Human Airway Monohydroxyeicosatetraenoic Acid Generation and Mucus Release

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ABSTRACT The effects of 5-, 8-, 9-, 11-, 12-, and 15-monohydroxyeicosatetraenoic acid (HETE) (0.1–100 nM) on mucous glycoprotein release from cultured human airways were determined. Each of the HETE was an active secretagogue of mucus at concentrations >1–10 nM with 12- and 15-HETE, the most active. Both 5- and 9-hydroperoxyeicosatetraenoic acid (HPETE) were also active as secretagogues at 100 nM, although of somewhat lower potency. As cultured airways were capable of responding to HETE with mucous glycoprotein release, it was of interest to identify and quantitate airway HETE formation. Accordingly, airways were incubated with tracer quantities of [14C]arachidonate for 16–48 h, and the spontaneous formation of 5-, 12- and 11- and/or 15-HETE was measured by high-pressure liquid chromatography. Indeed, sizeable quantities of 11- and/or 15- > 5- > 12-HETE were generated. This HETE generation was increased by the addition of 25 μg/ml of arachidonate and was reduced somewhat after 18–21 d in continuous tissue culture. Reversed anaphylaxis of human airways using anti-human IgE markedly increased the HETE formation, resulting in the production of micromolar concentrations of 5- and 11- and/or 15-HETE. Thus, human airways not only are capable of responding to the presence of HETE with mucous glycoprotein release, but also generate (both spontaneously and in response to anaphylaxis) at least three species of HETE, and do so in quantities capable of acting as mucus secretagogues.

INTRODUCTION

Allergic asthma is thought to develop from immunologic stimulation of mast cells, the release and/or generation of the mediators of allergy, and the elicitation by these mediators of the pathologic responses causing airflow obstruction (1, 2). Of the pulmonary responses incurred by allergic provocation that contribute to asthma, mucus production appears to play an important role in determining both morbidity and mortality. To examine the mechanisms possibly contributing to mucus production from human airways, a model consisting of cultured human bronchi (3) secreting biosynthetically labeled (4) mucous glycoproteins has been developed (5). Recent studies with this model have revealed that mast cell degranulation and mediator production lead to increased mucus glycoprotein release (5). Of the mast cell-derived mediators, histamine (through an H-2 receptor) (5), several prostaglandins (6), a mixture of synthetic monohydroxyeicosatetraenoic acids (HETE)1 (6), pure 12-HETE (6), prostaglandin-generating factor of anaphylaxis (PGF-A) (7), and the slow-reacting substances of anaphylaxis (SRS-A), leukotriene C4 and D4 (LTC4 and LTD4) (8), are all secretagogues of mucus. Whereas the precise relative contribution of each of these mediator-secretagogues to allergen-induced mucus production is unknown, the extraordinary responsiveness of mucus secretion in vitro to the lipoxygenase products of arachidonate (the HETE and leukotrienes) suggests that these agents may be active in vivo as well.

Stimulation of rat mast cells leads to the generation of prostaglandin (PG)D2 and 12-HETE (9), while purified human lung mast cells may also generate SRS-A (10). Anaphylaxis of human lung tissue generates PGF2α, PGE2, PGF2, PGD2, and thromboxane A2 (11–

1 Abbreviations used in this paper: GC, gas chromatography; HETE, monohydroxyeicosatetraenoic acid; HPETE, hydroperoxyeicosatetraenoic acid; HPLC, high-pressure liquid chromatography; LTC4, leukotriene C4; LTD4, leukotriene D4; m/e, mass/energy ratio; PC, prostaglandin; PGF-A, prostaglandin-generating factor of anaphylaxis; SRS-A, slow-reacting substances of anaphylaxis.
and, as yet, little is known of the specific lipox- 
genase products. Both human and guinea pig lung are 
capable of converting arachidonic acid to HETE, 
human lung generating 15-HETE (14) and guinea pig 
lung 12-HETE (15). Certainly, human lung must also 
have the capability of generating 5-HETE, as acti-
vation of a 5-lipoxygenase is the initial step in SRS-A 
formation (16). However, the quantities and specific 
identification of lipoxygenase products of human air-
ways generated during anaphylaxis are unknown.

The purpose of this paper is to describe the specific 
HETE generated by human airways during anaphyl-
axis and to demonstrate the capability of these HETE 
to act as secretagogues of mucus.

METHODS

[14C]Arachidonic acid (200 mCi/mmol) was obtained from 
New England Nuclear, Boston, MA. All other reagents and 
their sources have previously been described (5, 6). 5-, 8-, 
9-, 11-, and 12-HETE and 5- and 9-hydroperoxypenta-
tetraenoic acid (HPETE) were kindly provided by Dr. F. 
Sun (Upjohn Co., Kalamazoo, MI). 15-HETE was prepared by 
Dr. R. W. Bryant, and Dr. J. Vanderhook, of George 
Washington University School of Medicine (Wash., DC).

Preparation of human airways for culture. Human lungs 
were obtained at surgery primarily from tumor resec-
tion. Normal-appearing airways 2–10 mm in diameter were 
fragmented into 3–5 mm replicates and cultured as described 
(5, 6). The airway explants were maintained in CMRL-1066 
medium with penicillin (158 μg/ml), streptomycin (250 μg/ 
ml), and aureomycin (25 μg/ml) in a controlled atmosphere 
chamber gassed with 45% O2, 50% N2, and 5% CO2, and 
icubated at 37°C (3).

Radiolabeling of mucous glycoproteins. Mucous glyco-
proteins were radiolabeled by incorporating [3H]glucosamine 
(1 μCi/ml) into the culture medium. Explants were initially 
icubated for 16 h in the absence of [3H]glucosamine, washed 
twice with media, and then incubated with [3H]glucosamine 
for a 16- to 20-h base-line period (period I). The period I 
explants were harvested, fresh culture media without 
[3H]glucosamine was added for an additional 4-h period 
(period II), and these supernatants were subsequently har-
vested. Glucosamine-labeled mucous glycoproteins were 
precipitated from the samples by adding an equal volume of 
95% ethanol. The precipitates were filtered through me-
tricul membrane filters (6), and the retained radioactivity 
was determined. Analysis by gel filtration of airway super-
natants before or after precipitation indicates that >90% of 
all macromolecular radiolabeled glycoproteins were precipi-
tated and filtered by this procedure.

Effects of HETE on mucous glycoprotein release. The effect of HETE on the release of [3H]glucosamine-labeled 
mucous glycoproteins was determined by adding these 
agents to cultures at the beginning of period II. A ratio of 
radiolabeled mucous glycoproteins released in period II/pe-
riod I was determined for each sample and termed the se-
cretory index. The effects of pharmacologic agents were 
determined by comparing the secretory indices of manip-
ulated samples to matched unmanipulated control samples. 
HETE were maintained in absolute methanol at −70°C and 
preserved immediately before use by evaporation to dryness 
under a stream of nitrogen and resuspension in cul-
ture media.

Generation of HETE by cultured human airways. Air-
ways to be used for the generation of HETE were cultured 
with [14C]arachidonic acid (0.05–1.0 μCi/plate) and/or un-
labeled arachidonic acid (Nu Chek Prep, Elysian, MN) (10– 
25 μg/plate) for 4–48 h. The culture supernatants were im-
mEDIATELY frozen and maintained at −70°C in pyrex glass 
tubes sealed under nitrogen. The hydroxy fatty acids were 
extracted from the pooled supernatants by reverse-phase 
chromatography on silicic acid columns (17). ODS Silica 
columns (10 × 10 mm, SepPaks, Waters Associates, Millipore 
Corp., Milford, MA) were washed sequentially with meth-
anol and water before the samples were applied. The super-
natants were mixed with appropriate amounts of deu-
terated 5-HETE, 11-HETE, and 12-HETE standards to give 
a final concentration of 50 ng/ml each. The pH was adjusted 
to 4.0 with 1 M H3PO4, and the medium was passed through 
the SepPak column. The column was washed with 5 ml of 
hexane and eluted with 10 ml of diethyl ether. The ether 
extract was evaporated to dryness with a stream of nitrogen. 
Recovery of HETE exceeded 90%.

High-pressure liquid chromatography (HPLC). The resi-
due was dissolved in 0.05 ml of methanol and purified by 
reversed-phase HPLC. HPLC was performed on a Varian 
Model 5000 liquid chromatograph (Varian Associates, Inc., 
Palo Alto, CA) linked to a Varichrom detector (Varian As-
.sociates, Inc.) operated at a wavelength of 235 nm. The 
mobile phase was delivered at a flow rate of 1 ml/min−1. 
The column was washed isocratically with methanol/water/
acetic acid (80:20:0.1) for 5 min, which was followed 
with a linear gradient to straight methanol developed over 30 min 
(18). The HETE fraction was collected and evaporated to 
dryness with a stream of nitrogen.

The dried residue was dissolved in 1 ml of diethyl ether/ 
methanol (10:1) and esterified with diazomethane. The 
methyl ester was converted to trimethylylated ether deriv-
avative by gas chromatography-mass spectrometry. Samples in which 14C]arachidonic acid were added were 
analyzed after gas chromatography by monitoring radio-
activity.

Gas chromatography (GC) and mass spectrometry. GC 
was carried out with a Varian 2700 (Varian Associates, Inc.) 
coupled to a Packard model 594 proportional counter (Pack-
ard Instrument Co., Inc., United Technologies, Downers 
Grove, IL). Mass spectrometry was carried out on a Hewlett 
Packard 5992 GC-MS (Hewlett-Packard Co., Palo Alto, CA) 
operated in the selected ion monitoring mode. The GC col-
umn was 1% SE-30 ultraphase on Chromasorb W (HP) main-
tained at 210°C. The signal of the masses at mass/energy 
ratio (m/e) 229 vs. 225 for 11- and/or 15-HETE, m/e 301 
vs. 295 for 12-HETE, and m/e 313 vs. 305 for 5-HETE were 
recorded. Quantitative estimates were computed by refer-
ce to calibration curves derived from appropriate stan-
dards. This procedure does not distinguish between 11- 
and 15-HETE and, therefore, the data will be presented for 
either or both products.

Octadeuterated (D12) HETE and unlabeled HETE stand-
dards were synthesized from 5,6,8,9,11,12,14,15-D12-arachi-
donic acid by photochemical oxygenation. They were sep-
parated and purified by preparative HPLC with a μPorasil 
column (Waters Associates). The synthetic method used was 
especially the same as that described previously (19, 20).
(We are grateful to Drs. C. L. Bundy, W. P. Schneider, 
R. C. Kelly, and N. A. Nelson of the Department of Exper-
imental Chemistry at The Upjohn Co. for their generous 
support.)

Comparison of supernatant vs. tissue HETE concen-
tration. To estimate the relative proportion of HETE that were 

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generated and remained associated with the airways in comparison with that released into the supernatant, airways were incubated for 4 h either in media alone or media containing 10 μg/ml of arachidonic acid. After completing the incubation, the tissue was resuspended in fresh medium, chilled to 4°C, and homogenized with a polytron kinematika (Brinkman Instruments, Inc., Westbury, NY) at maximum speed for 90 s. The homogenate was centrifuged (2,500 g, 15 min, 4°C) and the supernatant was decanted, frozen at −150°C in dry ice/acetone, and maintained at −70°C under nitrogen until assayed.

**Anaphylaxis of human airways.** Airways anaphylaxis was produced with rabbit anti-human IgE. Antiserum directed at an IgE myeloma (PS) was partially purified by ammonium sulfate fractionation and protein A-Sepharose affinity chromatography. This material was absorbed on an IgG immunosorbent column to remove anti-light chain activity, and it formed a precipitin band against IgE heavy chain; precipitin bands against light chains or other immunoglobulins by Ouchterlony analysis were not present. A 1:100 dilution of this purified antiserum produced 20–40% histamine release in vitro from both peripheral human lung or airways. Anaphylaxis of airways with the preparation was initiated at the onset of period II, used 1:100–1:400 dilutions, and involved 4-h incubations. Histamine was analyzed by the automated fluorometric assay (21). The identity of the fluorescent material as histamine was confirmed by diamine oxidase digestions (22).

**Statistical analysis.** The results for mucus release are expressed as percent change from control and represent the comparison of the secretory indices (period II/period I) of experimentally manipulated samples with matched controls. Data are presented as mean±SEM. The n indicates the number of separate experiments combined to generate the results. In each experiment, 5–10 culture plates were used to generate each experimental point. The data were analyzed by paired sample t tests with a P value of <0.05 considered statistically significant.

**RESULTS**

**Effects of HETEs on mucus release.** 5-, 8-, 9-, 11-, 12-, and 15-HETE (0.1–100 nM) were assessed for their effects on mucus release (Fig. 1); all of these compounds increased mucus release in a dose-dependent fashion as compared with control. Nine separate lung cultures have been exposed to these compounds, and increased mucus release has been produced in each instance with 1–100-nM concentrations. All the HETE tested were equipotent secretagogues at concentrations of 1 and 10 nM, whereas 12- and 15-HETE were the most effective stimulants at 100 nM (12-HETE = +73±6.7%; n = 4).

In two separate experiments, the effects of 5- and 9-HPETE on mucus release were studied. The only
significant increase in mucus release was observed at concentrations of 100 nM: 20±2.6% increase above control (P < 0.05) for 5-HPETE and 28±1.9% (P < 0.01) for 9-HPETE. Thus, these two HPETE preparations were somewhat less active than their corresponding hydroxy compounds.

**Generation of HETE by human airways in vitro.** The data indicated that cultured human airways secrete mucus in response to exogenously added HETE and HPETE. It was therefore of interest to examine the capacity of these same airway preparations to generate HETE. Incubation of airways with [14C]arachidonic acid for 16 h, followed by silicic acid chromatography and GC, revealed the presence in the supernatant of two major peaks of radioactivity: the earlier peak cochromatographed with arachidonic acid while the second peak fractionated with HETE standards. Deuterated specific HETE standards were added to the incubation media as carriers, and the samples were purified by SepPak and HPLC. Mass fragmentography (selective ion monitoring) clearly showed prominent signals at m/e 225, 295, and 305, which occurred at the identical retention times as the deutered standards at m/e 229, 301, and 313. Therefore, it is likely that 11-HETE and/or 15-HETE, 12-HETE, and 5-HETE were present in the incubation medium.

Analysis of tissue culture media alone or tissue culture media to which arachidonic acid was added and incubated for 16 h failed to reveal the presence of any detectable HETE. However, airways incubated for 4 h in the presence of 25 μg/ml of arachidonic acid generated 32 ng/sample of 11- and/or 15-HETE and 10 ng each of 5- and 12-HETE. Thus, airways make at least three separate HETE. In a separate experiment, airways cultured for a longer period (48 h) in the presence of small amounts of [14C]arachidonic acid (0.5 μCi) generated 100 ng/sample of 5-HETE, 500 ng of 12-HETE, and 1,400 ng of 11- and/or 15-HETE.

Supernatants from airway samples that were cultured 1–3 d generated considerably more HETE than did the samples cultured 2 wk. For example, on the second day of culture, one airway preparation stimulated with 25 μg/ml of arachidonic acid generated 1,200 ng of 11- and/or 15-HETE/sample, 500 ng of 12-HETE, and 120 ng of 5-HETE, while after 18 d in culture, the same airways under the same conditions generated only 90 ng of 11- and/or 15-HETE, 90 ng of 12-HETE, and 85 ng of 5-HETE. Thus, the quantity and relative ratio of HETE generated varied with the time in culture. By 21 d, only 5-HETE was still being produced in sizable quantities (8 ng/sample).

To compare the relative quantities of HETE released into the supernatant vs. those associated with the tissue, tissue incubated in buffer alone or buffer plus 25 μg/ml of arachidonic acid was homogenized, extracted, and assayed for HETE. The results of these homogenates were compared with the supernatants from the same samples handled in parallel. The lung tissue homogenates generated <10 ng each of 11- and/or 15-HETE, 5-HETE, and 12-HETE, while the supernatant contained >100 ng of each of the three HETE. Thus, the HETE generated during short-term culture are released into the supernatant.

**Effect of airway anaphylaxis upon mucus release and HETE formation.** In separate reports, both reversed anaphylaxis of airways with rabbit anti-IgE (anti-IgE) and the addition of exogenous arachidonic acid have been shown to stimulate increased mucus release (5, 6). It was of interest to examine the effects of the simultaneous administration of both stimuli upon mucus release (Table I). Anti-IgE provoked histamine release and a dose-related increase in mucus release, while arachidonic acid also acted as a secretagogue of mucus. The combination of agents induced somewhat more mucus release than either maneuver alone and did so without an accompanying increase in histamine release.

The effects of reversed anaphylaxis upon airway HETE generation was examined in three separate experiments involving three individual lung preparations (Table II). Media containing [14C]arachidonic acid cultured for 4 h in the absence of lung tissue had no detectable HETE while airways cultured for 4 h in the presence of [14C]arachidonic acid (1 μCi/plate) produced <5 ng each of the three HETE. However, anaphylaxis (1:100 dilution of anti-IgE) in each instance induced sizable increases in the production of 11- and/or 15-, 12-, and 5-HETE with the average amount generated being 163±51, 88±23, and 255±109 ng, respectively. Thus, anaphylaxis of human airways leads to HETE generation.

<table>
<thead>
<tr>
<th>Agent</th>
<th>Concentration</th>
<th>Mucus release (Increase above control)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arachidonic Acid</td>
<td>75 μg/ml</td>
<td>21±2.7</td>
</tr>
<tr>
<td>Anti-IgE</td>
<td>1:400</td>
<td>26±1.9</td>
</tr>
<tr>
<td>Anti-IgE</td>
<td>1:100</td>
<td>59±1.2</td>
</tr>
<tr>
<td>Arachidonic Acid plus Anti-IgE</td>
<td>75 μg/ml</td>
<td>67±4.2</td>
</tr>
</tbody>
</table>

* Histamine release for each sample was the following: control, 9±0.4 ng/ml; arachidonic acid, 9.0±0.2 ng/ml; anti-IgE (1:100), 21±4 ng/ml; arachidonic acid plus anti-IgE, 24±3 ng/ml. Anti-IgE (1:400) was not assayed. Each result is the mean±SEM of five replicate samples.

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TABLE II
Effects of Anaphylaxis on HETE Formation by Human Airways

<table>
<thead>
<tr>
<th>Tissue culture preparation</th>
<th>HETE (ng/ml)</th>
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<tbody>
<tr>
<td></td>
<td>11- and/or 15-</td>
</tr>
<tr>
<td>Media with [14C]arachidonic acid (1 μCi/plate) but no airways</td>
<td>0</td>
</tr>
<tr>
<td>Airways plus [14C]arachidonic acid</td>
<td>Exp A</td>
</tr>
<tr>
<td></td>
<td>Exp B</td>
</tr>
<tr>
<td></td>
<td>Exp C</td>
</tr>
<tr>
<td>Airways plus [14C]arachidonic acid plus anti-IgE (1:100)</td>
<td>Exp A</td>
</tr>
<tr>
<td></td>
<td>Exp B</td>
</tr>
<tr>
<td></td>
<td>Exp C</td>
</tr>
<tr>
<td>Mean±SEM</td>
<td>163±51</td>
</tr>
</tbody>
</table>

Experiments A, B, and C are three separate experiments using different lungs. All incubations were for 4 h.

DISCUSSION

In the course of IgE-mediated allergic reactions, mast cell and basophil activation lead to the formation of an array of lipid mediators derived predominantly from arachidonic acid. The specific eicosanoids generated are determined by the tissue involved in the allergic reaction. Thus, certain cells and tissues have specific profiles of eicosanoids that they generate. For instance, mast cells themselves produce predominantly PGD_2 and 12-HETE (19) while platelets produce thromboxane A_2 and 12-HETE (23). Human lung tissue is known to release PGE_2, thromboxane A_2, PGE_2, PGD_2, PGD_2, and the leukotrienes LTC_4, LTD_4, and LTE_4 (11-13, 24, 25). Although not previously documented, it was clear that lung was also capable of producing monohydroxy oxidative products of arachidonic acid as 5-HPETE, the first stage in leukotriene formation.

Evidence is presented herein that human airways produce at least three distinct HETE: 11- and/or 15-, 5-, and 12-HETE. The synthesis of these HETE occurs spontaneously, without the need for exogenous stimulation, or may be accelerated by adding exogenous arachidonic acid or by anaphylaxis. Airways maintained in culture for 48 h with only trace amounts of exogenous arachidonic acid generate sizable quantities of 11- and/or 15- > 12- > 5-HETE, producing microgram quantities in the process. This synthesis may be markedly increased by adding 10-25 μg/ml of arachidonic acid, although the relative concentrations of the individual HETE produced remains the same: 11- and/or 15- > 12- > 5-HETE. HETE production decreased with time in culture, so that by 3 wk only trace amounts of 5-HETE were still being produced. Accordingly, all the results reported herein derive from airways cultured <7 d.

Although the observation that airways are capable of converting arachidonic acid to at least three species of HETE extends and confirms earlier observations involving guinea pig lung (14) and human peripheral lung (13), the capacity of anaphylaxis to stimulate HETE formation from airways is a new and potentially important observation. In three separate lung cultures undergoing anaphylaxis, 5- > 11- and/or 15- > 12-HETE synthesis was appreciated. The quantities synthesized were between 88 and 250 ng/culture, easily achieving concentrations known to be capable of stimulating mucus secretion (6, 26) and influencing neutrophil degranulation (27) as well as possibly modulating other airway-related events. Whereas the precise roles these HETE play in airway physiology are not known, the capacity of each of these products to stimulate mucus glycoprotein secretion suggests that they may contribute to at least this facet of allergic airways disease.

In earlier work, the ability of arachidonic acid to stimulate mucus secretion from human airways was appreciated (6). The arachidonic acid appeared to act through interaction with lipoygenase enzymes, leading to the formation of HETE. In this earlier work, complex mixtures of synthetic oxidative products of arachidonic acid as well as biosynthetic 12-HETE were shown to be capable of stimulating mucus secretion. In this study, these observations are extended to include 5-, 9-, 11-12-, and 15-HETE as well as 5- and 9-HPETE. Thus, HETE at concentrations >1–100 nM can act as secretagogues of mucus, and these are the concentrations achieved by airways undergoing anaphylaxis.

Other eicosanoids, including a variety of PG and LTC_4 and LTD_4, are also secretagogues (6, 7). The PG act in vitro at micromolar or higher concentrations and, thus, may not be physiologically important. The leukotrienes, however, are potent secretagogues at nanomolar concentrations and, clearly, may also contribute to the mucus secretion accompanying allergic airways disease. Although the precise mechanisms controlling the initial steps in leukotriene formation are not clearly defined as yet, it appears that anaphylaxis of lung tissues is generally required to activate the specific pathways involved. HETE formation from lung tissue may therefore occur more readily than SRS-A formation, as we observed that simply culturing airway tissue or adding exogenous substrate was suf-
sificent to lead to sizable HETE formation. Thus, it is possible that HETE production by airways occurs more commonly than does leukotriene formation and, therefore, that HETE may play a larger role in mucus production (and other lung functions as well). We conclude that airways in culture produce at least three species of HETE, that this synthesis is accelerated by anaphylaxis, and that the HETE formed act as potent mucus secretagogues.

ACKNOWLEDGMENTS
The authors gratefully acknowledge the fine typing of Ms. Joni Stefanelli, the editorial assistance of Karen Leighty, and the cooperation of the pathology and surgical departments at the Washington Hospital Center, Sibley Hospital, Suburban Hospital, Holy Cross Hospital, Walter Reed Army Medical Center, Montgomery General Hospital, and Washington Adventist Hospital in providing surgical specimens for this study.

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