A Novel Autotaxin Inhibitor Reduces Lyosphosphatidic Acid Levels in Plasma and the Site of Inflammation

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ABSTRACT

Autotaxin is the enzyme responsible for the production of lysophosphatidic acid (LPA) from lysophosphatidyl choline (LPC), and it is up-regulated in many inflammatory conditions, including but not limited to cancer, arthritis, and multiple sclerosis. LPA signaling causes angiogenesis, mitosis, cell proliferation, and cytokine secretion. Inhibition of autotaxin may have anti-inflammatory properties in a variety of diseases; however, this hypothesis has not been tested pharmacologically because of the lack of potent inhibitors. Here, we report the development of a potent autotaxin inhibitor, PF-8380 [6-(3-(piperazin-1-yl)propanoyl)benzo[d]oxazol-2(3H)-one] with an IC50 of 2.8 nM in isolated enzyme assay and 101 nM in human whole blood. PF-8380 has adequate oral bioavailability and exposures required for in vivo testing of autotaxin inhibition. Autotaxin’s role in producing LPA in plasma and at the site of inflammation was tested in a rat air pouch model. The specific inhibitor PF-8380, dosed orally at 30 mg/kg, provided >95% reduction in both plasma and air pouch LPA within 3 h, indicating autotaxin is a major source of LPA during inflammation. At 30 mg/kg PF-8380 reduced inflammatory hyperalgesia with the same efficacy as 30 mg/kg naproxen. Inhibition of plasma autotaxin activity correlated with inhibition of autotaxin at the site of inflammation and in ex vivo whole blood. Furthermore, a close pharmacokinetic/pharmacodynamic relationship was observed, which suggests that LPA is rapidly formed and degraded in vivo. PF-8380 can serve as a tool compound for elucidating LPA’s role in inflammation.
lating the mammalian K<sup>?</sup>2P<sub>2</sub>1 (KCΝK2, TREC-1) channel (Cohen et al., 2009) and β-fiber-mediated spinal transmission, which underlies neuropathic pain (Xie et al., 2008).

Autotaxin is found in three main isoforms, each with a different splice variant. The most stable and common β isoform is a 95-kDa protein secreted by inflammatory cells and responsible for the production of LPA by the hydrolysis of LPC (produced through secretory phospholipase A2) (Giganti et al., 2008).

Autotaxin has been shown to be responsible for the production of LPA in plasma. Autotaxin heterozygote knockout mice have a 50% reduction of circulating LPA compared with wild-type mice (van Meeteren et al., 2006). Early autotaxin inhibitors have shown that plasma LPA, generated from endogenous LPC by incubating whole blood for several hours, can be nearly 100% inhibited, suggesting that autotaxin is the primary enzyme responsible for LPA production (Ferry et al., 2008). These inhibitors, however, were not suitable for in vivo use because of their limited solubility and bioavailability.

One critical question remains: does inhibition of autotaxin reduce LPA levels at the site of inflammation? And if so, what are the consequences of that inhibition?

To date, no autotaxin inhibitors have been developed that regulate LPA levels in vivo; however, there have been several reports of inhibitors that modulate autotaxin in vitro or in whole blood. Inhibitors that have been described are based on substrate analogs (Prestwich et al., 2008). These compounds are weak autotaxin inhibitors and are also LPA1 receptor agonists/antagonists. Recently, S32826 [(4-tetradecanoylamo)benzyl]phosphonic acid disodium salt has been described as a potent inhibitor of autotaxin in vitro and in whole blood, but it is not bioavailable or useful for in vivo work (Ferry et al., 2008).

In this article we describe the development of an autotaxin inhibitor that modulates LPA levels in vitro and in vivo through the direct inhibition of autotaxin. The compound is capable of blocking inflammation-induced LPA synthesis, both in plasma and at the site of inflammation. This will allow further testing of the hypothesis that disruption of the autotaxin/LPA axis will have beneficial anti-inflammatory effects that could affect pain and an array of diseases such as RA and cancer.

Materials and Methods

Materials

LPC and LPA were obtained from Avanti Polar Lipids (Birmingham, AL), all standard buffer reagents and solvents were obtained from Sigma-Aldrich (St. Louis, MO).

tert-Butyl 4-(3-oxo-3-(2-oxo-2,3-dihydrobenzo[d]oxazol-6-yl)propyl)piperazine-1-carboxylate. 6-(3-chloropropanoyl)benzo[d]oxazol-2(3H)-one (19.87 g, 94.4 mmol) was added dropwise. The solution was slowly warmed to RT overnight. The solution was then quenched with water (300 ml), and the resulting precipitate was filtered. In a separate flask (3,5-dichlorophenyl)methanol (13.0 g, 72.26 mmol) was dissolved in DMF (50 ml), and sodium hydride (3.96 g, 99.0 mmol) was added. The solution was stirred for 10 min and then cooled to 0°C. The nitro-precipitate was dissolved in DMF (50 ml) and then added dropwise to the (3,5-dichlorophenyl)methanol solution. The solution was stirred for 10 min and then quenched with a small amount of water to ensure quenching of excess sodium hydride. The DMF was distilled off to give a bright yellow oil, which was purified by flash chromatography (5% MeOH/dichloromethane) to give PF-8380 an off-white solid (14.56 g, 30.65 mmol, 42%). 1H NMR (400 MHz, DMSO-d<sub>6</sub>) δ ppm 7.83–7.89 (2 H, m), 7.57 (1 H, s), 7.42 (2 H, s), 7.18 (1 H, d, J = 7.9 Hz), 5.07 (2 H, s), 3.37 (3 H, br. s.), 3.19 (2 H, t, J = 7.1 Hz), 2.89 (1 H, s), 2.73 (1 H, s), 2.70 (2 H, t, J = 7.1 Hz), 2.42 (4 H, t, J = 4.7 Hz) C13 NMR (400 MHz, DMSO-d<sub>6</sub>) 197.35, 154.71, 153.98, 143.37, 141.20, 135.90, 134.03, 130.81, 127.36, 126.08, 124.97, 109.33, 108.77, 74.64, 72.83, 52.25, 43.47, 35.38, mz = 478.1.

Murine Autotaxin Cloning

The full-length murine autotaxin cDNA was purchased from Open Biosystems (Huntsville, AL). The following primers purchased from Integrated DNA Technologies (Coralville, IA) were used to introduce cloning sites HindIII and NotI by polymerase chain reaction to allow cloning into pcDNA3.1 Zeo: 5′-AGC TAG CTA AGC TTA TGG CAA GAC AAG GCT GTG TT'C′ and 5′-ATG ACT GAC TGC GGC CGC TCA TTA GTG ATG ATG ATG GTG ATG ATG AAT CCT GCT CCT ATC TGT ATG CAG G3′. The 5′ primer introduced the addition of a His<sub>6</sub> tag to the C terminus of the protein. The murine autotaxin gene was amplified by using Accucerp Pfx DNA polymerase (Invitrogen, Carlsbad, CA). The DNA sequence of the final construct was confirmed by sequencing performed at Cogenics (Houston, TX).

Autotaxin Expression

Ten-liter batches of human and murine autotaxin-conditioned media were prepared by using a HEK 293 FreeStyle (Invitrogen) and Wave (GE Healthcare, Little Chalfont, Buckinghamshire, UK) bioreactor transient expression system. Approximately 2.5 × 10<sup>8</sup> HEK cells were seeded into a Wave Bioreactor CellBag20 containing 5 liters of 293 FreeStyle media. The bag was maintained at 37°C, 8% CO<sub>2</sub>, and a constant rocking of 25.5 rpm. When the cell density reached 2 × 10<sup>7</sup>/ml, 4.5 liters of fresh 293 FreeStyle media was added. The cells were transfected by the sterile addition of 640 ml of Opti-MEM (Invitrogen) containing 10 mg of plasmid DNA and 13 ml of 293fectin (Invitrogen). The bioreactor was maintained under the initial growth conditions for another 6 days after transfection, and the media were harvested for purification.
containing autotaxin with purity more than 95% were pooled, concentrated, and stored at −80°C.

Recombinant Enzyme Assay
FS-3 substrate (Echelon L-2000; Echelon Biosciences, Logan, UT) was solubilized in assay buffer at 500 μM and frozen at −20°C in single-use aliquots for up to 4 weeks. Recombinant autotaxin was diluted in Tris-buffered saline (140 mM NaCl, 5 mM KCl, 1 mM CaCl₂, 1 mM MgCl₂, 50 mM Tris, pH 8.0) and incubated with compound diluted 3:7 with 66% methanol/water plus 0.1% triethylamine further in preparation for LC-MS/MS. Autotaxin LPC assay was published method by Chen et al. (2008). In brief, 50 μg of recombinant autotaxin was diluted in Dulbecco’s modified Eagle’s medium (DMEM) and added to compound in DMSO or DMSO alone (final 1% DMSO), and the reaction was started with the addition of FS-3 at a final concentration of 1 μM. The reaction was allowed to proceed at 37°C for 30 min and monitored at 520 nm until the uninhibited control compared with a no-enzyme control gave a Z’ >0.5. IC₅₀ values were determined in triplicate by using a four-parameter fit.

Recombinant Enzyme Assay Using LPC LC-MS/MS
C17:0 LPC and C16:0 LPA were dissolved in MeOH (10 and 100 mg/ml, respectively) and maintained at −20°C. Recombinant autotaxin was diluted in Dulbecco’s modified Eagle’s medium (DMEM) and added to compound in DMSO or DMSO alone (final 1% DMSO), and the reaction was started with the addition of LPC at a final concentration of 15 μM/ml followed by incubation at 37°C for 30 min. The reaction was stopped by the addition of two volumes of organic (50:50:1 acetoniitrile/methanol/acetic acid + 15 ng/ml C16:0 LPC, internal standard) and either frozen at −80°C or diluted further in preparation for LC-MS/MS. Autotaxin LPC assay was diluted 3:7 with 66% methanol/water plus 0.1% triethylamine (TEA) and analyzed by LC-MS/MS.

Fetal Fibroblast Cell Assay
Human fetal fibroblasts were prepared from foreskin and cultured as described previously (Hawley-Nelson et al., 1980; Raz et al., 1989). Cells were grown in DMEM containing 10% fetal bovine serum, 4 mM l-glutamine, 25 mM HEPES, 100 U/ml penicillin, and 100 μg/ml streptomycin (Invitrogen) and were passed when confluent. The cells were suspended in fetal bovine serum-free DMEM and plated at density of 5 × 10⁴ cells/well on 96-well culture plates (Corning Glassworks, Corning, NY). After 24 h of incubation, various concentrations of compound were added with tyrosine protein phosphatase inhibitor cocktail (Millipore Bioscience Research Reagents, Temecula, CA). After 15-min incubation, the substrate C17:0 LPC (final 15 μg/ml) was added to start the autotaxin enzyme reaction. After 30 min at 37°C, the supernatant of cell culture was collected and mixed with organic solution, acetonitrile/methanol/acetic acid (50:50:1, v/v/v) (EMD Chemicals, Gibbstown, NJ), to stop the reaction. C17:0 LPA content was measured by LC-MS/MS.

Autotaxin Inhibition in Human Whole Blood
Venous blood from healthy human donors was collected in heparinized tubes. Blood (0.5 ml) was incubated with compound dissolved in DMSO (1% final concentration of DMSO) in 96-well collection plates for 2 h at 37°C. The reaction was stopped by centrifugation at 800g for 10 min to pellet the cells. Plasma supernatant (100 μl) was precipitated with three volumes of methanol/acetoniitrile/acetic acid (60:40:1) and spun at 4,000g for 10 min. Supernatants were recovered and diluted. LPA was measured by LC-MS/MS. Ten concentrations of compound starting at 10 μM with 3-fold dilutions were examined in duplicate. IC₅₀ values were generated by fitting the data with a four-parameter logistic regression fitting then determining the point that intersects 50% of the difference between negative (time 0) and positive (2 h) uninhibited controls (control IC₅₀).

LPA Mass Spectrometry
Panels of LPA species including C16:0, C18:0, C18:1, and C20:4 were analyzed by using an LC-MS/MS method modified from the published method by Chen et al. (2008). In brief, 50 μl of sample was extracted with 350 μl of acetonitrile/methanol/acetic acid (50:50:1) containing C17:0 LPA at 5 ng/ml as internal standard. Then, 300 μl of supernatant was removed and 700 μl of methanol/water/TEA (66:34:0.1) was added before analysis. Quantitation experiments were performed by two-dimensional LC-MS/MS by using an HP 1200 LC system (Agilent Technologies, Santa Clara, CA) composed of a quaternary pump, a CTC Analytics HTS PAL autosampler (LEAP Technologies, Carrboro, NC), and a switching valve, plumbed in-line with a HP 1100 high-performance liquid chromatography pump that was interfaced to an API 4000 Qtrap mass spectrometer (MDS-Scieix, Toronto, Canada) operated in the negative ion electrospray and multiple-reaction monitoring modes. Typically, 300 μl of sample was injected onto a Betasil C18 column (Thermo Fisher Scientific, Waltham, MA) with 1 ml/min of 25 mM ammonium acetate/methanol (50:50) mixture. After 3 min, the valve was switched, and the captured analytes were eluted onto a 3.0 × 50-mm, 3.5-μm (particle size) XBridge C8 analytical column (Waters, Milford, MA) with a 3-min gradient at a flow rate of 0.8 ml/min. Gradient mobile phases were methanol/methylene chloride/water (55:5:40) with 0.1% TEA and methanol/methylene chloride (50:50) with 0.1% TEA. Selected analytes were specifically detected by monitoring high-performance liquid chromatography retention times and ion pairs corresponding to the parent and specific fragment ion mass-to-charge ratios.

Rat Pharmacokinetics
Male Lewis rats weighing 275 to 300 g were purchased from Charles River Laboratories, Inc. (Wilmington, MA) and acclimated to their surroundings for approximately 1 week with food and water provided ad libitum. A minimum of 1 day before study, animals were anesthetized with isoflurane (to effect) and implanted with Culex (BASi, West Lafayette, IN) vascular catheters in the carotid artery. Animals were acclimated in Culex cages overnight before dosing. Patency of the carotid artery catheter was maintained by using the “tend” function of the Culex automated blood sampler. Animals were dosed with PF-8380 at 1, 3, 10, 30, and 100 mg/kg by oral gavage after an overnight fast. Blood collections were obtained from the carotid artery and performed by the Culex automated blood sampler at 0.25, 0.5, 1, 2, 4, 6, 8, and 24 h after administration. Blood was centrifuged, and plasma was collected for analysis of PF-8380 and LPA concentrations.

Pharmacodynamic Modeling
The LPA data obtained during the rat pharmacokinetic study were fit to the inhibitory Eₘₐₓ pharmacodynamic model described by the following equation:

\[ E = E_{\text{max}} - \left( E_{\text{max}} - E_{\text{p}} \right) \times \frac{C_p}{C_p + \text{EC}_{50}} \] (1)

where the effect (E) is Eₘₐₓ at a plasma concentration of zero (Cₚ = 0) and Eₚ as the plasma concentration approaches infinity (Cₚ = w). EC₅₀ represents the concentration required to achieve 50% of the maximal inhibitory effect. Pharmacodynamic modeling was conducted by using all plasma concentration and LPA concentration data with WinNonlin version 5.2 (Pharsight, Mountain View, CA).

Sample Preparation and LC-MS/MS Analysis of Plasma for PF-8380
Fifty microliters of plasma was mixed with 150 μl of acetonitrile containing LPA 17:0 internal standard and centrifuged to pellet-precipitated proteins. Supernatant from each sample was analyzed on a Sciex API4000 LC-MS/MS system equipped with two Shimadzu (Kyoto, Japan) LC-10AD VP binary pumps and LEAP HTS PAL autosampler. A 5-μl sample extract was injected onto a Thermo Fisher Scientific Aquosil C18 (2.1 × 30 mm, 3 μm) analytical column and eluted by using a linear gradient with mobile phase consisting of 0.1% formic acid and 0.1% formic acid in acetonitrile. PF-8380 was extracted by using all plasma concentration and LPA concentration data with WinNonlin version 5.2 (Pharsight, Mountain View, CA).
analyzed by using Turbo IonSpray (MDS-Sciex) in ESI+ mode with multiple-reaction monitoring of parent/daughter ions at m/z 478.1/161.0.

Carrageenan-Induced LPA in the Rat Air Pouch

Male Lewis rats (175–200 g; Charles River Laboratories) were used in the study. Air pouches were produced by subcutaneous injection of 20 ml of sterile air into the intrascapular area of the back. Pouches were allowed to develop for 1 day. Animals (six per group) were fasted with free access to water for 16 to 24 h before drug administration. Drug, vehicle, or 1 mg/kg dexamethazone were administered by oral gavage 1 h before injection of 2 ml of a 1% suspension of carrageenan (FMC BioPolymer, Philadelphia PA) dissolved in saline into the pouch. At 3 h after carrageenan injection, the pouch fluid was collected by lavage with 2 ml of phosphate-buffered saline containing 10 μM sodium fluoride (Sigma-Aldrich). The fluid was centrifuged at 800g for 10 min at 4°C, and the supernatants were collected for LPA analysis by LC-MS/MS. Blood samples were collected into 10-ml heparinized tubes (Vacutainer tubes; BD Biosciences, Franklin Lakes NJ) 4 h after dosing and immediately centrifuged at 800g for 10 min at 4°C and then frozen at −80°C. For analysis of autotaxin activity, samples of both pouch and plasma were incubated at room temperature for 4 h. LPA was produced from existing LPC and autotxin levels in those fluids. The difference in LPA levels from time 0 to 4 h was used to calculate autotxin activity.

Rat Adjuvant-Induced Arthritis Hyperalgesia

Inflammation was established in female Lewis rats (150–200 g; Harlan, Indianapolis, IN). Five animals per dose group were injected with 3 mg/kg heat-killed Mycobacterium butyricum suspended in paraffin oil subcutaneously on the base of the tail (day 0). By days 10 to 14 rats manifested signs of disease, and vocalization baselines differed by substrate and sources/species of enzyme. The data were fit to an inhibitory model, and the model parameters (EC50, E max, and C20:0 LPA levels in the plasma. The plasma LPA levels indicated a direct response (Supplemental Fig. 1–4). Data show that LPA concentrations declined rapidly and reached their maximal inhibition between 15 and 30 min after administration for all doses levels, corresponding with PF-8380 plasma t max (Table 2). It is interesting to note

| TABLE 1 |
| In vitro potency of PF-8380 |
| Assay | IC50 (nM) |
| Human enzyme FS-3 | 2.8 ± 0.16 (n = 4) |
| Human enzyme LPC | 1.7 ± 0.8 (n = 3) |
| Murine enzyme FS-3 | 1.16 ± 0.07 (n = 3) |
| Fetal fibroblast cell LPC | 1.15 ± 0.18 (n = 3) |
| Human whole blood | 101 ± 0.036 (n = 3) |

PF-8380. Screening of our compound collection identified numerous leads for our medicinal chemistry program. Our medicinal chemistry effort rapidly optimized the lead matter by improving potency and pharmacokinetics properties, resulting in the identification of PF-8380 as a suitable tool compound for in vivo evaluation.

In Vitro Potency. Potency of PF-8380 was confirmed as low nanomolar by using several assay methods. Each method differed by substrate and sources/species of enzyme. The data are summarized in Table 1. Recombinant human enzyme (β isoform) purified from HEK293 cells was used with the high-throughput screening assay of the fluorescent substrate FS-3 and LPC 17:0 as substrates. Both substrates were run at similar enzyme concentrations (approximately 5 nM) and incubation and reaction times. Both substrates gave similar low-nanomolar IC50s. PF-8380 also inhibited rat autotxin with an IC50 of 1.16 nM with FS-3 substrate. Potency of PF-8380 was maintained when using enzyme produced from fetal fibroblasts used in combination with LPC as a substrate. In human whole blood incubated with compound for 2 h autotxin was inhibited with an IC50 of 101 nM.

Pharmacokinetics. The pharmacokinetic profile of PF-8380 was evaluated at an intravenous dose of 1 mg/kg and oral doses of 1 to 100 mg/kg out to 24 h (Fig. 2, A and B). PF-8380 had mean clearance of 31 ml/min/kg, volume of distribution at steady state of 3.2 l/kg, and effective t 1/2 of 1.2 h (Table 2). Oral bioavailability was moderate, ranging from 43 to 83% (Table 2). Plasma concentrations increased with single oral escalating doses (Fig. 2B), but C max increased at a rate that was approximately proportional to dose from 1 to 10 mg/kg and less than proportional to dose from 10 to 100 mg/kg (Fig. 2C). PF-8380 exposures estimated by area under the curve were approximately proportional to dose and linear up to 100 mg/kg (Fig. 2C). Plasma C16:0, C18:0, and C20:0 LPA levels were measured immediately after collection (Fig. 2D shows C18:0 LPA levels at all doses). Maximal reduction of LPA levels was observed by the 3 mg/kg dose at 0.5 h with all LPA returning at or above baseline at 24 h.

PF-8380 Pharmacodynamics. The pharmacodynamic effect of PF-8380 was measured as decreasing C16:0, C18:0, and C20:0 LPA levels in the plasma. The plasma LPA levels and PF-8380 concentrations were fit to an inhibitory E max model, and the model parameters (EC50, E max, and E 0) for each LPA are presented in Table 3. The effect of PF-8380 on plasma LPA levels indicated a direct response (Supplemental Figs. 1–4). Data show that LPA concentrations declined rapidly and reached their maximal inhibition between 15 and 30 min after administration for all doses levels, corresponding with PF-8380 plasma t max (Table 2). It is interesting to note...
Inhibitory $E_{\text{max}}$ model parameters

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Units</th>
<th>C16.0 (95% CI)</th>
<th>C18.0 (95% CI)</th>
<th>C20.0 (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$EC_{50}$ (nmol/l)</td>
<td>54.7 (31.1–78.3)</td>
<td>84.6 (19.7–149)</td>
<td>51.7 (29.1–74.3)</td>
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<tr>
<td>$E_{\text{max}}$ (ng/ml)</td>
<td>19.6 (18.4–20.8)</td>
<td>15.9 (14.2–17.6)</td>
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</tr>
<tr>
<td>$E_{50}$ (ng/ml)</td>
<td>3.25 (2.30–4.21)</td>
<td>3.10 (1.64–4.56)</td>
<td>48.3 (34.8–61.7)</td>
<td></td>
</tr>
</tbody>
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that the $EC_{50}$ Values obtained (means = 51.7–84.6 nM) were similar to the $IC_{50}$ value obtained from human whole blood in the in vitro potency assay (Table 1).

**Discussion**

A high-throughput screening assay for autotaxin inhibitors using FS-3 produced many potent chemical series. A handful of inhibitors of autotaxin have appeared in the literature (Parrill and Baker 2008). A recent report claimed a series of autotaxin inhibitors, inspiring us to engage in a structure–activity relationship aimed at identifying a suitable tool compound (Schiemann et al., 2009). Our medicinal chemistry effort rapidly explored the structure–activity relationship in this lead series with a focus on improved potency and pharmacokinetics properties, resulting in the identification of PF-8380 as a suitable tool compound for in vivo evaluation.
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were much less potent. The series, which included PF-8380, substrate or enzyme produced from a cell line, many series
of formation of LPA 18:1 for 4 h at RT. Human whole blood is the
and 3 h after carrageenan. Autotaxin activity was determined as the rate
of autotaxin activity inhibition in human whole
blood, rat plasma, and rat inflammatory air pouch 4 h after oral dosing
(n = six rats per group). Three hours after carrageenan injection plasma and
pouch fluid were collected, and LPA was analyzed by mass spectrometry.
Plotted are total LPA of four major species immediately after collection from
plasma (A) and air pouch (B) and 4-h incubation at RT for plasma (C) and air
pouch (D).

Fig. 3. In vivo inhibition of LPA generation by PF-8380. Various amounts of
PF-8380 or dexamethasone (Dex) (1 mg/kg) were dosed orally 1 h before injection
of carrageenan in rat air pouch (n = six rats per group). Three hours after carrageenan injection plasma and
pouch fluid were collected, and LPA was analyzed by mass spectrometry.
Plotted are total LPA measured by mass spectrometry from six animals per
group. dex, dexamethasone.

Fig. 4. In vivo inhibition of autotaxin activity by PF-8380. Autotaxin activity
inhibition was measured by incubation of rat plasma (A) and pouch
fluid (B) for 4 h at RT. Plotted is the average total LPA measured by mass
spectrometry from six animals per group. dex, dexamethasone.

When these were tested further using LPC either as a substrate or enzyme produced from a cell line, many series
were much less potent. The series, which included PF-8380,
maintained their potency both when using LPC or FS-3 as a
substrate and either recombinant or enzyme secreted by fetal
fibroblast cells. Subsequent analogs synthesized within this
series produced PF-8380, with nanomolar potency in enzyme
assays and 100 nM potency in human whole blood. Several
observations suggest that PF-8380 is a tight-binding inhibi-
tor of autotaxin with subnanomolar potency. PF-8380 seems
to be competitive with substrate (data not shown). Tradition-
ally kinetic analysis was complicated by apparent sub-
strate inhibition in the FS-3 substrate assay with a

300
350
400
450

Fig. 5. Correlation of autotaxin activity inhibition in human whole blood
(HWB), rat plasma, and rat air pouch. Percentage of inhibition of auto-
taxin activity is plotted versus concentration of PF-8380 for human whole
blood, rat plasma, and rat inflammatory air pouch 4 h after oral dosing
and 3 h after carrageenan. Autotaxin activity was determined as the rate
of formation of LPA 18:1 for 4 h at RT. Human whole blood is the average ± S.E.M. for three donors. In vivo plasma and pouch is the percentage of inhibition of an average of six animals per group.

When these were tested further using LPC either as a
substrate or enzyme produced from a cell line, many series
were much less potent. The series, which included PF-8380,
Inhibition of hyperalgesia associated with adjuvant-induced arthritis by PF-8380. Female Lewis rats (five animals per dose group) were treated orally b.i.d. with 10, 30, or 100 mg/kg PF-8380 or 30 mg/kg naproxen (positive control), and vehicle and vocalization were assessed on day 3, 2 h after administration. Plotted is the average percentage of inhibition of vehicle versus control ± S.E.M.

$K_i$ of approximately 0.04 nM and an enzyme concentration of 2.2 nM (data not shown). In addition, a direct fit of enzyme activity versus inhibitor concentration data using the Morrison equation for tight-binding inhibitors yielded a $K_i$ of 0.02 nM and an enzyme concentration of 2.6 nM (data not shown). These observations indicate that under the conditions of the recombinant enzyme assays PF-8380 is essentially titrating the enzyme active site.

Because most preclinical animal models are done in rodents, the potency of PF-8380 was evaluated against rat autotaxin. Rat and murine autotaxin sequences are identical, so these measurements can be used for either species. The potency in the recombinant enzyme assay was comparable with human. A determination was made that human whole-blood assays could be used to estimate potency in a rodent system. Human whole-blood potency was shifted as expected with human. A determination was made that human whole-blood assays could be used to estimate potency in a rodent system. Human whole-blood potency was shifted as expected because the LPA biomarkers in this study were peripheral caused by adjuvant-induced arthritis is primarily peripheral, because the LPA biomarkers in this study were peripheral measures of LPA reduction.

Analysis of a pharmacokinetic/pharmacodynamic (PK/PD) relationship was attempted by measuring plasma LPA. Our initial observations with human whole blood and the observation in autotaxin heterozygote knockout animals indicate that autotaxin is the major source of LPA in plasma. To determine a PK/PD effect, however, the rate of biosynthesis/degradation of LPA needs to be taken into account. Whole-blood assays depend on an incubation period from several hours up to 24 h to produce LPA from plasma autotaxin. Our analysis was careful to limit the time of incubation of the plasma after sampling, but with many time points it was difficult to obtain samples at precisely the right time, meaning that there was most certainly some incubation time. In any event, LPA levels were rapidly reduced after oral dosing, indicating that LPA present at basal levels in the plasma is rapidly degraded.

Analysis of the in vivo effects of PF-8380 on production of LPA in plasma and the inflamed air pouch indicates a close correlation between plasma, pouch, and ex vivo whole-blood inhibition of autotaxin activity. Plasma and pouch samples were collected immediately and processed for LPA analysis. Plasma samples without incubation to determine autotaxin activity did not show a dose-dependent decrease in LPA as was seen in the PK study. Plasma that was incubated for 4 h at RT showed a very good dose response. This indicated that basal LPA levels are not as sensitive to autotaxin inhibition and that analysis of autotaxin activity needs to be performed to determine a PK/PD relationship. Pouch samples, on the other hand, either with 4 h of incubation or no incubation, showed a similar dose response. Presumably the pouch, with a robust inflammatory condition, produces much more autotaxin, hence higher basal levels of LPA, and it is sensitive to autotaxin inhibition.

To estimate doses necessary to inhibit LPA formation in vivo at the site of inflammation, a correlation of human whole-blood inhibition of plasma autotaxin activity and air pouch autotaxin activity indicate that there is a close correlation of the three. This suggests that a whole-blood assay can be an accurate measure of subsequent doses needed to completely inhibit the production of LPA at the site of inflammation. Autotaxin’s role in the production of LPA at the site of inflammation was then tested with PF-8380. Immediately after samples were taken from the pouch LPA levels were analyzed, which minimized LPA formation by autotaxin activity in the fluid. LPA levels in the pouch were inhibited >90%, indicating that autotaxin is the primary source of LPA at the site of inflammation. PF-8380 was evaluated for its inhibition of inflammatory hyperalgesia in rat adjuvant-induced arthritis. Maximal efficacy in hyperalgesia was equal to naproxen at 30 mg/kg. In previous studies, naproxen dosed orally at 10 mg/kg was determined from biomarker data to be efficacious to inhibit PGE2 and thus both forms of cyclooxygenase. A dose of naproxen three times higher was used in this study, which represents maximal effect based on cyclooxygenase. Naproxen at 100 mg/kg produces side effects in rats, such as a gastrointestinal side effect, that will not provide mechanism-based data; therefore, we did not test doses higher than 30 mg/kg. This indicated that hyperalgesia caused by adjuvant-induced arthritis is primarily peripheral, because the LPA biomarkers in this study were peripheral measures of LPA reduction.

In summary, this is the first report of an inhibitor of autotaxin that reduces LPA levels in vivo. PF-8380 has been demonstrated to be a potent inhibitor of autotaxin in vitro by using native enzyme and substrate and also in ex vivo human whole blood. PF-8380 is bioavailable and can be dosed in vivo to levels that are several multiples above the $ED_{50}$, which can be accomplished with twice a day dosing at 10 mg/kg. Autotaxin’s role in the production of LPA at the site of inflammation was confirmed in the rat air pouch model of inflammation. Autotaxin was inhibited >90% both in plasma and at the site of inflammation by PF-8380, indicating that autotaxin is the primary enzyme responsible for the production of LPA at the site of inflammation. Measurement of autotaxin activity in ex vivo whole blood or plasma can be used to calculate dose projections, and PF-8380 can now be used as a pharmacological tool to evaluate the role that LPA plays in inflammation and cancer.

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References


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A Novel Autotaxin Inhibitor Reduces Lysophosphatidic Acid Levels in Plasma and the Site of Inflammation

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**Figure S1. Plasma PF-8380 and LPA concentrations following a 30 mg/kg oral dose in rats.** PF-8380 concentrations are represented by black closed symbols and solid lines and LPA concentrations are represented by gray open symbols and dashed lines. PF-8380 and LPA concentrations are plotted as a function of time.

A.

B.

C.
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**Figure S2. C16:0 LPA concentrations as a function of plasma PF-8380 concentration for doses ranging from 0-100 mg/kg.** Data points on the graphs are the corresponding blood collection time points.
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Figure S3. C18:0 LPA concentrations as a function of plasma PF-8380 concentration for doses ranging from 0-100 mg/kg. Data points on the graphs are the corresponding blood collection time points.
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Figure S4. C20:0 LPA concentrations as a function of plasma PF-8380 concentration for doses ranging from 0-100 mg/kg. Data points on the graphs are the corresponding blood collection time points.