A broad-spectrum lipidomics screen of antiinflammatory drug combinations in human blood

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Current methods of drug screening in human blood focus on the immediate products of the affected pathway and mostly rely on approaches that lack sensitivity and the capacity for multiplex analysis. We have developed a sensitive and selective method based on ultra-performance liquid chromatography–tandem mass spectrometry to scan the effect of drugs on the bioactive eicosanoid lipidome in vitro and ex vivo. Using small sample sizes, we can reproducibly measure a broad spectrum of eicosanoids in human blood and capture drug-induced substrate redirevision and unexpected shifts in product formation. Microsomal prostaglandin E synthase-1 (mPGES-1) is an antiinflammatory drug target alternative to COX-1/-2. Contrasting effects of targeting mPGES-1 versus COX-1/-2, due to differential substrate shifts across the lipidome, were observed and can be used to rationalize and evaluate drug combinations. Finally, the in vitro results were extrapolated to ex vivo studies by administration of the COX-2 inhibitor, celecoxib, to volunteers, illustrating how this approach can be used to integrate preclinical and clinical studies during drug development.

Introduction

NSAIDs are among the most commonly prescribed drugs. They include aspirin, used in low doses for cardioprotection, and drugs such as ibuprofen and celecoxib, used for the relief of pain and inflammation (1). They target the prostaglandin (PG) G/H synthase enzymes, colloquially known as COXs (COX-1 and COX-2). Constitutively expressed COX-1 accounts largely for formation of PGs subserving “housekeeping” functions, such as thromboxane A2 (TxA2) in hemostasis and PGE2 and PGI2 in the maintenance of gastrointestinal epithelial integrity. Platelets express only COX-1, and suppression of platelet COX-1–dependent TxA2 by low-dose aspirin accounts for its cardioprotective effects (2). COX-2 is more readily regulated in its expression, especially by mitogens and cytokines, and accounts largely for elaboration of the PGE2 and PGI2 that mediate pain and inflammation. While some older NSAIDs, like ibuprofen, inhibit both COX-1 and COX-2, a series of COX-2–selective NSAIDs, including celecoxib, valdecoxib, and rofecoxib, were developed and brought to market. It was hoped that they would conserve the efficacy of mixed inhibitors while minimizing gastropathy, which was largely attributed to inhibition by NSAIDs of housekeeping PGs formed by COX-1. Indeed, the results of controlled trials ultimately supported this notion (3).

While suppression of COX-2–derived PGE2 and PGI2 explained the analgesic and antiinflammatory efficacy of these newer drugs, basic and clinical experimentation revealed that these same PGs, particularly PGI2, now formed in the cardiovascular system, afforded protective functions. Randomized trials ultimately demonstrated that COX-2–selective NSAIDs conferred a cardiovascular hazard, and rofecoxib and valdecoxib were withdrawn from the market (3). These discoveries shifted interest downstream in the PG biosynthetic pathway to the microsomal prostaglandin E synthase-1 (mPGES-1), the major source of PGE2 formation, as an alternative drug target. Here, studies in mice suggest that, rather than conferring cardiovascular risk by suppressing PGI2, inhibition of mPGES-1 might avoid this hazard or even confer cardiovascular benefit due to a shift of the PGH2 substrate to PGI2 synthase (Figure 1), augmenting PGI2 biosynthesis (4).

PGs and other products of arachidonic acid (AA) metabolism are formed in pM to fM amounts, acting locally as evanescent mediators (1). Furthermore, the capacity of tissues or cells to form these compounds, collectively termed eicosanoids, greatly exceeds actual rates of biosynthesis in vivo. Two
approaches to their analysis have informed drug development in this pathway. First, actual biosynthesis has been estimated by measurement (most accurately with mass spectrometry [MS]) of biologically inactive, but chemically stable, metabolites. The preferred target analyte is urine, as this avoids artefactual ex vivo cellular activation, is acquired noninvasively, and affords a time-integrated estimate of biosynthesis. A limitation of this approach is that the tissue and cellular origin of the measured metabolite is unknown, although this can be estimated indirectly in the case of the platelet (5).

A second approach is to study, in vitro or ex vivo, the capacity of a defined cell type to generate a particular eicosanoid. The power of this approach is best illustrated by measurement of the inactive Tx hydrolysis product, TxB2, in stimulated platelet-rich plasma (6). This approach, particularly combined with measurement of urinary Tx metabolites, elucidated the pharmacology of low-dose aspirin, affording the basis for clinical trials that established its cardioprotective properties (2).

Here, we describe a MS-based capacity-related assay, applied in vitro or ex vivo to stimulated human whole blood but encompassing a broad swathe of the eicosanoid lipidome, not just a single compound. This permits us, in remarkably small sample sizes, to scan for the consequences of substrate rediversion after blockade of different enzymes or shifts in product formation consequent to receptor blockade in the eicosanoid biosynthetic/response pathways. We demonstrate unexpected consequences across the lipidome of inhibition of COXs and mPGES-1 and of receptor blockade and reveal how this approach can be

Figure 1. Antiinflammatory drug targets from the COX and 5-lipoxygenase pathways. Arachidonic acid (AA) is metabolized by 3 major enzymatic cascades: COX (COX-1/-2), lipoxygenase (5-LOX, 12-LOX, 15-LOX), and cytochrome P450 (CYP). The 2 COX isoforms, COX-1 and COX-2, oxidize AA to the hydroperoxy-endoperoxide PGG2 and then reduce PGG2 to the unstable intermediate hydroxy-endoperoxide PGH2. PGH2 is transformed to eicosanoids PGI2, TxA2, PGE2, PGD2, and PGF2α by the tissue-specific isomerases prostaglandin I synthase (PGIS), thromboxane synthase (TxS), prostaglandin E synthases (cPGES, mPGES-1, mPGES-2), prostaglandin D synthases (L/H-PGDS), and prostaglandin F synthase (PGFS), respectively. Each eicosanoid acts at the corresponding receptor distributed in various tissues. The 5-LOX enzyme, together with the accessory protein 5-LOX–activating protein (FLAP), catalyzes the conversion of AA to 5-hydroperoxyeicosatetraenoic acid (5-HpETE) and then to the unstable intermediate leukotriene A4 (LTA4). 5-HpETE can be reduced by glutathione peroxidases into 5-HETE alcohol. The LTA4 hydrolase (LTA4H) metabolizes LTA4 to leukotriene B4 (LTB4), which acts at BLT1 and BLT2 receptors on target cells. Alternatively, LTA4 can be conjugated with glutathione (GSH) by LTC4 synthase (LTC4S) to yield cysteinyl leukotrienes LTC4, LTD4, and LTE4, which act at CysLT receptors. Enzyme inhibitors and receptor antagonists are shown in green, while drug targets are depicted in red. cPGES, cytosolic PGE synthase; mPGES-1 and mPGES-2, microsomal PGE synthase-1 and -2, respectively; L/H-PGDS, hematopoietic and lipocalin-type PGD synthases; IP, prostacyclin receptor; TP, thromboxane receptor; EP, E prostanoid receptor; DP, D prostanoid receptor; FP, F prostanoid receptor; EET, epoxyeicosatrienoic acid.
used to rationalize further evaluation of drug combinations and to scan for the biochemical consequences of combinatorial approaches to disruption of eicosanoid metabolism. Finally, using celecoxib as an example, we illustrate how the approach can be used to integrate preclinical and clinical studies by extrapolation of the in vitro results to whole blood ex vivo in patients dosed with the NSAID.

**Results**

**Perturbations of the human plasma lipidome triggering COX and lipoxygenase product formation.** Stimulation of whole blood with 100 μg/ml LPS at 37°C demonstrated that PGs and Tx peaked at 24 hours and remained elevated throughout the rest of the time course of observation (Figure 2A, Figure 3B, and Supplemental Figure 1; supplemental material available online with this article; doi:10.1172/jci.insight.87031DS1). The production of PGE2, rose to 21.5 ± 9.3 ng/ml (mean ± SD, n = 4, P = 0.004), while PGF2α and TxB2 averaged 5.7 ± 2.3 ng/ml (P = 0.004), respectively (Supplemental Figure 1, A–C, and Supplemental Table 3). In addition, LPS was a potent stimulus for formation of 15.3 ± 2.3 ng/ml (P = 0.01) total 15-hydroxyeicosatetraenoic acid (15-HETE) at 24 hours (Supplemental Figure 1E). Chiral analysis revealed that LPS induced both (S)- and (R)-enantiomers of 15-HETE, which averaged 8.98 ± 1.9 ng/ml (n = 7, P = 0.04) and 4.6 ± 1.2 ng/ml (P = 0.03), respectively (Supplemental Figure 7, A and B). If 15-HETE production derived solely from the 15-lipoxygenase (15-LOX) enzyme, 15(S)-HETE would have been overwhelmingly the dominant product. These observations are consistent with its formation by COX enzymes (7, 8) or by autoxidation of eicosanoids (9). The production of total 12-HETE was time dependent and robust, resulting in a concentration of 1,292 ± 190 ng/ml in activated blood by 48 hours (Supplemental Figure 1D); however,
LPS stimulation did not affect 12-HETE levels, suggesting that a prolonged incubation of whole blood alone at 37°C resulted in spontaneous 12-HETE release, most likely by activated platelets (10). Here, chiral analysis revealed that 12(S)-HETE was the predominant product, consistent with its formation by 12-LOX, and its formation was unaltered in human plasma by LPS (Supplemental Figure 8D). Finally, chiral analysis also revealed that LPS stimulated 11(R)-HETE production, whereas 11(S)-HETE, as well as both (S)- and (R)-enantiomers of 8-HETE, were not detected in whole blood (Supplemental Figure 8A). Production of 5-LOX–derived eicosanoids in the presence of LPS was at the lower limits of detection and was comparable to the vehicle control (data not shown). On the contrary, activation of whole blood with zymosan led to a time-dependent leukotriene (LT) and 5-HETE production, consistent with its effects on granulocytes, particularly neutrophil 5-LOX, in vivo (11, 12). Time-course and dose-response studies revealed that 125 μg/ml zymosan induced a robust increase in 5-LOX products that peaked at 4 hours and averaged 8.43 ± 4.12 ng/ml for LTB4, 9.62 ± 7.88 ng/ml for LTE4, and 15.1 ± 6.37 for 5-HETE (n = 22) (Figure 2B, Figure 3A, Supplemental Figure 2, and Supplemental Table 3). Costimulation with LPS and zymosan for 4 hours resulted in a lipid profile with effects comparable to those from zymosan alone, suggesting that, at early times, eicosanoid release was driven by zymosan regardless of LPS. At 24 hours, coincubation with both stimuli led to a combined profile, consisting of lipids triggered by zymosan early on and eicosanoids whose biosynthesis depended on induction by LPS over a long-term incubation (Figure 3, C and D, and Supplemental Figure 4).
A 4-hour incubation with the vehicle PBS resulted in detectable levels of PGF$_{2\alpha}$ (0.31 ± 0.34 ng/ml, mean ± SD, $n = 22$), TxB$_2$ (1.1 ± 0.42 ng/ml), 5-HETE (1.43 ± 0.6 ng/ml), 12-HETE (21.7 ± 20.6 ng/ml), 15-HETE (0.5 ± 0.2 ng/ml), and AA (2,613 ± 1,327 ng/ml). By 24 hours, most lipids were unaltered, but 12-HETE and AA increased 6-fold (141 ± 115 ng/ml) and 2-fold (5,216 ± 2,140 ng/ml), respectively (Supplemental Figure 3, A and B), consistent with continuous secretion of 12-HETE by platelets and a dynamic release of AA by all blood cell types. PGD$_2$ and PGI$_{2\alpha}$ production was not detected in human plasma. Amounts of 20-HETE did not differ between stimulated or vehicle-treated whole blood (Supplemental Figure 1F). Low (pg/ml) levels of 12-hydroxy-5,8,10E-heptadecatrienoic acid (12-HHT) were detected in plasma in response to LPS or zymosan, while a more robust production was detected in sera (12.38 ± 1.41 ng/ml, mean ± SD, $n = 4$), and inhibited by diclofenac, consistent with the origin of 12-HHT from the platelet-derived COX-1/thromboxane synthase (COX-1/TxS) axis (Supplemental Figure 5, C and D). The “specialized proresolving mediators,” resolvins D1 (RvD1) and D2 (RvD2), maresin 1, and protectin, were not detected in this study. Lipoxin A$_4$ (LXA$_4$) was not detected at baseline but could be induced by zymosan at 24 hours; while modest levels of resolin E1 (RvE1) were measured in untreated plasma, it was not induced by either stimulus (Supplemental Figure 5, A and B). The fatty acids eicosapentaenoic acid (EPA), docosahexaenoic acid (DHA), and AA were not significantly affected by whole blood stimulation (data not shown). Thus, for the purposes of increased LC-MS efficiency, incubation with authentic standards was reduced from 21 to 9 analytes (PGE$_{2\alpha}$, PGF$_{2\alpha}$, TxB$_2$, LTE$_{4\alpha}$, LTE$_{4\beta}$, 5-HETE, 12-HETE, 15-HETE, and AA). In summary, a 24-hour incubation of human whole blood with LPS (100 μg/ml) maximized activation of the COX-1/-2 pathways, while incubation with zymosan (125 μg/ml) for 4 hours yielded maximum activation of the 5-LOX pathway.

**Substrate rediversion consequent to inhibition of mPGES-1.** To determine how pharmacological blockade of specific targets in the COX and LOX pathways affects the human plasma lipidome in vitro, we screened the selective COX-2 inhibitors (COX-2i), celecoxib and GW406381; the COX-1/-2 inhibitor (COX-1/-2i), diclofenac; aspirin (at concentrations corresponding to low-dose (<100 mg/d) administration in vivo, which would preferentially target COX-1); and the E prostanoid receptor 4 (EP4) antagonist, CJ-042794, and compared their effects on the lipidome to those of the mPGES-1 inhibitor (mPGES-1i), MF-63 (Figure 4). We observed large interindividual variability in the capacity to form eicosanoids among donors in each treatment group (Supplemental Tables 4 and 5) that constrained comparison of absolute eicosanoid concentrations in studies of small sample size. However, expression of drug effects as a percentage of their own stimulation group (Supplemental Tables 4 and 5) that constrained comparison of absolute eicosanoid concentrations in studies of small sample size. However, when the same data were expressed as a percentage of stimulus control, a 60% reduction in PGE$_2$ relative to the untreated, stimulated control ($n = 5$, $P = 0.003$) was observed with 10 μM MF-63. Concurrently, there was an elevation in PGF$_{2\alpha}$ (by an average 50%, $P = 0.008$) and TxB$_2$ (by 37%, $P = 0.007$), consistent with rediversion of the PGH$_2$ substrate of mPGES-1 to PGF synthase (PGFS) and TxS, respectively. Although total 15-HETE was apparently elevated by approximately 10% on average ($P = 0.048$), this was not confirmed by chiral analysis (vide infra). The marginal effect of mPGES-1 inhibition on 15-HETE was in contrast to the effect of COX-1/-2 blockade. At 10 μM, the COX-2i, celecoxib ($n = 5$) and GW406381 ($n = 5$), and the COX-1/-2i, diclofenac ($n = 4$), profoundly reduced PGE$_2$ and PGF$_{2\alpha}$ (both by more than 90%, $P < 0.0001$), TxB$_2$ (by more than 80%, $P < 0.0001$), and total 15-HETE (by more than 70%, $P < 0.001$) (Figure 4, A–C). Chiral analysis confirmed the inhibitory effect of celecoxib on in vitro plasma levels of 15(S)-HETE and 15(R)-HETE, rather than solely the 15-LOX product, 15(S)-HETE. Suppression of 11(R)-HETE, but not the platelet 12-LOX product, 12(S)-HETE, was observed with celecoxib in vitro, in contrast to the mPGES-1i, MF-63, that had no effect on 11(R)-HETE (Supplemental Figure 8B). Neither 15(R)- nor 15(S)-HETE were suppressed by MF-63. Celecoxib also inhibited LTE$_4$ by 40% ($P = 0.02$) and LTE$_{4\beta}$ by 30% ($P = 0.043$) but had no effect on 5-HETE levels, suggesting that LT inhibition happened downstream of the 5-LOX enzyme. Similarly to COX-2i, aspirin ($n = 4$) inhibited PGF$_{2\alpha}$ (by 70%, $P < 0.0001$), TxB$_2$ (by 65%, $P < 0.0001$), and total 15-HETE (by 35%, $P = 0.0003$), but unlike the COX-2i, the low concentration of aspirin had no effect on PGE$_2$, suggesting its predominant formation from COX-2 in this setting (Figure 4D). Interestingly, aspirin potentiated total 12-HETE by 30% ($P = 0.01$), consistent with inhibition of platelet COX-1, making more substrate available for the platelet 12-LOX enzyme. Finally, the EP4 antagonist ($n = 5$) did not affect the plasma lipidome, suggesting that such a pharmacological perturbation did not result in feedback mechanisms to compensate for the lack of signaling through EP4 (data not shown). At their IC$_{50}$, MF-63 reduced PGE$_2$ and elevated
TxB2, while celecoxib and diclofenac reduced PGE2, PGF2α, and total 15-HETE (Supplemental Figure 6 and Supplemental Table 5). Similar trends were observed after targeted inhibition of the 5-LOX pathway (Supplemental Figure 9). Inhibition of LTA4 hydrolase (LTA4H) resulted in a dose-dependent reduction in LTB4 (by 25%, \( P = 0.05 \), and 90%, \( P = 0.0001 \), at IC50 and 10 \( \mu M \), respectively, \( n = 5 \)), with a concurrent elevation in LTE4 levels (by 38%, \( P = 0.02 \), and 170%, \( P = 0.01 \), at IC50 and 10 \( \mu M \), respectively, \( n = 4 \)). Such an effect is consistent with shunting of the common substrate LTA4 from hydrolysis to LTB4 toward the enzyme LTC4 synthase (LTC4S), which converts LTA4 and glutathione to LTC4 and then subsequently forms LTE4.

As expected, inhibition of both 5-LOX and 5-LOX–activating protein (FLAP) reduced levels of LTB4, LTE4, and 5-HETE. Interestingly, antagonism of the BLT2 (\( n = 5 \)), but not the BLT1 (\( n = 5 \)), receptor for LTB4 (by 20%, \( P = 0.02 \) and LTE4 (by 22%, \( P = 0.0009 \)) at their IC50s, but not at the fixed concentration of 10 \( \mu M \) (Supplemental Figure 9).

Treatment of whole blood with the vehicle DMSO for 4 hours led to detectable levels of total 5-HETE (0.7 ± 0.1 ng/ml), 12-HETE (13.5 ± 12 ng/ml), 15-HETE (0.4 ± 0.1 ng/ml), and AA (1,405 ± 238 ng/ml). Similarly to PBS, prolongation of incubation to 24 hours did not further alter 5-HETE and 15-HETE, while 12-HETE and AA increased 10-fold (134 ± 85 ng/ml) and 2-fold (2,724 ± 84 ng/ml), respectively (Supplemental Figure 3, C and D). DMSO constituted 0.4% of total sample volume and did not trigger hemolysis or otherwise visible cell toxicity.
Overall, despite only 4 to 5 donors in each experiment, the approach captured dose-dependent effects of the studied compounds, demonstrated substrate shunting within and between both COX and LOX pathways, and segregated the lipidomic profiles consequent to whole-blood perturbation with COX-1/-2 versus mPGES-1 inhibitors.

Screening of combinatorial approaches to antiinflammatory drug therapy. Considering that products of AA metabolism from both COX and LOX pathways may contribute to inflammation (13, 14), dual inhibitors of 5-LOX and COX cascades have been investigated (15). Here, we tested how coincubations of a COX-2i, an mPGES-1i, or an EP4 antagonist with inhibitors of specific components of the 5-LOX cascade (5-LOX, FLAP, or LTA4H) would affect the plasma lipidome. Results for combinations with the mPGES-1i demonstrate common features of this approach (Figure 5). Time-dependent effects on plasma lipidome were clearly evident. A 4-hour coincubation of the mPGES-1i with the FLAP inhibitor (FLAPi) (n = 5) resulted in almost complete abrogation of LT formation (P < 0.0001) and an 80% reduction in 5-HETE levels (P < 0.0001), consistent with inhibition of the FLAP, while total 12-HETE and 15-HETE were elevated by 85% (P = 0.02) and 20% (P = 0.006), respectively (Figure 5A). However, these elevations were only transient and undetected by 24 hours (Figure 5B). At 24 hours, the inhibitory effect on 5-LOX products was sustained and PGE2 was reduced on average by 52% (P < 0.0001), while PGF2α and TxB2 were increased by 53% (P = 0.002) and 28% (P = 0.0004), respectively. The latter effects on PGs and Tx are characteristic of mPGES-1 inhibition by MF-63 (Figure 4). In contrast to combination with the FLAPi, coincubation of MF-63 with the 5-LOXi ABT-761 did not result in elevation of plasma lipids but rather had profound inhibitory effects on various eicosanoids. A 4-hour cotreatment with the 5-LOXi resulted in a reduction in LTs (by 95%, P <
5-HT (by 76% \( P < 0.0001 \)), 5-HETE (by 76% \( P < 0.0001 \)), and PGE\(_2\) (by 60% \( P < 0.0001 \)) and a mild and transient decrease in PGF\(_2\alpha\) (by 14% \( P = 0.004 \)). Total 12-HETE and 15-HETE were also inhibited on average by 54% \( P = 0.002 \) and 16\% \( P = 0.02 \) at 24 hours, accordingly, which might be due to nonspecific effects of the 5-LOXi on 12-LOX and on COX-2 or 15-LOX after a prolonged incubation. Chiral analysis to make the latter distinction was not performed. Surprisingly, with this drug combination, we did not observe the elevation in PGF\(_2\alpha\) and TxB\(_2\) observed with mPGES-1 inhibition by MF-63 alone (Figure 4).

Coincubation of MF-63 with the LTA\(_4\)Hi SC-57461A for 4 or 24 hours reduced LTB\(_4\) by 90\% \( P < 0.0001 \) but elevated LTE\(_4\) by more than 200\% \( P = 0.0006 \) consistent with the LTA\(_4\)H inhibition observed in a single-compound screen (Supplemental Figure 9). Similar to the combination with the FLAP, short-term coincubation of MF-63 with the LTA\(_4\)H inhibitor (LTA\(_4\)Hi \( n = 5 \)) led to transient increases of total 12-HETE (by 52\% \( P = 0.03 \)) and 15-HETE (by 16\% \( P = 0.02 \)) that had dissipated by 24 hours, by which time PGF\(_2\alpha\) and TxB\(_2\) had increased by 44\% \( P = 0.03 \) and 38\% \( P = 0.015 \), respectively. PGE\(_2\) was reduced throughout the whole experiment by more than 50\% \( P = 0.001 \) (Figure 5).

The effect of drug combinations on plasma lipidome was stimulus related (Supplemental Figure 10), consistent with different blood cell types that are activated and engaged in the response to drug inhibition.

Validation of the in vitro whole-blood assay in an ex vivo setting. We validated our in vitro assay by its application ex vivo after administration of celecoxib to healthy volunteers. A single oral dose led to a reduction in the ex vivo LPS-stimulated plasma levels of PGE\(_2\) (49.9\% \( \pm 27\% \) of pre-dose control, mean \( \pm \) SD, \( P = 0.001 \), \( n = 10 \)), PGF\(_{2\alpha}\) (45.5\% \( \pm 20.5\% \), \( P = 0.0005 \)), TxB\(_2\) (52.2\% \( \pm 16.9\% \), \( P = 0.0002 \)), and total 15-HETE (49\% \( \pm 18.7\% \), \( P = 0.003 \)) (Figure 6). These findings are consistent with the in vitro celecoxib results (Figure 4). Chiral analysis of the ex vivo samples showed that celecoxib tended to inhibit both 15(S)-HETE and 15(R)-HETE, although the data reached significance ex vivo only with the (S)-enantiomer (Supplemental Figure 7). Here, despite suppression of 11(R)-HETE by celecoxib in vitro, it was not suppressed by celecoxib ex vivo (Supplemental Figure 8). Urinary metabolites (M), PGI-M and PGE-M, indices of systemic PGI\(_2\) and PGE\(_2\) biosynthesis, respectively, were reduced in the celecoxib-treated group (Supplemental Figure 11). However, urinary Tx-M was comparable to placebo control, indicating that although the capacity of blood cells to generate Tx in vitro and ex vivo was modestly reduced, actual biosynthesis of Tx was not
significantly altered by administration of a COX-2i as previously observed (16). Indeed, consistent with an effect on platelet capacity, we have previously reported that celecoxib inhibits serum TxB<sub>2</sub> ex vivo but to a degree insufficient to inhibit platelet aggregation (17). Urinary metabolites of PGD<sub>2</sub> and the isoprostane, iPF<sub>2</sub>α-VI, were not affected by celecoxib treatment (data not shown).

**Discussion**

A broad-spectrum, targeted lipidomics approach based on ultra-performance liquid chromatography–tandem MS (UPLC-MS/MS) is reported. It is a rapid, sensitive, analytical method that allows simultaneous profiling and high-throughput, quantitative analysis of COX and LOX products in human whole blood in vitro and ex vivo. Using small sample sizes, we can reproducibly screen compounds targeting specific eicosanoid biosynthetic enzymes or receptors for their dose-dependent effects, mechanisms of action, and combinatorial potential. The method was validated by extrapolating in vitro data to ex vivo results after in vivo administration of a COX-2i to volunteers. Thus, the method can be applied to proof-of-concept experiments in drug development and can be used to integrate preclinical and clinical studies.

Differential perturbations of the lipidome with LPS and zymosan allow for scanning a wide range of diverse products of the COX and LOX pathways. The whole-blood model provides near physiological conditions in a plasma protein-rich environment and in the presence of blood cells that participate in transcellular biosynthesis of eicosanoids (18). Stimulus-dependent effects of drug combinations on the plasma lipidome vividly depict how activation of different blood cell types affects the response to drug inhibition. The capacity of cells to generate a particular class of eicosanoids is limited by their ability to respond to a particular agonist and the availability of the relevant enzymatic machinery to engage in single-cell or transcellular lipid biosynthesis (18, 19). Human peripheral blood monocytes express both COX isoforms and provide a major source of eicosanoids in capacity-related assays in plasma (20, 21). Monocyte-derived TxB<sub>2</sub> and PGE<sub>2</sub> largely depend on the induction of COX-2 by exogenous agonists, such as LPS (20, 21). Polymorphonuclear neutrophils respond to LPS with induction of COX-2 (22–24). Zymosan also induces neutrophil COX-2 expression and production of TxB<sub>2</sub> and PGE<sub>2</sub> (25). Platelets express COX-1 and 12-LOX and are the major source of TxA<sub>2</sub> and 12-HETE in serum. Besides TxA<sub>2</sub>, platelet TxS catalyzes production of 12-HHT, a potential ligand for the LTB<sub>4</sub> receptor BLT2 (26, 27).

Although this high-throughput assay quantitates precisely a range of lipids, it is not designed to attribute them to their cellular origins in a complex matrix, such as whole human blood. Nevertheless the combination of drug intervention and chiral analysis allows for some tentative conclusions. Suppression of induced levels of TxB<sub>2</sub> and 12-HHT and elevation of 12-HETE by aspirin at concentrations corresponding to peak plasma levels after administration of low doses, points to these compounds being products of the COX-1 pathway in platelets. While induced total 15-HETE was suppressed by COX-2i, this applied to both enantiomers, not just 15(S)-HETE, the product of 15-LOX, which is expressed in monocytes and eosinophils (28–30). 15(R)-HETE is increased by LPS with kinetics that correspond to COX-2 induction and is suppressed, along with 11(R)-HETE, by celecoxib, consistent with its formation by COX-2. 15(S)-HETE may also be formed by COX-2 (7). However, while the nonenzymatic products 8(R)-HETE and 8(S)-HETE were not detected in human plasma, the possibility that some of 15(S)-HETE was made via this route cannot be excluded. A more likely explanation is that it may be the product of eosinophil 15-LOX, activated by formation of COX-2–derived eicosanoids in monocytes (31). Celecoxib, diclofenac, and the COX-2i GW406381 may thus suppress both 15-HETE enantiomers in stimulated whole blood by directly inhibiting COX-2, indirectly suppressing eosinophil 15-LOX, and also by curbing LPS-triggered reactive oxygen species generation (32). While 11(R)-HETE can be formed by both COX isoforms (7, 33), it was induced by LPS in plasma, unlike 12-HHT and 12(S)-HETE, and suppressed by COX-2i in vitro, all of which is consistent with it being derived predominantly from COX-2. The failure of celecoxib to reduce 11(R)-HETE ex vivo may reflect the variance between individuals relative to the signal, even when adjusted for their own controls. Overall, the suppressive effects of the COX-2i on 15- and 11-HETEs in human whole blood are marks of distinction from the lipidomic effects of mPGES inhibition.

Expression of the 5-LOX protein coincides with FLAP, LTA<sub>4</sub>H, and LTC<sub>4</sub>S and is restricted to cells of myeloid origin, including neutrophils, eosinophils, monocytes/macrophages, and mast cells (19). In whole blood, neutrophils are the primary source of LTs and 5-HETE; peripheral blood mononuclear cells and, to a lesser degree, eosinophils may also contribute (34). Zymosan is a naturally occurring proinflammatotry ligand that elevates intracellular calcium and thus triggers LT synthesis in cells possessing the 5-LOX.
enzyme (35). Although LPS does not directly activate 5-LOX synthesis, LPS can prime neutrophils for increased production of LTB₄ in response to subsequent stimulation with zymosan (36). Thus, it is possible that costimulation of whole blood with LPS and zymosan in our study potentiated the capacity of neutrophils to produce LTs. Platelets and red blood cells do not possess 5-LOX or FLAP, but they contain secondary enzymes, such as LTA₄H or LTC₄S, and may contribute to LT production through transcellular biosynthesis (18). Despite the capacity-related feature of this assay, with provocation by inflammatory stimuli, only modest amounts of LXA₄ and RvE1 were evident, while most specialized proresolving mediators were not detected. We and others have also largely failed to detect endogenous production of these mediators in plasma or urine (37, 38) and failed to relate alterations in specialized proresolving mediator formation in vivo to the evocation of inflammation with LPS or its resolution or the dose-dependent provision of their theoretical substrate (38).

Expression of data as a percentage of a subject’s own stimulus control permits assessment of drug effect on the plasma lipidome in small sample sizes, despite variability in absolute values between donors. Standardization of data for individual responses lets us capture evidence consistent with substrate rediversion during screening of drugs targeting mPGES-1 or LTA₄H. Substrate shunting or rediversion may occur when inhibition of one terminal synthase makes more of the common substrate PGH₂ available for other PG synthases (Figure 1). Targeting of a terminal PG synthase can also alter AA substrate availability to other pathways. Thus, we observed that blockade of mPGES-1 inhibited PGE₂ production but elevated products of PGFS, TxS, and, potentially, 15-LOX. Similarly, LTA₄H substrate may be shunted between LTA₄H and LTC₄S in the 5-LOX pathway (Figure 1). Here, inhibition of LTA₄H resulted in reduced levels of LTB₄ but augmentation of LTC₄S product LTE₄.

Considering the involvement of both COX and LOX pathways in common inflammatory diseases, such as atherosclerosis (13, 14), abdominal aortic aneurysm (39–41), asthma (42, 43), arthritis, and inflammatory pain (13, 44, 45), dual inhibitors of both pathways are of interest (46). Concurrent blockade of mPGES-1 with inhibition of various components of the 5-LOX pathway revealed both predicted and unexpected effects on the human plasma lipidome. Coincubations of an mPGES-1i with inhibitors of FLAP or LTA₄H led to initial shunting of the substrate toward 12-LOX and possibly 15-LOX, with a subsequent rediversion toward the COX pathway (Figure 4). In the combination with the FLAPI, sustained inhibition of 5-HETE, LTs, and PGE₂ coincided with a concurrent increase in PGF₂α and TxB₂, characteristic of the mPGES-1i. Thus, each drug retained features of their distinct mechanism of action when combined in whole blood. Surprisingly, when the mPGES-1i was combined with the 5-LOXi, which would be expected to recapitulate the effect of FLAP inhibition, there was early suppression of both PGs and 5-LOX products, with later reductions in total 12-HETE and 15-HETE but no evidence of substrate rediversion. Thus, combination of the mPGES-1i with compounds that have apparently similar mechanisms of action, FLAP and 5-LOX inhibitors, has distinct effects on the plasma lipidome.

In summary, using a small-scale in vitro whole-blood assay and broad-spectrum lipidomics analysis, we can study drug effects on the COX and LOX pathways in plasma samples of limited quantity. The method complements in vivo analysis of eicosanoid metabolites and MS-based analysis of target modification, such as acetylation of COX-1 by aspirin (47). It can be applied to in vitro screening of drug combinations and can be readily extended for the quantitation of other eicosanoids, as the system is applicable to lipid metabolites with diverse hydrophobicity.

**Methods**

**Reagents.** Deuterated analogs of PGE₂, PGD₂, PGF₂α, 6-keto-PGF₁α (the hydrolysis product of PGI₂), TxB₂ (the hydrolysis product of TXA₂), LTE₄, LTB₄, 5-HETE, 12-HETE, 15-HETE, 20-HETE, AA, EPA, DHA, 7-hydroxy-5,11-diketotetranorprostane-1,16-dioic acid (PGE-M), 11,15-dioxo-9α hydroxy-, 2,3,4,5-tetranorprostanoic acid, 2,3-dinor TxB₂, 2,3-dinor 6-keto PGF₁α (PG-M), isoprostane 8,12-iso-IP₂α-VI, and creatinine and authentic standards for 11(α)-HETE, 11(β)-HETE, (±)12-HETE, and (±)15-HETE were purchased from Cayman Chemical ([d]-PGE₂, [d]-PGD₂, [d]-PGF₂α, [d]-TxB₂, [d]-LTB₄, [d]-LTE₄, [d]-5-HETE, [d]-12-HETE, [d]-15-HETE, [d]-AA, [d]-EPA, [d]-DHA, [d]-tetranor-PGEM, [d]-tetranor-PGDM, [d]-11-dehydro-TxB₂, [d]-8,12-iso-IP₂α-VI, and [d]-creatinine). Stock solutions were prepared in acetonitrile and stored in glass vials at ~80°C. We utilized purified deionized water (Milli-Q water purification system, EMD Millipore) in the preparation of all aqueous solutions and mobile phases. UPLC-grade acetonitrile and ethyl acetate were purchased from...
the 5-LOXi ABT-761, the FLAPi MK-0591, or the LTA4Hi SC-57461A. To ensure a concurrent, maximum
Thus, COX-2i celecoxib, mPGES-1i MF-63, and EP4 antagonist CJ-042794 were coincubated, each with
a compound targeting the COX pathway was coincubated with a compound targeting the 5-LOX pathway.
Fisher Scientific) immediately after the blood draw, served as untreated controls. For drug combinations,
the addition of zymosan. Blood samples, spun down in polypropylene tubes (Sarstedt, 5 ml, 75
μg/ml final concentration) for 4 hours in the presence of 2
μg/ml final concentration of inhibitors were simulated with both LPS and zymosan at 4 and 24 hours. Each test compound was delivered
alone were combined in drug combination screening experiments. Thus, whole blood containing two types
of inhibitors was simulated with a compound targeting the 5-LOX pathway. Thus, COX-2i celecoxib, mPGES-1i MF-63, and EP4 antagonist CJ-042794 were coincubated, each with the 5-LOXi ABT-761, the FLAPi MK-0591, or the LTA4Hi SC-57461A. To ensure a concurrent, maximum
activation of the COX and 5-LOX pathways, optimal stimulating conditions for each of these pathways alone were combined in drug combination screening experiments. Thus, whole blood containing two types of inhibitors was simulated with both LPS and zymosan at 4 and 24 hours. Each test compound was delivered in 1 μl DMSO to the well bottom, avoiding mixing of the drugs, to the final concentration of 10 μM for each compound. Then, 500 μl of heparinized blood was added to the well and incubated for 30 minutes at 37°C. After that, 5 μl LPS and 5 μl zymosan were added to the final concentrations of 100 μg/ml and 125 μg/ml, respectively, and incubated for 4 and 24 hours. Drug combinations stimulated with either LPS
or zymosan alone served as internal controls for stimulating conditions. For the in vitro assay, data were normalized to plasma volume and expressed as a percentage of patient’s stimulus + DMSO control using the formula: percentage of stimulus control = \( \frac{C_{\text{sample}}}{C_{\text{stimulus}}} \times 100\% \), where \( C \) represents eicosanoid concentration in ng/ml. For the ex vivo assay, data were normalized to plasma volume and expressed as a percentage of patient’s pre-dose control using the formula: percentage of pre-dose control = \( \frac{C_{\text{post-dose}}}{C_{\text{pre-dose}}} \times 100\% \), where \( C \) represents eicosanoid concentration in ng/ml.

**Analytical method development for broad-spectrum eicosanoid quantification in human plasma.** A UPLC-MS/MS assay was developed to measure a wide range of COX and LOX products in plasma in a single run. In addition to \( \text{PGE}_2 \), inhibition of which contributes substantially to the therapeutic effects of NSAIDs (13), we were interested in other COX products as possible products of substrate rediversion after \( \text{mPGES-1} \) inhibition in whole blood. Thus, we focused on \( \text{PGD}_2 \), \( \text{PGF}_2\alpha \), 6-keto-\( \text{PGF}_1\alpha \), \( \text{TxB}_2 \), and 12-HHT. Considering that substrate shifts could occur beyond the COX pathway, for instance, toward LOX or cytochrome P450 (CYP) cascades, we measured 5-HETE and 15-HETE, products of 12- and 15-LOX, respectively, and 20-HETE, an AA-derived metabolite made by CYP enzymes. We also measured resolvins (RvD1, RvD2, RvE1), lipoxin A4, maresin 1, and protectin. We measured AA levels to assess how different stimulating conditions and inhibition with drugs affect the substrate in the whole-blood matrix. Finally, we were interested to determine whether activation of blood cells affected plasma levels of EPA and DHA. Thus, our initial mix of standards contained 17 compounds to monitor 21 analytes. Extensive kinetics (0, 0.5, 1, 2, 4, 8, 24, 32, and 48 hours) and stimulus dose-response experiments were done to establish optimal conditions for activation of COX and LOX pathways in human whole blood. For COX activation, we selected bacterial endotoxin LPS, a known trigger of PG production in whole blood (60, 61). For LOX activation, we selected a yeast-derived TLR2 agonist, zymosan. Once optimal conditions for COX and LOX pathways were determined, the spike was reduced to 9 standards.

**UPLC-MS/MS analysis of plasma samples.** The analysis was performed on a Waters ACQUITY UPLC system in-line with a Waters Xevo TQ-S Triple Quadrupole Mass Spectrometer. Plasma samples (200 \( \mu \)l) were spiked with stable isotope-labeled internal standards ([d]$_4$-PGE$_2$ [5 ng]; [d]$_4$-PGF$_2$$_\alpha$ [2.5 ng]; [d]$_4$-TxB$_2$ [10 ng]; [d]$_4$-LTB$_4$ [1 ng]; [d]$_4$-LTE$_4$ [2.5 ng]; [d]$_4$-12-HETE [25 ng]; [d]$_4$-15-HETE [1 ng]; [d]$_4$-AA [2,500 ng]; [d]$_5$-RvD1 [0.2 ng]; [d]$_5$-RvD2 [1.25 ng]; [d]$_5$-LXA$_4$ [1 ng]) in 900 \( \mu \)l of acetonitrile. Then, they were acidified by addition of formic acid to a final concentration of 1%. The samples were vortexed, incubated at room temperature for 15 minutes, vortexed again, and sonicated for 1 minute. After that, they were spun down, and the supernatant was transferred to Phree cartridges for phospholipid and protein removal (Phenomenex, 8B-S133-TAK). The samples were eluted with a slight vacuum (<20 kPa). They were then dried under a gentle stream of nitrogen at the ambient temperature and reconstituted with 25 \( \mu \)l of methanol.

Before injection, 25 \( \mu \)l of water was added to each sample and an aliquot of 20 \( \mu \)l was injected into a C18 UPLC column (Waters ACQUITY UPLC BEH 2.1 \times 150 mm \times 1.7 \mu \)m) and eluted at a flow rate of 350 \( \mu \)l/min, with a linear gradient from 20% solvent B to 90% in 20 minutes. Mobile-phase and MS parameters were optimized to obtain maximum sensitivity for respective product ions. Mobile phase A consisted of water/mobile phase B, 95:5 (v/v), with 0.5% formic acid; mobile phase B consisted of acetonitrile/methanol, 95:5 (v/v), with 0.5% formic acid. The UPLC system was directly interfaced with the negative-mode electrospray ionization (ESI) source of the mass spectrometer using multiple reaction monitoring (MRM) of all transitions (Supplemental Table 2). A single run took 26 minutes. Quantitation was done by peak area ratio, and results were normalized to plasma volume.

**LC-MS/MS analysis of urine samples.** Systemic urinary production of PGE$_2$, PGD$_2$, TxA$_2$, and PGI$_2$ was determined by quantification of their major urinary metabolites (PGE$_M$, PGD$_M$, TxA$_M$, PGI$_M$), and isoprostane 8,12-iso-iPF$_{20 \alpha}$-VI was measured as an index of lipid peroxidation. Stable isotope-labeled internal standards ([d]$_4$-tetranor-PGEM [25 ng]; [d]$_4$-tetranor-PGDM [25 ng]; [d]$_4$-2,3-dinor-6-keto-PGF$_{1 \alpha}$ [5 ng]; [d]$_5$-11-dehydro-TxB$_2$ [5 ng]; [d]$_8$-8,12-iso-iPF$_{20 \alpha}$-VI [5 ng]) were added to 1 ml of human urine in 50 \( \mu \)l of acetonitrile and allowed to equilibrate for 15 minutes. 500 \( \mu \)l of MO HCl was added and allowed to equilibrate at pH 3 for 30 minutes. The samples were purified by solid-phase extraction using Strata-X 33 \mu \)m polymeric reversed phase, 30 mg/ml cartridges (Phenomenex, 8B-S100-TAK). The SPE cartridge was conditioned with 1 ml of acetonitrile and equilibrated with 0.25 ml of water. The sample was applied to the cartridge, which was then washed with 1 ml of water and dried with a vacuum for 15 minutes. The analyte
endogenous creatinine (m/z 114.0 > 86.0 amu, 15.0 eV) and [d3]-creatinine (m/z 117.0 > 89.0 amu, 15.0 eV) were analyzed using MRM of a Thermo Electron Corporation TSQ Quantum Ultra tandem instrument (Thermo Fisher Scientific) equipped with a triple quadrupole analyzer. The HPLC system (Thermo Scientific Accela Pump) was directly interfaced with the positive-mode ESI source of the mass spectrometer. Mobile phase A consisted of 100% AcN; mobile phase B was 5 mM ammonium formate (pH 3.9). The flow rate was 350 μl/min. Separations were carried out with various linear solvent gradients. The Thermo Finnigan TSQ Quantum Ultra tandem instrument was operated in negative-mode ESI, and the analyzer was set in the MRM mode for the analysis of the urinary metabolites. To adjust for urine production, data were normalized to creatinine levels. Here, 20 μl of urine was added to 1 ml of [d3]-creatinine in AcN. Next, 40 μl of spiked urine was further diluted with 360 μl of AcN, and 20 μl of the sample was injected into a HPLC column (Waters XBridge BEH HILIC 2.1 × 50 mm × 2.5 μm). Samples were eluted at a flow rate of 350 μl/min with an isocratic 12% solvent B in 1.5 minutes. Mobile phase A was 100% AcN; mobile phase B consisted of 5 mM ammonium formate (pH = 4). The HPLC system was directly interfaced with the positive-mode ESI source of the mass spectrometer (Thermo Electron Corporation TSQ Quantum Ultra) using MRM of endogenous creatinine (m/z 114.0 > 86.0 amu, 15.0 eV) and [d3]-creatinine (m/z 117.0 > 89.0 amu, 15.0 eV). Urine data are reported as a percentage of the volunteer’s own pre-dose control using the formula: percentage of pre-dose control = (Cpost-dose/Cpre-dose) × 100%, where C represents metabolite concentration in ng/mg creatinine.

Chiral UPLC–electron capture atmospheric pressure chemical ionization/high-resolution MS analysis. Chiral products of 8-HETE, 11-HETE, 12-HETE, and 15-HETE were quantified by a targeted chiral lipidomics approach using UPLC–electron capture atmospheric pressure chemical ionization/high-resolution MS (ECAPCI/HRMS) methodology based on a technique reported previously with a Thermo Scientific Q-Exactive HF high-resolution mass spectrometer replacing the triple quadrupole instrument (62). The pentfluoro-2-benzyl (PFB) derivatives of 8 lipids and a heavy isotope analog internal standard of 15(S)-HETE were prepared by dissolving the residues from extracted plasma in 100 μl diisopropylethylamine in acetonitrile (1:19, v/v) followed by 50 μl PFB-Br in acetonitrile (1:9, v/v). The solution was incubated at 60°C for 30 minutes, allowed to cool, evaporated to dryness under a nitrogen stream at room temperature, and redisolved in 100 μl hexane/ethanol (97:3, v/v) for chiral UPLC-HRMS analysis. A 5-μl aliquot was injected for each sample. Normal-phase chiral chromatography was performed using an UltiMate 3000 binary UPLC equipped with a refrigerated autosampler (6°C) and a column heater (35°C). Gradient elution was performed in the linear mode. A Chiralpak AD-H column (250 × 4.6 mm i.d., 5 μm; Chiral Technologies) was employed with a flow rate of 1 ml/min. Solvent A was hexane and solvent B was 2-propanol/methanol (5:5, v/v). The linear gradient was as follows: 2% B at 0 minutes, 2% B at 3 minutes, 8% B at 11 minutes, 8% B at 13 minutes, 95% B at 17 minutes, 2% B at 18 minutes, and 2% B at 22 minutes. MS was conducted on a Thermo Scientific Q-Exactive HF Hybrid Quadrupole-Orbitrap mass spectrometer. The column effluent was diverted to waste 3 minutes before and 14 minutes after the run. The mass spectrometer was equipped with an APCI source in the negative electron-capturing ion mode (62). The operating conditions were as follows: vaporizer temperature, 450°C; heated capillary temperature, 320°C; corona discharge needle, set at 30 μA. The sheath gas (nitrogen) and auxiliary gas (nitrogen) pressures were 40 psi and 10 arbitrary units, respectively. The S-lens was set at 60. The Q-Exactive-HF instrument was alternating between a full scan (100–600 m/z) at a resolution of 30,000 and parallel reaction monitoring (PRM) at 60,000 resolution with a precursor isolation window of 2 m/z and normalized collision energy of 20%. Molecular (M-PFB) precursor ions for 15-HETE and [d3]-15(S)-HETE were 319.2323 and 321.2781, respectively. The most intense product ions for 15-HETE and [d3]-15(S)-HETE were 216.1372 and 208.1815, respectively. The PRM transitions for all chiral products are summarized in Supplemental Table 2. Quantification was based on the product ion with m/z ± 3 ppm of theoretical. Data analysis was performed using Xcalibur software, version 2.0 SR2 (Thermo Scientific), from raw mass spectral data. Calibration standard samples were prepared with charcoal-stripped FBS. Calibration samples were spiked with authentic standards of 11(S)-HETE, 11(R)-HETE, (±)-12-HETE, and (±)-15-HETE in the amounts of 0, 0.1, 0.25, 0.5, 1, 2.5, 5 ng, and 1 ng of the internal standard [d3]-15(S)-HETE. Oxidized lipids were extracted, purified, derivatized, and analyzed as described above for the analytical samples. Calibration curves were plotted using a linear regression of
peak area ration of analytes against internal standard (Supplemental Figure 12). Typical regression lines for
11(R)-HETE, 11(S)-HETE, 12(R)-HETE, 12(S)-HETE, 15(R)-HETE, and 15(S)-HETE were $y = 4.60723x + 1.46655$ ($r^2 = 0.9959$), $y = 4.44665x + 1.38262$ ($r^2 = 0.9914$), $y = 1.09631x + 0.748153$ ($r^2 = 0.9959$), $y = 1.06212x + 1.09477$ ($r^2 = 0.9850$), $y = 0.338872x + 0.189865$ ($r^2 = 0.9955$), and $y = 0.752994x + 0.393947$ ($r^2 = 0.9986$), respectively. Concentrations of the chiral products were calculated by interpolation from the calculated regression lines.

**Statistics.** Statistical analyses were performed using GraphPad Prism software version 5.0 for Mac OS X. The statistical significance of the differences between various treatments was determined by 1-sample, 2-tailed $t$ tests for the in vitro assays and by unpaired, 2-tailed $t$ tests for the ex vivo assay. $P < 0.05$ was considered significant. Plots for plasma and urine data from the ex vivo study were built using GraphPad Prism software. The investigators were blinded to sample allocation during the ex vivo study and outcome assessment. Visualization of broad-spectrum lipidomic changes in in vitro stimulated and drug-treated whole blood was generated using the open-source Circos software package (63).

**Study approval.** All clinical studies were conducted according to Declaration of Helsinki principles. The study protocols were approved by the Institutional Review Board of the University of Pennsylvania and by the Advisory Council of the Clinical and Translational Research Center of the University of Pennsylvania. All study volunteers provided written informed consent prior to inclusion in the study.

**Author contributions**

LLM, TG, ER, and GAF designed the study and experiments. LLM, ER, and GAF analyzed data. LLM performed the experiments and coordinated the clinical studies. JAL developed the LC-MS method and analyzed MS data for the in vitro study. XL analyzed MS data for drug combinations experiments and the ex vivo study. CM and IAB developed and performed chiral UPLC-ECAPCI/HRMS analyses. GG created Circos plots. LLM and GAF wrote the manuscript.

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