic formation and enable accurately evaluating the underlying oxidative stress [14,15]. Moreover, the parallel analysis of different IsoP isomers from individual as well as multiple biological relevant PUFA - as carried out in the present study - allows drawing comprehensive conclusions on the oxidative stress status independent from the origin of the analyzed specimen.

In addition to IsoP a multitude of autoxidative PUFA products is generated during lipid peroxidation. This includes products with epoxy moieties resulting from hydroperoxides by intramolecular homolytic substitution of the peroxide bond by an adjacent carbon radical [4]. Epoxy-PUFA are well characterized highly potent lipid mediators which possess important biological properties, e.g. acting anti-inflammatory and vasodilatory [16] and are present in body fluids and tissues (e.g. plasma, red blood cells, liver, kidney, lung, heart) [17,18]. In mammals they are generated from enzymatic conversion of the naturally occurring all-cis PUFA by cytochrome P450 monooxygenases (CYP) leading to regioisomeric cis-epoxy-PUFA (R,S- and S,R-enantiomers, Fig. 1) for each of the double bonds with distinct stereo- and regioselectivity depending on the respective involved CYP isoform [19,20]. Similarly to COX only the free, non-esterified PUFA are enzymatically converted by CYP, whereas in contrast to PG, epoxy-PUFA are mainly incorporated in lipids [21], e.g. predominantly the sn-2 position of phospholipids [22].

Apart from the CYP derived *cis*-epoxy-PUFA also *trans*-epoxy-PUFA (*S*,*S*- and *R*,*R*-enantiomers, Fig. 1) have been detected in red blood cells (RBC) [23] and heart tissue from untreated healthy animals [24]. Exposure of RBC with the oxidative stress causing agent *tert*-butylhy-droperoxide (*t*-BOOH) led to an increase of ARA derived epoxy-PUFA compared to control [25,26]. Regarding stereochemistry in response to *t*-BOOH treatment both diastereomers (i.e. *cis* and *trans*) increased in RBC, though a greater increase of *trans*-epoxy-ARA was observed [26].

A favored formation of *trans*- over *cis*-epoxy-ARA was also observed after exposing ARA to radical starter in benzene and liposomes [24]. However, so far no information about the time course of *trans*-epoxy-PUFA formation in biological settings and its correlation to the extent of oxidative stress is available.

In the present study we examined the formation of IsoP and epoxy-PUFA following oxidative stress to investigate the potential use of *trans*epoxy-PUFA and the *trans/cis*-epoxy-PUFA ratio as marker of oxidative stress. For this purpose, we extended an established LC-ESI(-)-MS/MS method covering enzymatically and non-enzymatically formed oxylipins to enable the separation and parallel quantification of *cis*- and *trans*-epoxy-PUFA. The concentration and time dependent generation of the oxidative lipid metabolites was examined in three cell lines from different human tissues and the model organism *Caenorhabditis elegans (C. elegans)* causing oxidative stress by treatment with *t*-BOOH, and in murine kidney tissue after renal ischemia reperfusion injury (IRI). The parallel assessment of both, IsoP, known biomarker of oxidative stress, and epoxy-PUFA allowed us to demonstrate the applicability of the *trans/cis*-epoxy-PUFA ratio as oxidative stress marker.

2. Materials and methods

2.1. Chemicals

Standards of regioisomeric *cis*-epoxy-PUFA (mixture of *R*,*S*- and *S*,*R*enantiomers), isoprostane 15-F_{2t}-IsoP and the deuterated internal standards (IS) ²H₄-15-F_{2t}-IsoP (²H₄-8-*iso*-PGF_{2α}), ²H₁₁-5(*R*,*S*)-5-F_{2t}-IsoP (²H₁₁-(\pm)5-iPF_{2a}-VI), ²H₄-9(10)-EpOME, ²H₁₁-14(15)-EpETrE were purchased from Cayman Chemicals (local distributor: Biomol, Hamburg, Germany). Other IsoP and IsoF standards from the biologically relevant PUFA α-linolenic acid (18:3n3, ALA), ARA, eicosapentaenoic acid (20:5n3, EPA), adrenic acid (22:4n6, AdA), docosapentaenoic acid (22:5n6, DPA_{n6}), docosahexaenoic acid (22:6n3, DHA) [27] were synthesized according to our chemical strategies already described in the literature [28–32].

LC–MS-grade methanol (MeOH), LC–MS-grade acetonitrile (ACN), LC–MS-grade isopropanol and LC–MS-grade acetic acid were purchased from Fisher Scientific (Schwerte, Germany). Disodium hydrogen phosphate dihydrate and *n*-hexane (HPLC Grade) were obtained from Carl Roth (Karlsruhe, Germany). Potassium hydroxide (85%) was obtained from Gruessing GmbH (Filsum, Germany). All other chemicals were purchased from Sigma Aldrich (Schnelldorf, Germany).

2.2. Cell culture assay

HCT-116 human colorectal carcinoma cells, Caki-2 human kidney carcinoma cells and HepG2 human liver carcinoma cells were grown in 10 cm dishes and incubated with 50 μ M and 200 μ M of *t*-BOOH (Sigma Aldrich, Schnelldorf, Germany) for 30 min, 1 h and 2 h. The cells were harvested using trypsin as described and the cell pellets were stored at -80 °C until analysis [27]. Cell pellets typically contained 5 × 10⁶ (HCT-116), 4 × 10⁶ (Caki-2) and 20 × 10⁶ (HepG2) cells.



Fig. 1. Stereochemistry of epoxy-PUFA. Each of the four double bonds can be epoxygenated resulting in the formation of four regioisomeric epoxy-PUFA from ARA each comprising two diastereoisomers, i.e. *cis*- and *trans*-epoxy-PUFA isomers with two enantiomers (with *S*,*R* and *R*,*S* or *S*,*S* and *R*,*R* configuration, respectively).

Cytotoxicity was assessed by the MTS assay (3-(4,5-dimethylthiazol-2-yl)-5-(3-carboxymethoxyphenyl)-2-(4-sulfophenyl)-2H-tetrazolium) as described [27,33]. The used *t*-BOOH concentration showed no relevant effect on cell viability (> 84%) (SI Fig. S1, [27]).

2.3. C. elegans treatment

The N2 Bristol strain, provided by the Caenorhabditis Genetics Center (CGC; University of Minnesota), was propagated at 20 °C on Nematode Growth Medium (NGM) plates spotted with the *Escherichia coli* strain OP50-1 [34]. Age-synchronized L4 stage worms were obtained by treating worms with an alkaline bleach solution (1% NaClO and 0.25 M NaOH) and growing the hatched L1 larvae on OP50-1seeded NGM plates for approximately 48 h. Treatment was performed using 12 000 L4 stage worms. Therefore, the nematodes were exposed to 0, 2.5 mM t-BOOH for 60 min or 6.5 mM t-BOOH for 15, 30, 45 or 60 min in 85 mM NaCl containing 0.01% Tween (6.5 mM represents the lethal dose 50% (LD50) following 1 h exposure (data not shown)). Worms were then pelleted by centrifugation at 1600 rpm for 2 min and washed four times in 85 mM NaCl containing 0.01% Tween. The remaining nematode pellet was snap-frozen in liquid nitrogen and stored at -80 °C until analysis.

2.4. Renal ischemia reperfusion injury in mice

C57BL/6J^{ham-ztm} male mice (12–13 weeks of age) were purchased from the institute of laboratory animal science (Hannover Medical School, Germany). Animals were cared for in accordance with the institution's guidelines for experimental animal welfare and with the guidelines of the American Physiological Society. The animal protection committee of the local authorities (Lower Saxony state department for food safety and animal welfare, LAVES) approved all experiments (approval 33.14–42502-04-14/1657). Mice were housed under conventional conditions with a 12 h light/dark cycle and had free access to food (Altromin 1324 standard mouse diet) and domestic quality drinking water *ad libitum*.

Surgery to induce renal IRI was done in general isoflurane anesthesia (5% induction, 2% maintenance) in combination with iv analgetic treatment with butorphanol. IRI was induced by transient unilateral renal pedicle clamping for 35 min using a non-traumatic vascular clamp [35]. Mice were sacrificed 2 h, 4 h and 24 h after reperfusion by deep general anesthesia and total body perfusion with ice cold PBS. Kidneys were collected, immediately shock frozen and stored at -80 °C until oxylipin analysis.

2.5. Quantification of oxylipins

For the analysis of total, i.e. free and esterified oxylipins formed under different treatment conditions cell pellets (comprising 5×10^6 HCT-116, 4×10^6 Caki-2 or 20×10^6 HepG2 cells) and nematode pellets as well as kidney tissue samples (25 ± 3 mg) were extracted following alkaline hydrolysis using anion exchange Bond Elut Certify II SPE cartridges (Agilent, Waldbronn, Germany) as described [27,36].

In brief, to cell pellets (HCT-116, HepG2, Caki-2), *C. elegans* pellets and kidney tissue samples 10 µL of IS solution (100 nM of ${}^{2}H_{4}$ -15- F_{2t} -IsoP, ${}^{2}H_{11}$ -5-(*R*,*S*)-5- F_{2t} -IsoP, C19-17-*epi*-17- F_{1t} -PhytoP, C21-15- F_{2t} -IsoP, ${}^{2}H_{4}$ -6-keto-PGF_{1α}, ${}^{2}H_{4}$ -PGE₂, ${}^{2}H_{4}$ -PGD₂, ${}^{2}H_{4}$ -TxB₂, ${}^{2}H_{4}$ -LTB₄, ${}^{2}H_{4}$ -9 – HODE, ${}^{2}H_{8}$ -5-HETE, ${}^{2}H_{8}$ -12-HETE, ${}^{2}H_{6}$ -20-HETE, ${}^{2}H_{11}$ -14,15-DiHETrE, ${}^{2}H_{11}$ -14(15)-EpETrE, ${}^{2}H_{4}$ -9(10)-EpOME and ${}^{2}H_{4}$ -9,10-DiHOME), 10 µL of antioxidant solution (0.2 mg/mL BHT and EDTA, 100 µM indomethacin, 100 µM of the soluble epoxide hydrolase inhibitor *trans*-4-[4-(3-Adamantan-1-yl-ureido)-cyclohexyloxy]-benzoic acid (*t*-AUCB) [37] in MeOH/water (50/50, ν/ν) and 50 µL water were added. The samples were homogenized after addition of 400 µL isopropanol with a vibration ball mill (MM 400, Retsch, Haan, Germany) using two stainless steel beads (3 mm, 10 min, 25 Hz). After hydrolysis (300 µL 1.5 M KOH (25/75, H₂O/MeOH, ν/ν) for 30 min at 60 °C) the samples were immediately cooled, neutralized using acetic acid (50%) and mixed with 2000 µL 0.1 M disodium hydrogen phosphate buffer (pH = 6) followed by SPE. After sample loading, the cartridges were washed with one column volume of each water and MeOH/water (50/50, ν/ν) and dried. The analytes were eluted with 2 mL of 75/25 (ν/ν) ethylacetate/*n*-hexane with 1% acetic acid, evaporated and the residue was resuspended in 50 µL MeOH (containing 40 nM of 1-(1-(ethylsulfonyl)piperidin-4-yl)-3-(4-(trifluoromethoxy)phenyl)urea as IS2 for the calculation of the extraction efficiency of the IS). After centrifugation (10 min, 4 °C, 20,000 × g) the samples were analyzed by RP-LC–MS/MS (6500 QTRAP, Sciex, Darmstadt, Germany) in scheduled selected reaction monitoring mode following negative electrospray ionization as described [27].

2.6. Data analysis

Oxylipin concentrations were quantified using external calibration with authentic standards (linear fitting with $1/x^2$ weighting) based on the analyte to corresponding IS area ratio as described [27]. For each epoxy-fatty acid regioisomer both *cis*- and *trans*-epoxy-PUFA isomers were quantified using the calibration curve of corresponding *cis*-epoxy-PUFA (identified with authentic standards). Based on that, the *trans/cis*epoxy-PUFA ratios were calculated. For 7(8)-EpDPE and 5(6)-EpETE due to absence of accurate quantitative standards only the *trans/cis*epoxy-PUFA ratio could be determined. *Trans*-epoxy-PUFA isomers eluting 0.14–0.3 min after their corresponding *cis*-isomers (SI Fig. S2) were identified based on retention time and identical MS-fragmentation pattern as described [23,24,26].

Data evaluation and statistical analyses were performed as indicated using GraphPad Prism version 6.01 for Windows (GraphPad Software, La Jolla California USA, www.graphpad.com).

3. Results

In the present study using a comprehensive LC–MS/MS method total, i.e. free and esterified, levels of IsoP as well as *cis*- and *trans*-epoxy-PUFA derived from different precursor PUFA were simultaneously quantified in cell pellets and *C. elegans* incubated with oxidative stress generating *t*-BOOH and in murine kidney tissue after renal ischemia reperfusion injury.

3.1. Formation of IsoP

3.1.1. IsoP in cells following treatment with t-BOOH

In cell pellets from colorectal (HCT-116), renal (Caki-2) and hepatic (HepG2) origin 6 IsoP regioisomers derived from 4 precursor PUFA (ARA, EPA, DHA, AdA) were quantified. Regarding detected isomers the regioisomers carrying the hydroxyl group in proximity to the carboxy function, i.e. ARA derived 5(R,S)-5-F_{2t}-IsoP, EPA derived 5(R,S)-5-F_{3t}-IsoP, DHA derived 4(R,S)-4-F_{4t}-NeuroP and AdA derived ent-7(R,S)-7-F_{2t}-dihomo-IsoP showed higher concentration than other regioisomers from the same PUFA or were the only regioisomers detected as it is the case for AdA derived F2t-dihomo-IsoP. Under basal conditions in HCT-116 cells only 5(R,S)-5- F_{2t} -IsoP, 5(R,S)-5- F_{3t} -IsoP and 4(R,S)-4- F_{4t} -NeuroP were detected while in Caki-2 and HepG2 cells additionally other regioisomers (15-F_{2t}-IsoP and 10(R,S)-10-F_{4t}-NeuroP) were found. Interestingly, basal levels of detected IsoP in Caki-2 cells were 2-7 fold higher than in HepG2 and 6-20 fold higher than in HCT-116 cells. In all cell lines incubation with t-BOOH led to an increase in IsoP formation, however the dose and time dependent pattern showed considerable differences between the cell types, though being similar for all regioisomers within the individual cell types (Fig. 2). In HCT-116 cells a dose dependent increase of IsoP level was observed leading to 3-5 fold and 10-17 fold higher IsoP concentrations compared to control with 50 µM and 200 µM t-BOOH respectively (Fig. 2I). While with 50

IsoP / *trans*-epoxy-PUFA □ 0.5 h □ 1 h □ 2 h *cis*-epoxy-PUFA ☑ 0.5 h ☑ 1 h ☑ 2 h



Fig. 2. Total levels of representative **(A)** IsoP and **(B)** epoxy-PUFA derived from ARA, EPA and DHA following incubation of **(I)** HCT-116, **(II)** HepG2 and **(III)** Caki-2 cells with 50 μ M and 200 μ M *t*-butyl hydroperoxide (*t*-BOOH) for 0.5 h, 1 h and 2 h. Shown are mean \pm SD (HCT-116: n = 4, HepG2: n = 3, Caki-2: n = 4). For epoxy-PUFA the *trans/cis*-epoxy-PUFA-ratio (*trans/cis*) is indicated above the corresponding bars.

 μ M *t*-BOOH no effect of the incubation time (0.5 h–2 h) was observed, 200 μ M *t*-BOOH induced a trend towards higher IsoP levels with longer incubation time. In HepG2 cells only 200 μ M of *t*-BOOH led to an increase of IsoP levels compared to control (Fig. 2II). Interestingly, with longer incubation time a decrease in IsoP concentrations was observed leading at 2 h to almost same concentrations as with 50 μ M *t*-BOOH. In Caki-2 cells only incubation with 200 μ M *t*-BOOH led to a slight increase of IsoP concentrations, however here no time dependent change of IsoP levels was observed (Fig. 2III).

3.1.2. IsoP in C. elegans following treatment with t-BOOH

In nematode pellets of wildtype N2 *C. elegans* only EPA derived 5(R,S)-5-F_{3t}-IsoP were detected in the control group. Incubation with *t*-BOOH led to an increase and additional formation of EPA derived 8(R,S)-8-F_{3t}-IsoP as well as ARA derived 15-F_{2t}-IsoP and 5(R,S)-5-F_{2t}-IsoP. A dose dependent increase in IsoP levels was observed with the applied *t*-BOOH concentration leading to 4–6 fold higher IsoP levels for 6.5 mM compared to 2.5 mM *t*-BOOH after 60 min incubation (Fig. 3I). Also, longer incubation time led to a gradual increase resulting in 46 fold higher IsoP levels for 60 min compared to 15 min incubation with 6.5 mM *t*-BOOH (Fig. 3II).

3.1.3. IsoP in murine kidneys after ischemia reperfusion injury

In murine kidney tissue ARA derived 5(R,S)-5- F_{2t} -IsoP and 15- F_{2t} -IsoP and DHA derived 4(R,S)-4- F_{4t} -NeuroP and 10(R,S)-10- F_{4t} -NeuroP were detected with higher levels of the regioisomers carrying the hydroxyl group in proximity to the carboxy function. While in IRI tissue 2 h after reperfusion higher levels of ARA derived IsoP were observed compared to unclamped contralateral kidney tissue, DHA derived NeuroP remain almost unchanged. 4 h after reperfusion higher levels for both, ARA and DHA derived IsoP were observed in IRI tissue compared to unclamped control. Interestingly, 24 h after unilateral IRI lower levels of DHA derived NeuroP were observed in the IRI kidney

compared to the contralateral unclamped kidney, while levels of ARA derived IsoP were similar in the IRI and unclamped kidney (Fig. 4).

3.2. Formation of epoxy-PUFA

3.2.1. Epoxy-PUFA in cells following treatment with t-BOOH

In cell pellets from colorectal (HCT-116), renal (Caki-2) and hepatic (HepG2) origin for all epoxy-PUFA derived from ARA (i.e. EpETrE), EPA (i.e. EpETE) and DHA (i.e. EpDPE) *cis*- and *trans*-isomers could be determined. However as a result of chromatographic interference in HCT-116 cells *cis*-19(20)-EpDPE and in HepG2 cells *cis*-8(9)-EpETE were not evaluable.

In control incubations the *trans/cis*-epoxy-PUFA ratios of the respective regioisomers were similar between the different cell types, however differed between the individual regioisomers. While for most regioisomers levels of *cis*-isomers were higher than of *trans*-isomers, for regioisomers carrying the epoxy moiety in the middle or front part of the carbon chain (i.e. 8(9)-EpETrE, 11(12)-EpETrE, 5(6)-EpETE, 7(8)-EpDPE and 10(11)-EpDPE) the ratios of *trans/cis*-isomers were approximately equal (0.7–1.4) or higher as for 5(6)-EpETrE (*trans/cis*-ratio > 1.5).

Incubation with *t*-BOOH led in all cell lines to an increase of *trans*epoxy-PUFA levels, while levels of respective *cis*-isomers remained almost constant, resulting in an increase of the *trans/cis*-epoxy-PUFA ratios. However, dose and time dependent formation of *trans*-epoxy-PUFA differed between the investigated cell lines with the individual regioisomers showing the same pattern within the particular cell types irrespective of the precursor PUFA (Fig. 2). In HCT-116 cells with 50 μ M *t*-BOOH *trans*-epoxy-PUFA remained almost constant, whereas 200 μ M *t*-BOOH led to 1.7–6.6 fold higher levels of *trans*-isomers compared to control. Regarding incubation time, only with 200 μ M *t*-BOOH a trend towards higher levels of *trans*-epoxy-PUFA with longer incubation was observed, leading to a 1.7–2.7 fold increase of the *trans/cis*-ratios



Fig. 3. Total levels of **(A)** IsoP and **(B)** epoxy-PUFA derived from ARA and EPA in *C. elegans* incubated for **(I)** 60 min with 2.5 mM and 6.5 mM *t*-BOOH and for **(II)** 15, 30, 45 and 60 min with 6.5 mM *t*-BOOH. Shown are mean \pm SD (n = 2–6). In case the levels do not exceed the lower limit of quantification (LLOQ) in one treatment group, the LLOQ is indicated as a dashed line. For epoxy-PUFA the *trans/cis*-epoxy-PUFA-ratio (*trans/cis*) is indicated above the corresponding bars. Statistical differences between control and the treatment groups were evaluated by one-way ANOVA followed by Tukey post-test. For dose dependent formation of 5-(*R*,*S*)-5-*F*_{2t}-IsoP unpaired *t*-test was used. For time dependent formation only differences between control and treatment groups are shown, results for differences between all treatments groups are summarized in SI Tab. S1 (* p < 0.05, ** p < 0.01, *** p < 0.001).



Fig. 4. Total levels of (A) IsoP and (B) epoxy-PUFA derived from ARA and DHA in murine kidney tissue following unilateral IRI for 35 min. Shown are mean \pm SEM (n = 6) in the unclamped and IRI kidneys 2h, 4h and 24 h after reperfusion. For epoxy-PUFA the trans/cis-epoxy-PUFA-ratio (trans/ cis-ratio) is indicated above the corresponding bars. Statistical differences between the unclamped and IRI kidneys were evaluated by paired t-test for 5-(R,S)-5-F_{2t}-IsoP, 4(R,S)-4-F_{4t}-NeuroP and the trans/cis-epoxy-PUFA ratios (* p < 0.05, p < 0.01, p < 0.001).

for 1 h and 2 h compared to 0.5 h incubation (Fig. 2I). In HepG2 cells 50 μ M *t*-BOOH led only to a slight increase of the *trans/cis*-ratios (1.5–1.8 fold), while with 200 μ M *t*-BOOH *trans/cis*-ratios were 2.0–5.2 fold higher compared to control. Similarly to HCT-116 cells with 50 μ M *t*-BOOH *cis*- and *trans*-isomer levels remained almost unchanged with longer incubation time. In contrast, with 200 μ M *t*-BOOH levels of *trans*-epoxy-PUFA increased after 0.5 h and remain similar after 1 h, however

after 2 h of incubation a trend towards lower levels of *cis*- and *trans*epoxy-PUFA was observed (Fig 2II). In Caki-2 cells only with 200 μ M *t*-BOOH a slight trend towards higher levels of *trans*-epoxy-PUFA was observed, however neither 50 μ M nor 200 μ M *t*-BOOH led to a time dependent change of epoxy-PUFA levels (Fig. 2III).

3.2.2. Epoxy-PUFA in C. elegans following treatment with t-BOOH

In non-exposed wildtype N2 C.elegans cis- and trans-isomers of all analyzed epoxy-PUFA regioisomers derived from ARA and EPA were detected. Levels of EPA derived epoxy-PUFA were markedly higher than ARA derived epoxy-PUFA. Interestingly, levels of the terminal regioisomers, i.e. 14(15)-EpETrE and 17(18)-EpETE, were different compared to the other regioisomers. While levels of ARA derived regioisomers were in the same range, 17(18)-EpETE showed 2.7-5.2 fold higher basal levels than the other EPA derived regioisomers. Incubation with t-BOOH led to a dose and time dependent increase in epoxy-PUFA levels. With 2.5 mM t-BOOH for 60 min both, cis- and trans-isomers showed a slight but similar trend towards higher levels compared to control resulting in unchanged *trans/cis*-ratios. Further increasing the applied t-BOOH concentration to 6.5 mM led to a favored formation of trans- over cis-isomers, leading to a 2.3-2.9 fold increase in trans/cisratios compared to control (Fig. 3I). A similar pattern was also observed regarding incubation time with 6.5 mM t-BOOH. While after 15 min cisand trans-isomers increased only slightly though to a similar extent, longer incubation (30-60 min) resulted in a favored formation of transover cis-isomers. Consequentially trans/cis-ratios increased gradually with longer incubation time resulting in 1.7-2.4 fold higher trans/cisratios for 60 min compared to 15 min incubation (Fig. 3II).

3.2.3. Epoxy-PUFA in murine kidney after ischemia reperfusion injury

In murine kidney tissue only for ARA and DHA derived epoxy-PUFA regioisomers cis- and trans-isomers were detected. For both precursor PUFA basal levels of individual epoxy-PUFA regioisomers showed a distinct distribution of trans- and cis-isomers depending on the position of the epoxy-moiety relative to the carboxy-function: the closer the epoxy-moiety to the carboxy terminus the higher the observed trans/cisratio (SI Fig. S3). Following unilateral IRI, 2 h after reperfusion levels of epoxy-PUFA remained almost unchanged, while 4 h after reperfusion both cis- and trans-epoxy-PUFA showed a slight increase compared to unclamped kidney resulting in almost unchanged *trans/cis*-ratios. 24 h after reperfusion levels of ARA derived epoxy-PUFA were similar in IRI and unclamped tissue. While in contrast for DHA derived epoxy-PUFA a decrease of cis-isomers was observed in IRI compared to unclamped kidney tissue with almost unchanged trans-epoxy-PUFA level. Interestingly, this decrease was more pronounced for terminal isomers, i.e. 19(20)- and 16(17)-EpDPE compared to 13(14)- and 10(11)-EpDPE (Fig. 4, SI Fig. S3).

3.2.4. Correlation of IsoP

In cell lines as well as in *C. elegans* following incubation with *t*-BOOH a clear correlation between IsoP formation and the *trans/cis*epoxy-PUFA ratio was found (Fig. 5, Table 1). Pearson correlations between the levels of IsoP regioisomers carrying the hydroxyl group in proximity to the carboxy-function and the ratio of *trans/cis*-epoxy-PUFA regioisomers from the corresponding PUFA were determined (Table 1): ARA derived 5(R,S)-5 F_{2t}-IsoP was correlated with EpETrE regioisomers, EPA derived 5(R,S)-5-F_{3t}-IsoP with EpETE regioisomers and DHA derived 4(R,S)-4-F_{4t}-NeuroP with EpDPE regioisomers. For all epoxy-PUFA regioisomers positive and highly significant correlations with IsoP levels were obtained in the three cell lines and *C. elegans* (Fig. 5, Table 1). As no significant increase in the *trans/cis*-ratio was found in murine renal IRI (Fig. 4, SI Fig. S3) consequently no correlation of IsoP levels and *trans/cis*-epoxy-PUFA ratios resulted.

4. Discussion

In the present study total levels of IsoP and *cis*- and *trans*-epoxy-PUFA were simultaneously quantified. Based on that, the correlation between these autoxidatively formed oxylipins during oxidative stress was evaluated in three models: (i) *t*-BOOH induced stress in three cultured cell lines and in (ii) *C. elegans* and (iii) following ischemic reperfusion injury in the kidney in mice.

ARA derived F2-IsoP are well established marker of oxidative damage formed during lipid peroxidation in vivo. Particularly 15-F_{2t}-IsoP $(8-iso-PGF_{2\alpha})$ is commonly used as biomarker in diseases and environmental exposures related to oxidative stress [8,9,11]. In line we found a strong increase of IsoP in response to t-BOOH in all cell lines as well as in C. elegans (Fig. 2 and 3). The modulation of IsoP was markedly different between the cell lines regarding both, dose and time dependency: while in HCT-116 cells already 50 µM t-BOOH led to an increase in IsoP levels, in HepG2 and Caki-2 cells a trend towards higher IsoP levels was only observed in incubations with 200 µM t-BOOH (Fig. 2). This could result from different intracellular RONS detoxifying mechanisms between the cell types, breaking down t-BOOH generated radicals and preventing oxidative damage. It seems that only if a threshold in intracellular ROS burden is reached, an increased IsoP formation results. For HCT-116 cells this level seems to be reached at t-BOOH levels above 50 µM while for Caki-2 cells only at a high concentration of 200 µM t-BOOH IsoP levels are elevated. A threshold of oxidative stress inducing noxae for the formation of IsoP depending on the concentration of oxidative stress inducer was also observed previously in HepG2 cells treated with the redoxcycler paraquat [38]. Here only incubation with concentrations equal or above 100 µM led to an increase in IsoP formation compared to control.

Noteworthy, with high concentration of t-BOOH (200 µM) for all detected regioisomers, IsoP level showed a distinct time dependent pattern in the different cell lines. In HCT-116 cells a trend towards higher level of IsoP with longer incubation time was observed (Fig. 2I). Consistently, a similar time dependent increase of IsoP was previously reported for copper induced peroxidation of ARA and EPA containing phospholipids or of cell lysates [38,39]. In contrast, in HepG2 cells levels of IsoP showed a decreasing trend with longer incubation and remained unaffected by the incubation time in Caki-2 cells (Fig. 2). A decrease in total IsoP levels as observed in HepG2 cells might result from metabolic degradation of the formed oxylipins. As formation of IsoP occurs in situ on phospholipids [7] hydrolysis of IsoP containing phospholipids by phospholipase A2 seems likely to be involved in the observed decrease of total IsoP level. Indeed, the intracellular type II platelet-factor-acetyl hydrolase exhibiting negligible phospholipase activity against long chain-PUFA containing membrane phospholipids [40,41] hydrolyses F₂-IsoP containing phospholipids [42] and was shown to translocate from the cytosol to the membrane upon oxidative stress stimulus [43]. Furthermore, a time course similar to the one observed in HepG2 cells was also observed following CCl4 induced peroxidation in the rat [7]. Here, total IsoP levels in the liver increased rapidly after CCl₄ was applied with a peak after 2 h, however were followed by a decrease over time leading to almost restored levels after 24 h. Correspondingly, plasma level of free IsoP arising from release of esterified IsoP showed a similar, however delayed time course, further supporting the hypothesis of phospholipase contribution.

Among the multitude of different IsoP regio- and stereoisomers that can be formed during lipid peroxidation (e.g. for ARA a total of 64 different F₂-isomers) [5], we observed a favored formation of selected regioisomers in all investigated cell lines, C. elegans as well as kidney tissue. Regarding IsoP isomers covered in our analysis, regioisomers carrying the side chain hydroxy group in proximity to the carboxy group, i.e. ARA derived 5(R,S)-5-F_{2t}-IsoP, EPA derived 5(R,S)-5-F_{3t}-IsoP, DHA derived 4(R,S)-4-F4t-NeuroP and AdA derived ent-7(R,S)-7-F_{2t}-dihomo-IsoP, were observed in higher abundance than its regioisomers. This is consistent with previous studies where a preferential formation of particular regioisomers was observed. Similarly to our results for ARA the 5- and 15-F2t-IsoP, for EPA the 5- and 18-F3t-IsoP [44] and for DHA the 4- and 20-F4t-NeuroP [45] regioisomers were observed in greater abundance. This may be explained by further oxidative conversion of the precursor involved in the formation of other IsoP regioisomers during autoxidation. One could also expect, that the regioisomers bearing the hydroxy-group at C4 or C5 position are also more stable towards β -oxidation which could contribute to higher



Fig. 5. Correlation between the *trans/cis*-ratio of individual epoxy-PUFA regioisomers (i.e. for ARA 11(12)-EpETrE, for EPA 11(12)-EpETrE, for DHA 10(11)-EpDPE and the most abundant IsoP isomer derived from the same precursor PUFA (i.e. for ARA 5(R,S)-5- F_{2t} -IsoP, for EPA 5(R,S)-5- F_{3t} -IsoP, for DHA 4(R,S)-4- F_{4t} -NeuroP). Shown are pearson correlation coefficients r and respective p-values for ARA, EPA and DHA derived oxidative metabolites in **(A)** HCT-116, **(B)** HepG2 and **(C)** Caki-2 cells, as well as **(D)** *C. elegans* incubated with *t*-BOOH.

levels of these regioisomers, as has been observed previously for other 5-hydroxyeicosanoids (e.g. leukotriens) [46,47].

Remarkably, in all cell lines, levels of these regioisomers were in a similar range for ARA, EPA and DHA derived IsoP (Fig. 2) while in *C. elegans* levels of EPA derived 5(R,S)-5-F_{3t}-IsoP were considerably higher than the corresponding ARA derived regioisomer (Fig. 3). As PUFA serve as substrates for IsoP formation, the fatty acid composition and PUFA pattern affect the level of respective formed IsoP. Consistently, the observed high level of EPA derived F₃-IsoP in *C. elegans* is in agreement with the predominant relative EPA content compared to

ARA [48].

Regarding formation of epoxy-PUFA, the *trans/cis*-epoxy-PUFA ratio increased with the applied *t*-BOOH concentration in all cell lines and *C. elegans* and showed a distinct time dependent pattern in the different cell lines (Fig. 2 and 3). The similar pattern in comparison with the observed IsoP level suggests the contribution of related mechanisms (discussed above). Overall the increase in the *trans/cis*-epoxy-PUFA ratio was mainly caused by a favored formation of *trans*- compared to *cis*-isomers during *t*-BOOH induced oxidative stress. This supports previous findings demonstrating formation of epoxy-PUFA during

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Table

	5(<i>R</i> , <i>S</i>)-5-F _{2t} -Iso	уР				$5(R,S)-5-F_{3t}-Ist$	Ъ				4(<i>R</i> , <i>S</i>)-4-F _{4t} -N	uroP	
ARA	HCT-116	HepG2	Caki-2	C. elegans	EPA	HCT-116	HepG2	Caki-2	C. elegans	DHA	HCT-116	HepG2	Caki-2
14(15)-	r = 0.9231	r = 0.7177	r = 0.8764p	r = 0.9121	17(18)-	r = 0.8207	r = 0.8231	r = 0.7832	r = 0.9379	19(20)-	*	r = 0.7461	r = 0.7327
	p < 0.0001	p < 0.0001	< 0.0001	p < 0.0001		p < 0.0001	p < 0.0001	p < 0.0001	p < 0.0001			p < 0.0001	p < 0.0001
11(12)-	r = 0.8851	r = 0.6760	r = 0.8298	r = 0.9526	14(15)-	r = 0.8610	r = 0.7929	r = 0.8303	r = 0.9015	16(17)-	r = 0.8621	r = 0.6950	r = 0.8899
	p < 0.0001	p = 0.0001	p < 0.0001	p < 0.0001		p < 0.0001	p < 0.0001	p < 0.0001	p < 0.0001		p < 0.0001	p < 0.0001	p < 0.0001
-(6)8	r = 0.9085	r = 0.6261	r = 0.848	r = 0.9150	11(12)-	r = 0.9104	r = 0.6915	r = 0.9008	r = 0.9261	13(14)-	r = 0.8342	r = 0.7489	r = 0.8942
	p < 0.0001	p = 0.0005	p < 0.001	p < 0.0001		p < 0.0001	p < 0.0001	p < 0.0001	p < 0.0001		p < 0.0001	p < 0.0001	p < 0.0001
5(6)-	r = 0.8952	r = 0.6906	r = 0.7978	r = 0.8950	8(9)-	r = 0.8649	*	r = 0.7839	r = 0.9535	10(11)-	r = 0.8846	r = 0.6651	r = 0.8876
	p < 0.0001	p < 0.0001	p < 0.0001	p < 0.0001		p < 0.0001		p < 0.0001	p < 0.0001		p < 0.0001	p = 0.0002	p < 0.0001
					5(6)-	r = 0.8335	r = 0.662	r = 0.749	r = 0.9675	7(8)-	r = 0.9006	r = 0.5821	r = 0.9042
						p < 0.0001	p < 0.0006	p < 0.0001	p < 0.0001		p < 0.0001	p = 0.0014	p < 0.0001

Prostaglandins and Other Lipid Mediators 144 (2019) 106334

autoxidation, as has been shown for ARA derived EpETrE in RBC incubated with *t*-BOOH [25,26]. Consistently, both, free radical induced peroxidation in benzene and liposomes as well as exposure of RBC to *t*-BOOH led to a favored formation of *trans*- over *cis*-epoxy-ARA [24,26]. As double bonds in naturally occurring PUFA are oriented all-*cis* a possible mechanism leading to the formation of both, *cis*- and *trans*epoxy-PUFA could involve PUFA carbon centered radicals with an adjacent peroxyl group. These are formed at the position of the double bond in the PUFA molecule during the radical chain reaction. These carbon radicals allow for free rotation of the σ -bond prior intramolecular attack of the peroxyl bond by the reactive carbon radical which leads via homolytic substitution to the formation of an epoxymoiety oriented either *cis* or *trans* [4,24].

Irrespective of the precursor PUFA the observed increase in the *trans/cis*-epoxy-PUFA ratio following *t*-BOOH induced oxidative stress was similar for all regioisomers, though being slightly higher for 5(6) EpETE. However, this has to be regarded with caution as 5(6)-EpETE, like 5(6)-EpETrE might rapidly degrade forming a six-membered δ -lacton [49]. Additionally, also high basal levels of *cis*-epoxy-PUFA derived from CYP as observed for the preferred formation of the ω 3-epoxygenation product 17(18)-EpETE compared to its other regioisomers in *C. elegans* (Fig. 3) might hamper the suitability of the respective *trans/cis*-ratio reflecting the oxidative damage.

Here, we tested for the first time if the *trans/cis*-epoxy-PUFA ratio can reflect oxidative stress. For this purpose we correlated the *trans/cis*-epoxy-PUFA ratio with level of IsoP which are established biomarker of lipid peroxidation. Considering the consistent time and *t*-BOOH concentration dependent pattern, quite clear, positive highly significant correlations of IsoP level with the *trans/cis*-epoxy-PUFA ratio were observed for all epoxy-PUFA regioisomers in *t*-BOOH induced oxidative stress in all three cell lines and *C. elegans* (Fig. 5, Table 1). Thus, we suggest that the *trans/cis*-epoxy-PUFA ratio might serve as alternative oxidative stress marker.

However, in the evaluated invivo model of murine renal IRI, the moderate increase of IsoP was not accompanied by an increase in the trans/cis-epoxy-PUFA ratio (Fig. 4, SI Fig. S3). Thus, here the applicability of the trans/cis-epoxy-PUFA ratio as oxidative stress marker cannot be evaluated based on comparison with IsoP levels. Even though IRI is linked to excess ROS formation [50,51] and an increase of oxidative damage derived IsoP following renal IRI has been observed in rat models [52,53], there are various factors influencing measured IsoP levels in this model. These include clamping time, since longer clamping seem to lead to higher IsoP levels [52], as well as the time point of sampling following reperfusion and the specimen analyzed, e.g. plasma, interstitial fluid, urine or tissue. Consistently to our observation of an increase of total 5(R,S)-5-F_{2t}-IsoP levels in the kidney 4 h after reperfusion and a decrease after 24 h (Fig. 4), in a rat model of unilateral IRI an increase of oxidized phosphatidylcholine species containing IsoP was observed 6 h after reperfusion which decreased after 24 h [53]. A similar time course of IsoP level was also observed in the liver of CCl₄ treated rats (see above) [7]. These studies show that the time point of measuring oxidative damage for the evaluation of oxidative stress based on IsoP and likewise probably also epoxy-PUFA is crucial [54]. Interestingly, while showing the same time dependent trend, ARA and DHA derived 5(R,S)-5- F_{2t} -IsoP and 4(R,S)-4- F_{4t} -NeuroP, respectively, were modulated to a different extent in the time course of IRI in mice (Fig. 4). Also species specific differences might result in different time courses or extent of changes. Taken together, probably the ischemia reperfusion injury induced oxidative stress does not cause strong lipid oxidation in the investigated model and at the investigated time points. Therefore, other oxidative stress models, e.g. CCl₄ induced liver toxicity [8,9,55] or paraquat induced lung toxicity [56,57] should be used to evaluate the biological significance of the modulated trans/ cis-epoxy-PUFA ratio during oxidative stress as seen for the t-BOOH induced oxidative stress models.

5. Conclusion

Based on the presented results, we suggest a potential new biomarker of oxidative damage: the *trans/cis*-epoxy-PUFA ratio. In the investigated human cell lines from hepatic, colorectal and renal origin, as well as in the model organism *C. elegans* an increase in the *trans/cis*epoxy-PUFA ratio with *t*-BOOH induced oxidative stress was observed. The *trans/cis*-epoxy-PUFA ratio correlates perfectly with common marker of oxidative stress, i.e. IsoP. However, in the investigated *in vivo* model of murine renal ischemia reperfusion injury an effect on the *trans/cis*-epoxy-PUFA ratio was absent, underlining the need for further investigations especially regarding different oxidative stress agents with different modes of inducing oxidative stress to validate its suitability. This analysis could also pave the way to a mechanistically understanding of *trans*-epoxy-PUFA formation and understanding of their biological role.

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Appendix A. Supplementary data

Supplementary material related to this article can be found, in the online version, at doi:https://doi.org/10.1016/j.prostaglandins.2019. 04.004.

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