Oxidative Metabolism of a Fatty Acid Amide Hydrolase-Regulated Lipid, Arachidonoyltaurine†

Melissa V. Turman,§ Philip J. Kingsley,‡ Carol A. Rouzer,§ Benjamin F. Cravatt,§ and Lawrence J. Marnett*,‡

A. B. Hancock, Jr. Memorial Laboratory for Cancer Research, Departments of Biochemistry, Chemistry, and Pharmacology, Vanderbilt Institute of Chemical Biology, and Center in Molecular Toxicology, Vanderbilt University School of Medicine, Nashville, Tennessee 37232, and The Skaggs Institute for Chemical Biology, Departments of Cell Biology and Chemistry, The Scripps Research Institute, La Jolla, California 92037

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ABSTRACT: A novel class of lipids, N-acyltaurines, was recently discovered in fatty acid amide hydrolase knockout mice. In some peripheral tissues, such as liver and kidney, N-acyltaurines with long, polyunsaturated acyl chains are most prevalent. Polyunsaturated fatty acids are converted to a variety of signaling molecules by cyclooxygenases (COXs) and lipoxygenases (LOXs). The ability of COXs and LOXs to oxygenate arachidonoyltaurine was evaluated to gain insight into the potential metabolic fate of N-acyltaurines. Although arachidonoyltaurine was a poor substrate for COXs, mammalian 12S- and 15S-LOXs oxygenated arachidonoyltaurine with similar or better efficiency than arachidonic acid. Products of arachidonoyltaurine oxygenation were characterized by liquid chromatography-tandem mass spectrometry (LC-MS/MS). The positional specificity of single oxygenation was retained for 15S-LOXs. However, platelet-type 12S-LOX produced 12- and 15-hydroxyeicosatetraenoyltaurines (HETE-Ts). Furthermore, LOXs generated dihydroxyeicosatetraenoyltaurines (diHETE-Ts). Metabolism of arachidonoyltaurine by murine resident peritoneal macrophages (RPMs) was also profiled. Arachidonoyltaurine was rapidly taken up and converted primarily to 12-HETE-T. Over prolonged incubations, RPMs also generated small amounts of diHETE-T. Oxidative metabolism of polyunsaturated N-acyltaurines may represent a pathway for the generation or termination of novel signaling molecules.

The endocannabinoid system has emerged as a critical therapeutic target in recent years (1). Numerous pathophysiological conditions, including mood disorders, osteoporosis, myocardial ischemia/reperfusion injury, and cancer, could potentially be treated by modulation of the endocannabinoid system (1–8). One approach to the manipulation of this system is the inhibition of fatty acid amide hydrolase (FAAH)† (9). FAAH has a central role in the regulation of fatty acid amide levels in vivo, including those of the endocannabinoid arachidonoylethanolamide (AEA; Figure 1). Discovery metabolite profiling has revealed that another, previously unidentified class of lipids, N-acyltaurines (NATs), are substantially increased by chemical or genetic knockout of FAAH (10, 11). NATs are able to activate members of the transient receptor potential family of calcium ion channels in a cellular system (10). With the exception of these reports, the signaling and catabolic pathways for NATs have not been elucidated.

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* To whom correspondence should be addressed: Department of Biochemistry, Vanderbilt University School of Medicine, Nashville, TN 37232-0146. Telephone: 615-343-7329. Fax: 615-343-7534. E-mail: larry.marnett@vanderbilt.edu.

‡ Vanderbilt University School of Medicine.

§ The Scripps Research Institute.

FIGURE 1: Structures of arachidonoyl derivatives.

An initial approach to determine potential metabolic pathways is to probe the interactions of a novel molecule against enzymes that are known to metabolize related species. In peripheral tissues, such as the liver and kidney, NATs possessing long, polyunsaturated acyl chains are most prevalent (10), Linoleoyl (18:2), arachidonoyl (20:4), and docosahexaenoyl (22:6) taurines were increased from 20- to more than 100-fold in the livers of FAAH knockout mice. The parent polyunsaturated fatty acids are susceptible to both
enzymatic and nonenzymatic peroxidation of the allylic chain, and this process plays a central role in the biology and pathophysiology of lipid signaling. Prostaglandins (PGs) and hydroxyeicosatetraenoic acids (HETEs) are generated from the oxygenation of arachidonic acid by cylooxygenases (COXs) and lipoxygenases (LOXs), respectively, and serve as critical local bioactive mediators of cellular signaling (12–14). Moreover, these enzymes have been implicated in the metabolism of arachidonoyl derivatives, including the endocannabinoids 2-arachidonoylglycerol (2-AG) and AEA (Figure 1) (15).

COX-1 and COX-2 are homologous proteins, differing primarily in their expression profiles and, consequently, their physiological and pathophysiological functions (13, 16). Both isoforms exhibit similar kinetics and product profiles with arachidonic acid (16, 17). However, the volume of the COX-2 active site is approximately 20% larger than that of COX-1 (18). As a result, the COX-2 active site is able to accommodate a variety of esters and amides of arachidonate, including 2-AG and AEA, as well as arachidonoyl amino acids, such as N-arachidonoylglycerine (NAGly) (19–24). In contrast, COX-1 uses these substrates inefficiently, if at all.

Multiple isoforms of LOXs have been cloned from mammals and are classified by the position and stereochemistry of oxygen addition to arachidonic acid (25). Although all LOXs catalyze a single dioxygenation of polyunsaturated fatty acids, the regio- and stereospecificity of oxygenation varies between isoforms, as does substrate specificity. 15S-LOX-1 and leukocyte-type 12S-LOX (lk12-LOX) are promiscuous enzymes, metabolizing a range of C18, C20, and C22 fatty acids (26–28). These isoforms will also oxygenate fatty acid amides and esters, such as 2-AG, AEA, and NAGly (23, 29–32). 15S-LOX-1 and lk12-LOX can even incorporate oxygen into polyunsaturated acyl chains of phospholipids (33, 34). In contrast, platelet-type 12S-LOX (pl12-LOX) shows a marked preference for C20 fatty acids and is unable to efficiently metabolize neutral amides and esters of arachidonate (26, 29, 30, 32, 34–36). Notably the lipoic acid amides NAGly, N-arachidonoylalanine (NAld), and N-arachidonoyl-γ-aminobutyric acid (NAGABA), which would bear a negative charge at physiological pH, are kinetically comparable to arachidonic acid; however, the positional specificity of pl12-LOX is altered, and the enzyme acts as a 12/15-LOX on these substrates (23). Human 15S-LOX-2 also uses the arachidonoyl amino acids efficiently but maintains its positional specificity.

The COX and LOX pathways may modify polyunsaturated NAs and thus alter the activity of NAs in vivo. We investigated the ability of these enzymes to metabolize arachidonoyltaurine. We demonstrate that 15-LOX-1, lk12-LOX, and pl12-LOX but not COX can catalyze the peroxidation of arachidonoyltaurine, although with some perturbations of typical product profiles. We further characterized the ability of murine resident peritoneal macrophages to metabolize arachidonoyltaurine and show that arachidonoyltaurine is able to enter the LOX pathway in cells. LOX metabolism of arachidonoyltaurine may represent a pathway of termination or biosynthesis of signaling molecules.

**MATERIALS AND METHODS**

**Reagents.** Arachidonic acid and arachidonoylchloride were purchased from NuChek Prep (Elysian, MN). Arachidonoyltaurine was purchased from Cayman Chemical (Ann Arbor, MI). Ram seminal vesicles were purchased from Oxford Biomedical Research (Oxford, MI), and oCOX-1 was purified as previously described (37). Expression of mCOX-2 was performed with baculovirus reagents from BD Biosciences (San Diego, CA). Human 15-LOX-2 (h15-LOX-2) was expressed in bacteria. Purification procedures for mCOX-2, h15-LOX-2, and human platelet 12-LOX (pl12-LOX) were described previously (22, 31, 38). All of the expressed proteins were over 95% pure by sodium dodecyl sulfate–polyacrylamide gel electrophoresis (SDS–PAGE) staining and analysis, except for h15-LOX-2, which was 70% pure. Cell lysates containing rabbit reticulocyte 15-LOX-1 (r15-LOX-1) were purchased from Calbiochem (La Jolla, CA), and porcine leukocyte 12-LOX (lk12-LOX) was purchased from Cayman Chemical (Ann Arbor, MI).

**COX Activity Assay.** Quantification of cylooxygenase activity was performed in a thermostatted cuvette at 37 °C and monitored using a polarographic electrode with a YS5300 oxygen monitor (Yellow Springs Instrument Co., Inc., Yellow Springs, OH). Substrates were solubilized in dimethyl sulfoxide (DMSO). Activity assays were performed in 100 mM Tris-HCl buffer containing 500 µM phenol, with hematin-reconstituted protein (50 nM). Maximal reaction velocity data were obtained from the linear portion of the oxygen uptake curves and normalized to the metabolism rate of arachidonic acid.

**LOX Activity Assay.** LOX activity was detected by monitoring the absorbance of the conjugated diene product at 236 nm. UV assays were monitored using a Hewlett-Packard 8453 diode array spectrophotometer equipped with a thermostatted cuvette at 25 °C, with stirring at 180 revolutions per minute. The enzyme reactions included reaction buffer [50 mM Tris-HCl (pH 7.4) with 0.03% Tween-20] and substrate and were initiated by the addition of enzyme (3.7 µg/mL r15-LOX-1, 170 µg/mL h15-LOX-2, 200 µg/mL lk12-LOX, and 2 µg/mL pl12-LOX). Compounds were dissolved in acetonitrile (ACN) containing 10% acetic acid before the addition to the reaction buffer; ACN was kept below 1% reaction volume (2 mL). For the initial metabolism screen, compounds were diluted to a final concentration of 25 µM. To determine Michaelis–Menten kinetic parameters, the concentration of substrate was varied (1–50 µM). Maximal reaction velocity data were obtained from the linear portion of the absorbance curves, and the data were analyzed by nonlinear regression with Prism 4.0 (GraphPad Software, San Diego, CA).

**Enzyme Incubations for Product Identification.** To characterize products of oxidation, 5 µg of enzyme was incubated with 50 µM arachidonoyltaurine in 50 mM Tris-HCl at pH 8.0. After allowing the reaction to proceed for 15 min, 1 volume of 1 M sodium thiosulfate was added to reduce hydroperoxides to corresponding alcohols. The reaction mixture was then extracted with 3 volumes of ethyl acetate containing 0.5% acetic acid and 1 µM arachidonic acid-d8. The organic layer was removed and dried under argon, and the resulting residue was stored at −20 °C until analysis. For liquid chromatography–mass spectrometry (LC–MS...
analysis, the residue was resuspended in 1:1 methanol/5 mM ammonium acetate (pH 6.9) and passed through a SpinX filter to remove the precipitate.

**RPMs Cultures.** These experiments were performed with the approval of the Vanderbilt IACUC. Female CD-1 mice (25–30 g) were obtained from Charles River Laboratories (Wilmington, MA). The mice were sacrificed by asphyxiation with carbon dioxide, and the peritoneal cavities were lavaged with a total of 3 mL of ice-cold calcium- and magnesium-free phosphate-buffered saline (PBS) (39). Peritoneal cells were collected by centrifugation of lavage fluid and resuspended at a concentration of 2–3 x 10⁶ cells/mL in α-minimal essential medium (α-MEM) supplemented with GlutaMAX (Invitrogen) containing 10% heat-inactivated fetal calf serum (FCS; Summit Biotechnologies, Fort Collins, CO) plus 100 units/mL penicillin and 0.10 mg/mL streptomycin (Sigma) (α-MEM/FCS). The cell suspension was plated onto 35 mm tissue culture dishes at 2 mL/dish or onto 60 mm dishes at 6 mL/dish and incubated for 2 h at 37 °C in a humidified 5% CO₂ atmosphere. Nonadherent cells were removed by washing the plates 4 times with PBS, and the cultures were then incubated overnight in fresh α-MEM/FCS. The mean protein content of RPM cultures (35 mm dish) was 100 ± 10 µg/dish ([8.2 ± 0.8] x 10⁵ cells/dish]. Cultures of RPMs were washed twice with PBS at 37 °C and then overlaid with 1 mL of fresh α-MEM containing 25 µM arachidonic acid or arachidonoyltaurine. Cells were incubated with 25 µM arachidonic acid or arachidonoyltaurine for either 30 s or 15 min. The medium was removed and extracted with 3 volumes of ethyl acetate containing 0.5% acetic acid and 1 µM arachidonic acid-d₆. Extracts were dried under argon and resuspended in 1:1 methanol/5 mM ammonium acetate (pH 6.9).

**LOX Product Characterization.** LC was conducted on a Surveyor Separation Module using a Luna C18(2) column (50 x 2 mm, 3 µm) equipped with a C18 (2) Security Guard. The mobile phase consisted of 5 mM ammonium acetate at pH 6.9 (solvent A) and 3% A in acetonitrile (solvent B), and flow was constant at 0.3 mL/min. The gradient increased from 30% B to 90% B over 5 min, followed by a hold at 90% B for 4 min. Flow from the column passed first through a UV detector set to monitor 236 nm before proceeding to the electrospray ionization source. MS and MS/MS were conducted on a Quantum triple quadrupole mass spectrometer. The instrument was operated in negative-ion mode using the electrospray ionization source. MS and MS/MS were monitored in negative mode from 30% B to 90% B over 5 min, followed by a hold at 90% B for 4 min. Flow from the column passed first through a UV detector set to monitor 236 nm before proceeding to the electrospray ionization source. MS and MS/MS were conducted on a Quantum triple quadrupole mass spectrometer. The instrument was operated in negative-ion mode using the electrospray ionization source. MS and MS/MS were monitored in negative mode.

**Protein Expression and Immunoblotting.** After removal of the medium from RPMs for product characterization, cells were scraped twice into 100 µL lysis buffer [50 mM Tris-HCl at pH 7.5 plus 150 mM NaCl, 4 mM ethylenediaminetetraacetic acid (EDTA), 50 mM NaF, 0.1 mM Na₃VO₄, 0.2% Triton X-100, 0.1% NP40, 0.5% sodium deoxycholate, 1 mg/mL 4-(2-aminoethyl)benzenesulfonyl fluoride, and 5 µg/mL each of antipain, leupeptin, chymostatin, and pepstatin] and stored at −80 °C. Protein concentrations were determined using the Bradford protein assay (Bio-Rad, Hercules, CA). Either 5 or 10 µg of protein from RPMs and lysis buffer (as a control) were subjected to SDS–PAGE. Proteins were then transferred to nitrocellulose membrane (0.2 µm), which was treated with Miser Antibody Extender Solution (Pierce, Rockford, IL) prior to blocking with 5% nonfat milk protein in 20 mM Tris-HCl and 100 mM NaCl (pH 7.5) containing 0.1% Tween-20 (TTBS). 12/15-LOX was detected following incubation with a rabbit polyclonal antiserum directed against purified murine recombinant leukocyte 12-LOX (Cayman Chemical, Ann Arbor, MI) diluted 1:3000 in TTBS with 0.1% milk protein; it should be noted that this antibody cross-reacts with p112-LOX and r15-LOX-1 from other species. FAAH was detected following incubation with a rabbit polyclonal antibody directed against a synthetic peptide of rat FAAH amino acids 561–579 conjugated to KLH (GE Healthcare, Piscataway, NJ) diluted 1:4000 in TTBS with 0.1% milk protein. After the membranes were overlaid with ECL detection reagents (GE Healthcare, Piscataway, NJ), the membranes were exposed to Hyperfilm ECL (GE Healthcare, Piscataway, NJ) to obtain photographic images.

**RESULTS**

**Oxygenation of Arachidonoyltaurine by COXs and LOXs.** COX-2 and several LOXs are able to metabolize a variety of arachidonoyl-containing substrates, including AEA, 2-AG, and NAGly (15, 22, 23). Given the structural similarity of these substrates, the ability of COXs and LOXs to metabolize arachidonoyltaurine was investigated. At a concentration of 25 µM, arachidonoyltaurine was a very poor substrate for COX-1 and COX-2, with oxygenation proceeding at a rate of less than 10% of that observed for arachidonic acid. In contrast, r15-LOX-1, p12-LOX, and lк12-LOX efficiently metabolized arachidonoyltaurine.

The kinetics of arachidonoyltaurine oxygenation by r15-LOX-1, p12-LOX, and lк12-LOX were determined by monitoring the formation of the conjugated diene system via absorbance at 236 nm. In some cases, the concentration dependence of rates could not be fit to the classical Michaelis–Menten equation, and vₘₐₓ/Kₐ was calculated from the SEM of vₘₐₓ and Kₐ. The concentration dependence of rates could not be fit to the Michaelis–Menten equation, and vₘₐₓ/Kₐ ± standard error of the mean (SEM) was estimated from the slope of linear regression. Concentration dependence could be fit to the Michaelis–Menten equation, and vₘₐₓ/Kₐ was calculated from Michaelis–Menten parameters. The SEM of vₘₐₓ/Kₐ was calculated from the SEM of vₘₐₓ and Kₐ.

### Table 1: Kinetic Parameters for Arachidonic Acid and Arachidonoyltaurine Metabolism by LOXs

<table>
<thead>
<tr>
<th>LOX</th>
<th>Vₘₐₓ/Kₐ (min⁻¹)</th>
<th>Vₘₐₓ/Kₐ (min⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>arachidonic acid</td>
<td>0.45 ± 0.08*</td>
<td>0.76 ± 0.10*</td>
</tr>
<tr>
<td>arachidonoyltaurine</td>
<td>0.38 ± 5*</td>
<td>0.41 ± 1*</td>
</tr>
</tbody>
</table>

*Concentration dependence of rates could not be fit to the Michaelis–Menten equation, and vₘₐₓ/Kₐ ± standard error of the mean (SEM) was estimated from the slope of linear regression.
of arachidonic acid for all LOX isoforms examined. In contrast to r15-LOX-1 and lk12-LOX, pl12-LOX exhibited apparent substrate inhibition at high concentrations of arachidonoyltaurine (Figure 2). Inhibition was at least partially abrogated by inclusion of low concentrations of 13-hydroperoxyoctadecadienoic acid (13-HpODE).

**Characterization of Products of Arachidonoyltaurine Oxygenation.** CID of oxidized lipids can provide detailed structural information and is particularly useful for assigning regiochemistry of oxygenation (40, 41). Lipid hydroperoxides are highly susceptible to in-source fragmentation, resulting in the neutral loss of water. To reduce ambiguity resulting from in-source fragmentation, samples were treated with sodium thiosulfate to reduce hydroperoxides to the corresponding alcohols. The reduced products were subjected to analysis by LC-UV and tandem mass spectrometry (MS/MS). In addition, a standard of arachidonoyltaurine (m/z410) was subjected to CID to establish fragment ions characteristic of the taurine moiety (Figure 3). Such fragment ions were observed at m/z 80 (sulfonate), 107 (ethenesulfonate), and 124 (taurine). Additional fragment ions resulting from cleavage along the arachidonoyl chain were also observed. It was anticipated that the LOX reaction should not alter the taurine moiety or C-1–C-4 of the acyl chain, and as such, fragment ions at m/z 80, 107, 124, 151, 165, 178, and 192 should serve as a signature for the taurine moiety in LOX products.

Incubation of arachidonoyltaurine with r15-LOX-1 and h15-LOX-2 yielded similar product profiles, generating one major mono-oxygenated product (m/z 426) with a retention time of 5.4 min and strong absorbance at 236 nm (Figure 4 and 5A). CID of m/z 426 produced fragmentation patterns consistent with those reported for HETEs (40) but bearing characteristic fragment ions of the taurine moiety. In the case of r15-LOX-1 and h15-LOX-2, fragment ions with m/z 354 and 326 were observed (Figure 5B). These ions are characteristic of fragmentation α to an alcohol at C-15.

12S-LOXs gave rise to quite different product profiles. pl12-LOX produced two distinct mono-oxygenated products with strong absorbance at 236 nm (Figure 4). The product at 5.4 min fragmented to give CID spectra similar to that for 15-HETE-T (Figure 5B). CID of the product at 5.6 min yielded a unique fragmentation pattern (Figure 5C). In this case, fragment ions with m/z of 314 and 286 were characteristic of fragmentation α to the alcohol of 12-HETE-T. Thus, the mass spectra suggested that the two major products were 15-HETE-T (～30%) and 12-HETE-T (～70%). The elution order of the putative 15- and 12-HETE-T is consistent with that observed for arachidonic acid products 15- and 12-
HETE (42). In contrast, lk12-LOX generated 12-HETE-T with no detectable 15-HETE-T.

In addition to HETE-T, LC-MS analysis revealed that each LOX produced one or two chromatographically distinct dioxygenated products with \( m/z \) 442 (Figure 6). These products appeared to account for less than 10% of the total products for r15-LOX-1, h15-LOX-2, and pl12-LOX. In contrast, diHETE-T accounted for almost 40% of total products formed by lk12-LOX. Positions of dioxygenation could not be definitively assigned by LC-MS/MS.

**Oxidative Metabolism of Arachidonoyltaurine in RPMs.** RPMs provide a well-characterized system for the study of cellular uptake and metabolism of arachidonate-derived substrates. The profile of eicosanoid production, its modulation by various stimuli, and the expression of many arachidonate-metabolizing enzymes are well-established in this system. RPMs were incubated with either arachidonic acid or arachidonoyltaurine for 30 s. Medium was removed and extracted, and extracts were analyzed by LC-UV-MS and MS/MS. As anticipated from *in vitro* studies, no prostanoids derived from arachidonoyltaurine were observed in incubations with RPMs. Within 30 s of incubation with arachidonoyltaurine, significant amounts of mono-oxygenated products were observed (Figure 7A). Similar results were obtained with arachidonic acid (data not shown). All mono-oxygenated products absorbed strongly at 236 nm and yielded fragmentation patterns corresponding to HETE-derived species. In the case of arachidonoyltaurine, the major metabolite was 12-HETE-T, although small amounts of 15-HETE-T were also observed. A 12/15-LOX was detected by immunoblotting. There was no evidence for oxygenation of arachidonoyltaurine at C-5.

Prolonged incubation of RPMs with arachidonic acid or arachidonoyltaurine afforded only modest increases (<20%) in mono-oxygenated products (Figure 7B). However, a dioxygenated product was observed following incubation of

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**FIGURE 5:** MS and CID of mono-oxygenated products of arachidonoyltaurine metabolism *in vitro*. (A) Full-scan MS of the product at 5.4 min from incubation of arachidonoyltaurine with r15-LOX-1. Products at 5.4 and 5.6 min from enzymatic incubations afforded similar full-scan MS. (B) CID of the product at 5.4 min. (C) CID of the product at 5.6 min. The proposed structure and fragmentation are also shown; ions in italic were also observed for arachidonoyltaurine (refer to Figure 3).

**FIGURE 6:** Selected ion chromatograms of dioxygenated products (\( m/z \) 442) of *in vitro* arachidonoyltaurine oxidation by LOXs. The substrate was incubated with each indicated enzyme for 15 min. Hydroperoxides were reduced *in situ* by the addition of sodium dithionite, and products were extracted and analyzed by LC-UV-MS. The extracted chromatograms for \( m/z \) 442 are shown, and the absolute ion intensity is shown at the left of each panel. Absolute ion intensities for HETE-T were on the order of \( 3-8 \times 10^7 \) for each enzyme.
RPMs with arachidonic acid or arachidonoyltaurine for 15 min. For arachidonic acid, the product with m/z 335 appeared to be oxygenated at C-5 and C-12. However, because of the low abundance of the diHETE-T product, which accounts for less than 5% of the total products of arachidonoyltaurine oxidation, definitive assignment of the positions of oxygenation could not be made. There was no evidence of hydrolysis of arachidonoyltaurine to arachidonic acid in RPMs after 15 min, and FAAH was not detectable by immunoblot.

Arachidonic acid metabolism in RPMs is altered by inflammatory stimuli; LPS increases prostanoid biosynthesis via induction of COX-2, and zymosan activates 5-LOX, resulting in increased synthesis of leukotrienes (43–50). To assess the effect of these stimuli upon arachidonoyltaurine metabolism, RPMs were preincubated with either LPS for 6 h or zymosan for 10 min. Cells were then washed and incubated with arachidonoyltaurine for 15 min, and the metabolic profiles were compared to those of unstimulated macrophages. Unlike arachidonic acid, arachidonoyltaurine metabolism was not affected significantly by LPS or zymosan (data not shown), suggesting that arachidonoyltaurine is not a good substrate for COX-2 or 5-LOX in cells (Figure 8).

**DISCUSSION**

Inhibition of FAAH provides an attractive approach to modulating the endocannabinoid system, because it enhances the tonic actions of the endocannabinoids rather than indiscriminately activating all cannabinoid receptors. However, the recent discovery of NATs points to the possibility of another effector molecule mediating some biological actions of FAAH inhibition. The pathways for the generation and degradation of NATs are only now being defined.

The current studies were aimed at investigating the interactions of a polyunsaturated NAT with important lipid-metabolizing enzymes, COXs and LOXs. COX-2 but not COX-1 is able to metabolize a wide range of ester and amide derivatives of arachidonate (15, 22, 23). However, neither COX-1 nor COX-2 is able to use arachidonoyltaurine as a substrate. In contrast, 12S- and 15S-LOXs are able to oxygenate arachidonoyltaurine. In fact, pl12-LOX and r15-LOX-1 use arachidonoyltaurine more efficiently than arachidonic acid in vitro, and lk12-LOX exhibits similar efficiencies with arachidonate and arachidonoyltaurine.

Although arachidonoyltaurine is a very efficient substrate for pl12-LOX, the enzyme exhibited apparent substrate inhibition at high concentrations of arachidonoyltaurine. Although alkenyl sulfate substrates, including arachidonyl sulfate, are able to inhibit human 15-LOX-1, the mechanism of inhibition appears to be linked to aggregation of the substrate at high concentrations and can be eliminated by the addition of detergent (51). This is likely not the cause of inhibition by arachidonoyltaurine, because detergent was...
dominantly, 12-HETE accounts for 10\% of arachidonoyltaurine but not arachidonic acid in the concentration range examined. 

Commercially supplied fatty acids and fatty acid amides typically contain trace amounts of lipid hydroperoxides that arise from autoxidation of the allylic chain. Typically these trace hydroperoxides are sufficient for activation of LOXs, and the subsequent product of the lipoxygenase reaction is able to activate the enzyme further. However, in some instances, at high substrate concentrations, the lipid hydroperoxide product may be unable to efficiently compete with unreacted substrate for the resting, ferrous form of the enzyme, preventing autoactivation. Inclusion of exogenous lipid hydroperoxide should attenuate substrate inhibition, which is the case for inhibition of pl12-LOX by arachidonoyltaurine. Differences in substrate affinity may account for why substrate inhibition is observed with arachidonoyltaurine but not arachidonic acid in the concentration range examined.

LOXs display distinct product profiles and specificities of oxygenation. Although 15-LOX-1 produces 15-HETE predominantly, 12-HETE accounts for 10–20\% of total products of arachidonate oxygenation (28, 53). Likewise, lk12-LOX will oxygenate arachidonate primarily at C-12 and, to a lesser extent, C-15 (56). In contrast, the regiospecificity of arachidonate oxygenation of 15-LOX-2 and pl12-LOX is more tightly controlled, with a single HETE isomer being formed almost exclusively (35, 56–58). When arachidonoyltaurine was provided as the substrate, both r15-LOX-1 and h15-LOX-2 generated one major product, 15-HETE-T. Similarly, the monohydroxy product formed by lk12-LOX is predominantly 12-HETE-T, although this enzyme forms diHETEs as well. Thus, the fidelity of the oxygenation reaction for 15-LOX-1, 15-LOX-2, and lk12-LOX is retained, indicating that binding of the arachidonoyl chain is not significantly altered by the addition of the taurine moiety.

However, pl12-LOX displays dual specificity with arachidonoyltaurine, generating both 12- and 15-HETE-T. In combination with previous studies of pl12-LOX and its substrate specificity, this provides potentially interesting implications to the model of substrate binding to pl12-LOX (23). Substrate alignment within the LOX active site is a key determinant of the product profile. As discussed above, the lipoxygenase reaction is initiated by abstraction of a bis-allylic hydrogen atom by the nonheme ferric iron center. The depth of the LOX substrate-binding channel determines the position from which hydrogen is most frequently abstracted (59–61). Arachidonic acid binds to 12- and 15-LOXs in a “tail-first” orientation, with the carboxylate binding at the opening of the channel, stabilized by an ion-pairing interaction (59, 62), and the acyl chain extending into the hydrophobic channel. Lk12-LOX and 15-LOXs efficiently use neutral esters and amides of arachidonate, indicating that a hydrogen bond can substitute for the ion-pairing interaction (29–32).

The ion-pairing interaction appears to be critical for pl12-LOX, because it cannot efficiently metabolize neutral arachidonoyl species. However, pl12-LOX readily oxygenates amides of arachidonate if the substrate possesses a negatively charged head group, such as is present in arachidonoyltaurine and the arachidonoyl amino acids NAGly, NAla, and NAGABA (23). Furthermore, with these extended, negatively charged substrates, pl12-LOX acts as a 12/15-LOX. For this to occur, the substrate must adopt two different positions in the channel, one with C-10 positioned for hydrogen abstraction (to generate 12-HETE) and one that is displaced out of the channel, such that C-13 is positioned above the iron center (to make 15-HETE). Thus far, this phenomenon of dual specificity of pl12-LOX has only been observed with negatively charged substrates of arachidonate, suggestive of a currently unidentified determinant of substrate binding for pl12-LOX.

For a substrate to be metabolized in vivo, the compound must be able to cross membranes and reach sufficient concentrations within intracellular compartments, where the potential metabolizing enzymes are located. Murine RPMs were used to study the cellular metabolism of arachidonoyltaurine. The eicosanoid metabolic pathways and the modulation of these pathways by inflammatory stimuli are well-established in this cell line (43–50). Furthermore, the lipoxygenase pathway represents a major metabolic pathway of arachidonate in unstimulated cells, although some prostaglandins are also observed. After incubation with arachidonoyltaurine, no COX-derived products were observed, even following preincubation with LPS to increase COX-2 expression. This is in agreement with in vitro studies, which demonstrated that arachidonoyltaurine is a poor COX substrate. However, arachidonoyltaurine is rapidly taken up by RPMs and enters the lipoxygenase pathway. Arachidonoyltaurine is first oxygenated at C-12 to give rise to 12-HpETE-T (Figure 8). On the basis of the position of oxygenation and Western-blotting results, this reaction is most likely catalyzed by lk12-LOX. 12-HpETE-T is subsequently reduced to 12-HETE-T by a peroxidase. Biosynthesis...
of diHETE-T proceeds more slowly and in lower yield. Preliminary data indicate that diHETE-T is likely derived from 12-HETE-T or its parent 12-HpETE-T. However, further investigation is required to unravel the exact mechanism of diHETE-T formation.

The importance of NATs is only now beginning to be explored. This study demonstrates that a polyunsaturated mechanism of diHETE-T formation.

Preliminary data indicate that diHETE-T is likely derived from 12-HETE-T and proceeds more slowly and in lower yield.

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