Innate immune responses are essential to host defense, yet if unchecked can lead to tissue injury and illness (1). PMN are the primary initial immune effectors of acute inflammation, and these cells use as many as fifty toxins for microbial killing in phagocytic vacuoles (2). Incomplete closure of phagolysosomes or aberrant extracellular release of reactive oxygen species (ROS), granule components, lipid mediators, hypochlorous acid, and other potentially toxic PMN products to surrounding tissues contribute to injury in several human diseases, including the devastating clinical entities of acute lung injury (ALI) and acute respiratory distress syndrome (ARDS) for which no disease-remitting therapy is currently available (3).

To prevent an overexuberant inflammatory response and limit damage to the host, PMN activation programs need to be tightly controlled (1). Select membrane-derived lipid mediators have recently been described as autacoid regulators of PMN functional responses in acute inflammation (4). One of these classes of antiinflammatory membrane lipids involves isoprenoid metabolism. Immune regulatory roles for isoprenoids are emphasized by the hyper-IgD and periodic fever syndrome that results from low polyisoprenyl phosphate (PIPP) levels secondary to defective mevalonate kinase activity (5). Recently, we identified a novel PIPP signaling pathway in PMN (6). One of its components is presqualene diphosphate (PSDP), present in freshly isolated human PMN in nanomolar quantities at baseline (i.e., 1.7 nmol/10⁷ cells) (6). PMN exposure to the chemoattractant and secretagogue leukotriene B₄ (LTB₄) initiates transient activation of this PIPP signaling pathway with conversion of PSDP to its monophosphate form, presqualene monophosphate (PSMP). PSDP carries biological...
activity as a potent counterregulatory mediator that prevents ROS generation (6, 7). In sharp contrast, PSMP is >100-fold less active than PSDP for inhibition (6, 7). Thus, incoming positive signals for PMN (e.g., LTB4) initiate rapid degradation and inactivation of an inhibitory lipid signal (i.e., PSDP) coincident with cell responses (e.g., ROS generation). PSDP levels quickly return to baseline in a time frame that parallels cellular deactivation. Intracellular targets for PSDP to control PMN activity remain to be elucidated. Select PSDP structural mimetics are also active in vivo, dampening mouse responses to zymosan A–induced peritonitis (8).

In addition to PSDP remodeling, LTB4 also initiates phosphatidylinositol 3–kinase (PI3K) activation in PMN to promote NADPH oxidase assembly and ROS production (9, 10). Phosphoinositide signaling initiated by PI3K is a critical early event in PMN responses, such as phagocytosis (11) and chemotaxis (12), and contributes to ALI pathogenesis (13). Because LTB4 initiates PMN PI3K activation and PIPP remodeling, we hypothesized that these signaling events were related in the regulation of PMN responses. Here, we report that PI3K activity and PSDP remodeling are linked during PMN activation and deactivation with direct inhibition of PI3K by PSDP to limit PMN responses and lessen the severity of experimental lung inflammation.

RESULTS AND DISCUSSION
PSDP and PI3K regulate LTB4–triggered O2− release by human PMN
To determine if PSDP and PI3K regulate LTB4-stimulated responses, we exposed freshly isolated human PMN to a new structural PSDP mimetic (Fig. 1 A) or a potent and specific inhibitor of PI3K activity (LY294002) before LTB4. A marked increase in the rate of O2− generation was observed within seconds after LTB4 addition that was transient, slowing considerably by 60 s (Fig. 1 B). The presence of either PSDP mimetic (100 nM) or PI3K inhibitor (3 μM) markedly blocked O2− generation. Exposure to LTB4 (10 min) induced 1.6 ± 0.3 nmol O2−/106 PMN (as compared with vehicle 0.5 ± 0.1 nmol O2−/106 PMN; P < 0.01). Both the PSDP mimetic and PI3K inhibitor led to >50% inhibition of LTB4–triggered O2− generation (0.8 ± 0.3 nmol/106 PMN and 0.7 ± 0.2 nmol/106 PMN, respectively; P < 0.05) (Fig. 1 C). The PSDP mimetic concentration in these experiments (i.e., 100 nM) was 1,000-fold lower than its critical micellar concentration (CMC) (~100 μM; Fig. S1, at http://www.jem.org/cgi/content/full/jem.20052143/DC1). To verify that the observed inhibition was not secondary to micelle formation and sequestration of the lipid agonist LTB4, in the next experiments, there was pretreatment with the mimetic, a wash, and stimulation with LTB4. Although this extra step reduced total ROS generation in response to LTB4, the PSDP mimetic still led to a >50% decrease (71.2 ± 27.7% inhibition, n = 4). Together, these results indicate that human PMN activation by LTB4 is highly dependent on PI3K activity and can be inhibited by a new PSDP structural mimic.
Materials and methods, available at http://www.jem.org/cgi/content/full/jem.20052143/DC1) that progressed to 27.1 ± 4.8% reductions (P = 0.001) within 30 s (Fig. 2 B).

These results indicate that LTB₄-triggered activation of PMN PI3K and PSDP remodeling was concurrent and temporally overlapped with initiation of ROS generation. Although both of these signaling events were rapid in onset, the kinetics for PI3K activation and deactivation differed from the time course for total PSDP remodeling during the initial 30-s interval. To determine if PSDP directly interacted with PI3K as its activity decreased, PMN were exposed to LTB₄ for 30 s and p110γ–PI3K was immunoprecipitated from cellular materials. Lipid extracts were performed and analyzed by gas chromatography/mass spectrometry (GC/MS) (6). Selective ion monitoring at m/z 137 (2 isoprenoid units) revealed a unique peak at 18.1 min in LTB₄-exposed PMN extracts. MS spectral diagnostic ions (Fig. 2 C), namely m/z 567 [M⁺ − (H₂O)], 488 [M⁺ − (H₃PO₄)], 410 [M⁺ − (HP₂O₇)], 341 [M⁺ − (HP₂O₄) − 69], 205 [M⁺ − (HP₂O₅) − 69 − (CH₃C(CH₃)CHCH₂ × 2)], 137 [C₆H₇], 97 [HP₂O₇], 81 [CH₂C(CH₃)CH(CH₂)₂ − H⁺] and 69 [base peak; (CH₃)₂CCHCH₂], were consistent with authentic PSDP in the PI3K immunoprecipitated material.

Direct inhibition of p110γ–PI3K by PSDP

Because PI3K activity and PSDP remodeling were both early signaling events in PMN with interaction between PSDP and p110γ–PI3K, next we questioned if PSDP could directly regulate PI3K activity. Recombinant human (rh) p110γ–PI3K activity was determined by PIP₃ formation in vitro in the presence of PSDP, PSMP, or the PI3K inhibitor LY294002. PIP₃ formation was significantly decreased by PSDP (800 pmol) with 94.7 ± 5.3% inhibition (P < 0.001) and a PI3K inhibitor (500 pmol) with 46.7 ± 6.7% inhibition (P = 0.01) (Fig. 3 A). PSDP inhibited p110γ–PI3K in a concentration-dependent fashion (Fig. 3 B). In sharp contrast, PSMP (8–800 pmol) did not significantly impact p110γ–PI3K activity. The IC₅₀ for PSDP (38 pmol) had a stoichiometry with PI3K of 9:1. These results indicate that PSDP is a potent direct inhibitor of p110γ–PI3K with a structure–activity relationship that suggests an important role for the diphosphate structure in PSDP’s action on p110γ–PI3K activity. The LTB₄-mediated PMN remodeling of PSDP corresponds to an ~50 pmol change in PSDP/10⁶ PMN, a decrease that is within the concentration range for regulation of p110γ–PI3K activity (Fig. 3 B). After cell activation, the percent change in total PSDP (i.e., 28%) is similar to the change in phosphatidylinositol (17%) that occurs in activated PMN membranes (14). Collectively, our new findings indicate that receptor-mediated agonists for PMN remodel PSDP in time-frames and amounts consistent with functional impact on PI3K activity and cellular responses.

PSDP remodeling in vivo during tissue injury and inflammation

Because PI3K activity occupies a central role in regulating PMN activation during lung injury and inflammation (13), next we determined PSDP remodeling in vivo in mouse lungs during an experimental model of mild ALI secondary
to aspiration of gastric acid (15), which is a common clinical event (3). To simulate acid aspiration, hydrochloric acid (HCl) (0.1 N, pH 1.5) was selectively instilled into the animals’ left lungs (15). Lung PMN infiltration was maximal 12 h after HCl injury (14.2 ± 1.8 vs. 5.5 ± 0.8 × 10⁴ PMN/mg lung; P < 0.01) (Fig. 4 A). Expression of class IA and IB PI3Ks in mouse lungs were both increased at 2 and 12 h after HCl (Fig. 4 B). Lungs were removed and lipid extracts were prepared for PSDP determination. Of interest, despite increased PMN numbers, PSDP levels were significantly lower in the left lungs of HCl-injured mice (4.6 ± 0.3 µg PSDP vs. 9.0 ± 1.6 µg PSDP in control lungs; P < 0.02). These results indicate that experimental lung injury led to decrements in PSDP concomitant with increased PMN, suggesting an inverse relationship in vivo between lung PSDP and inflammation.

PSDP mimetic blocks PMN infiltration and PI3K activity

To determine if PSDP can block pulmonary inflammation and PI3K in vivo, we administered a PSDP structural mimetic (0.8 µg/mouse, i.v.) or vehicle 15 min before HCl instillation into the left main-stem bronchus. PSDP markedly reduced lung PMN 12 h after injury (Fig. 5 A). Tissue morphometry on LY-6G–stained histological sections (for identification of mouse PMN) revealed significant inhibition with the PSDP mimetic (4.6 ± 0.3 µg PSDP vs. 9.0 ± 1.6 µg PSDP in control lungs; P < 0.02). These results indicate that experimental lung injury led to decrements in PSDP concomitant with increased PMN, suggesting an inverse relationship in vivo between lung PSDP and inflammation.

Figure 3. PSDP directly blocked PI3K activity. (A) rhp110γ–PI3K activity was determined in the presence or absence of PSDP (800 pmol), PSMP (800 pmol), or a PI3K inhibitor (500 pmol). (B) Concentration response for the impact of PSDP (●) or PSMP (■) on rhp110γ–PI3K activity (mean ± SEM; n = 3; *, P < 0.05 as compared with vehicle).

Figure 4. ALI induced PMN infiltration, PI3K, and PSDP remodeling. (A) 12 h after HCl-initiated ALI, PMN accumulation was determined by MPO activity in lung homogenates (n = 5; *, P = 0.002). (B) Semi-quantitative RT-PCR for p85 and p110γ–PI3K mRNA expression in the injured lungs 2 and 12 h after HCl instillation (n = 3; *, P < 0.05). (C) PSDP levels were measured in lung lipid extracts with or without HCl-initiated injury (12 h; mean ± SEM; n = 3–4; *, P < 0.05).

that increased markedly after ALI (Fig. 4 B), we next determined class IA PI3K activity after acid injury in the presence or absence of the PSDP mimetic. HCl injury induced significant increases in lung PI3K activity in p85 immunoprecipitates (0.59 ± 0.17 PIP₃/mg lung with HCl vs. 0.05 ± 0.02 pmol PIP₃/mg lung with PBS; P < 0.05). Administration of the PSDP mimetic blocked the HCl–induced increase in class IA PI3K activity to near basal levels (0.05 ± 0.01 pmol PIP₃/mg lung; P < 0.05) (Fig. 5 C). Thus, PSDP can regulate PMN activation, tissue accumulation, and total PI3K activity in vivo during experimental acid–initiated ALI.

During acute inflammation, PI3Ks orchestrate several cellular responses for host defense, including PMN ROS generation (12). Benefiting its central role in cell activation, several mechanisms are in place to restrain PI3K activity (16–18). Previous reports have suggested a link between decreased PIPP formation and increased PI3K (19, 20). Results presented here are the first to demonstrate direct inhibition of PI3K by a PIPP and inverse relationships between PI3K
activity and PSDP levels both in vitro and in vivo. In addition, PSDP bound to p110γ–PI3K in activated PMN and potently inhibited rhp110γ–PI3K in vitro and a PSDP structural mimetic blocked PI3K activity in vivo. Together, these new findings support a signaling relationship between PI3K and PIPPs in the regulation of leukocyte functions during inflammation.

Pivotal regulatory properties have been ascribed to isoprenoids. For example, polyisoprenyl glycolipids form antigen complexes with CD1 to activate T cells (21), and cholesterol is critical to PMN cell membrane organization and polarization in response to chemotactic stimuli (22). Although PIPPs are appreciated as cholesterol biosynthetic intermediates, PSDP is also present in cells, such as human PMN, that cannot use it for cholesterol biosynthesis because they lack squalene cyclase and other mixed function oxidase activities (23). There are now several lines of evidence to support a role for PSDP as a counterregulatory signal in PMN functional responses. PSDP is also present in cells, such as human PMN, that cannot use it for cholesterol biosynthesis because they lack squalene cyclase and other mixed function oxidase activities (23). There are now several lines of evidence to support a role for PSDP as a counterregulatory signal in PMN functional responses. PSDP is a constituent of PMN cell membranes (20), and receptor activation leads to rapid and transient PIPP remodeling. PSDP is a potent and direct inhibitor of phospholipase D (PLD) that is activated, in part, via isoprenylation (25). These shared isoprenoid moieties with PSDP may provide additional sites of interaction outside the PI3K catalytic center. Of interest, PSDP and PI3K are present in resting PMN in complementary intracellular locations (cell membranes and cytosol, respectively), and both translocate to intracellular organelles upon cell activation (20, 26).

Activated PMN contribute to the development and severity of tissue injury during inflammatory illnesses such as ALI. No specific therapy is currently available to modulate the inflammatory response to airway injury from acid and protect the lung in ALI. PI3K is an important enzyme in the proinflammatory PMN signaling program of ALI, as p110γ–PI3K deficiency dampens the severity of endotoxin-induced ALI (13). In addition to PMN, PI3Ks are also activated during ALI in lung-resident cells (27) and both endothelial cell and PMN PI3Ks contribute to PMN accumulation in the lung (28). Select lipid mediators, namely lipoxin A₄ (LXA₄), inhibit PI3K in structural cells (29). LXA₄ signaling also promotes resolution of experimental ALI (15) and inhibits PMN activation by LTB₄ in a postreceptor manner, in part by blocking PSDP remodeling (7). Here, PSDP levels were

Figure 5. PSDP reduced acid-initiated ALI. (A) PSDP structural mimetic (0.8 μg/mouse) or vehicle was administered i.v. 15 min before HCl injury. 12 h after HCl instillation, histological specimens were prepared and mouse PMN were identified by LY-6G immunostaining (arrows). Bar, 100 μm. (B) Tissue morphometry was performed to determine the percentage of LY-6G staining cells in mouse lungs (n = 4 measurements in each group; *, P < 0.05 as compared with HCl-injured left lung). (C) Class IA (p85-based) PI3K activity was determined in lung lysates after ALI. Data are mean ± SEM. n = 3; *, P < 0.05 as compared with control; **, P < 0.05 as compared with HCl-injured lung.
decreased in acid-injured lungs and a novel PSDP mimetic blocked PI3K, PMN ROS generation, and PMN accumulation in the lung. Because the PSDP mimetic was administered intravenously, regulation of cells other than PMN may have also contributed to the marked inhibition of leukocyte trafficking after acid injury.

In conclusion, the ability of PSDP and a new PSDP mimetic to directly inhibit PMN early intracellular activating signals, such as PI3K, and to lessen the inflammation associated with experimental ALI provides insight into new mechanisms for in vivo protection from excess PMN-driven inflammation and tissue injury. Together, our findings suggest that PIPP signaling pathways, and specifically PSDP, can serve as natural templates for the design of new therapeutic strategies in inflammatory diseases.

MATERIALS AND METHODS

Materials. PSDP and PSMP were isolated from human PMN or prepared by total organic synthesis (8). The bisphosphonate PSDP structural mimetic, tetraethyl presqualene carboxamido-methylene-diphosphonate was prepared from presqualene carboxylic acid. All synthetic compounds were characterized by NMR spectroscopy.

Human PMN incubations. Peripheral blood was obtained by venipuncture from healthy volunteers who denied taking any medications for at least 2 wk and had given written informed consent to a protocol approved by Brigham and Women’s Hospital’s Human Research Committee. PMN were isolated from whole blood as described previously (6). Freshly isolated PMN (1–5 × 10^6 PMN/ml HBSS plus 1.6 mM CaCl_2) were incubated (5 min, 37°C) in the presence of 3 μM LY294002, 100 nM PSDP mimetic, or vehicle (0.1% ethanol), then exposed to LTβ (100 nM) in the presence of 7 mg/ml cytochrome c. This concentration of LTβ was chosen because it initiates PMN NADPH oxidase assembly (7) and is similar to amounts measured in vivo at sites of acute inflammation (30). In some incubations, PMN in HBSS without calcium were exposed to PSDP mimetic or vehicle, pelleted by centrifugation (700 g, 3 min), and resuspended in HBSS plus 1.6 mM CaCl_2 without PSDP before the addition of agonist. Superoxide anion generation was determined (37°C) as superoxide dismutase-inhibitable cytochrome c reduction by monitoring (550 nm) at 5-s intervals in a continuously flowing water-bath–jacketed cuvette or after timed incubations. For PSDP identification, PMN (50–100 × 10^6 cells/ml) were activated (LTβ, 100 nM, 30 s) before disruption by N_2 cavitation (350 psi, 20 min, 4°C). Remaining intact cells and nuclei were pelleted (500 g, 10 min, 4°C) and supernatants were used for immunoprecipitation with anti-p110γ (15 min, 37°C). After overnight incubation, washed protein A agarose (60 μl) was added and incubated for 2 h at 4°C. After several washes, the immunoprecipitates were saponified, lipids were extracted, and materials were analyzed C. After several washes, the immunoprecipitates were saponified, lipids were extracted, and materials were analyzed. After several washes, the immunoprecipitates were saponified, lipids were extracted, and materials were analyzed. After several washes, the immunoprecipitates were saponified, lipids were extracted, and materials were analyzed.

Measurement of class IA and IB PI3K activity. After immunoprecipitation with either an anti-p85 or anti-p110γ-PI3K selective antibody, PI3K activity from PMN and lung homogenates was determined by PIP_2 formation in vitro using a competitive ELISA (Echelon Biosciences Inc.). For recombinant purified PI3K enzyme, activity was determined by a luminescent kinase assay (Kinase-Glo; Promega) that quantitated decrements in ATP by kinase utilization. rh p110γ–PI3K (4 pmol/reaction) was exposed (15 min, 37°C) to PSDP or PSMP (0.4–800 pmol) followed by 25 μg t-α-phosphatidylinositol and 1 μM ATP and allowed to react for 90 min at room temperature. Incubations were stopped with 50 μl of Kinase-Glo reagent and incubated an additional 10 min at room temperature.

Luminescence was measured with a FLx800 microplate luminometer (BioTek Instruments, Inc.).

Experimental model of ALI. All animal protocols were approved by the Harvard Medical Area Animal Institutional Review Board. Acid (0.1 N HCl, pH = 1.5, 50 μl) was instilled intratracheally into the left lung of anesthetized mice (FVB, male, 10–12 wk; Charles River Laboratories) (15). A PSDP mimetic (0.8 μg in 100 μl 0.9% saline) or vehicle (1% ethanol) was administered by tail vein 15 min before HCl instillation. After 12 h, lungs were removed, prepared for MPO (15) or PI3K assay, or were fixed in IHC zinc buffer and paraffin embedded for immunostaining with LY-6G (1:50 dilution). Area and number of positively staining cells was measured with National Institutes of Health Image software and percentage of positive cells/area calculated.

Statistical analysis. Results are expressed as the mean ± SEM. Statistical significance of differences was assessed by Student’s t test and one-way analysis of variance. P < 0.05 was set as the level of significance.

Online supplemental material. Fig. S1 shows the CMC determination for the PSDP mimetic and related compounds. Further information on materials and experimental protocols are supplied as the supplemental Materials and methods. Online supplemental material is available at http://www.jem.org/cgi/content/full/jem.20052143/DC1.

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