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P2Y6 signaling in alveolar macrophages prevents leukotriene-dependent type 2 allergic lung inflammation

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Conflict of interest:

Antagonists of the type 1 cysteinyl leukotriene receptor (CysLT1R) are widely used to treat asthma and allergic rhinitis, with variable response rates. Alveolar macrophages express UDP-specific P2Y6 receptors that can be blocked by off-target effects of CysLT1R antagonists. Sensitizing intranasal doses of an extract from the house dust mite Dermatophagoides farinae (Df) sharply increased the levels of UDP detected in bronchoalveolar lavage fluid of mice. Conditional deletion of P2Y6 receptors before sensitization exacerbated eosinophilic lung inflammation and type 2 cytokine production in response to subsequent Df challenge. P2Y6 receptor signaling was necessary for dectin-2-dependent production of protective IL-12p40 and Th1 chemokines by alveolar macrophages, leading to activation of NK cells to generate IFN-γ. Administration of CysLT1R antagonists during sensitization blocked UDP-elicited potentiation of IL-12p40 production by macrophages in vitro, suppressed the Df-induced production of IL-12p40 and IFN-γ in vivo, and suppressed type 2 inflammation only in P2Y6-deficient mice. Thus, P2Y6 receptor signaling drives an innate macrophage/IL-12/NK cell/IFN-γ axis that prevents inappropriate allergic type 2 immune responses on respiratory allergen exposure and counteracts the Th2 priming effect of CysLT1R signaling at sensitization. Targeting P2Y6 signaling might prove to be a potential additional treatment strategy for allergy.

Introduction

Type 2 immunity (T2I) mediates host defense against helminthic parasites (1) and facilitates epithelial barrier repair (2). If dysregulated or directed against innocuous antigens (allergens), T2I can also promote persistent eosinophilic inflammation and disease. Allergens in house dust mites (HDMs), fungi, and pollen carry endogenous adjuvants (endotoxin, lipids, proteases, and glycans) that activate barrier cells and antigen-presenting cells through pattern recognition receptors (3, 4). This innate response bridges to adaptive T2I involving effector and memory T cells, antigen-specific IgE, and the production of the type 2 cytokines IL-4, IL-5, and IL-13 (5). T2I to allergens is strongly associated with increasingly prevalent diseases such as asthma, rhinosinusitis, and allergic esophagitis, each of which involves end-organ dysfunction and remodeling due to T2I-dependent immunopathology (6–8). Recent studies using cytokine-targeted therapies and anti-IgE validate the importance of the pathologic processes regulated by T2I in these diseases (9, 10). Checking the destructive potential of T2I requires the presence of mechanisms that ensure that the initial innate response to allergens is tightly regulated and compartmentalized. The basis of this regulation remains incompletely understood but likely involves components of the innate immune response that control the magnitude of subsequent T2I responses.

In addition to exogenous adjuvants, endogenous danger-associated molecular patterns (DAMPs) (lipids, cytokines, and other extracellular signaling molecules) help to bridge innate to adaptive immunity. DAMPs derive from tissue-resident cells or recruited myeloid cells early following exposure to antigens (3, 11). They fine-tune signal strength and production of canonical cytokines by antigen-presenting cells, thus shaping instructions received by lymphoid effectors. Cysteinyl leukotrienes (cysLTs) are arachidonic acid–derived mediators generated by mast cells, eosinophils, DCs, and macrophages. In addition to their role as bronchoconstrictors in asthma (12, 13), they act as DAMPs that facilitate T2I through effects on DCs (3) and group 2 innate lymphoid cells (ILC2s) (14). These effects are mediated by the type 1 receptor for the cysLTs (CysLT1R), a GPCR that is the target for the widely prescribed drugs montelukast, zafirlukast, and pranlukast. The therapeutic efficacy of these drugs for the treatment of asthma and allergic rhinitis varies, with as few as 24% to as many as 78% of patients failing to show symptomatic improvement (15, 16). The reasons for this variable response are not clear but may involve individual variation in the contributions from cysLTs to T2I (17), as well as potential off-target effects of the drugs at receptors that exert a protective role (18).

The purinergic receptor (P2Y) class of GPCRs is a potential unintended target of CysLT1R antagonists. P2Y receptors recog-
nize extracellular nucleotides (19), are structurally homologous to CysLT1R (20), and are blocked in vitro by CysLT1R antagonists (21). P2Y receptors, along with nucleotide-selective ion channels (P2X receptors), mediate DAMP-like effects of nucleotides in T21. ATP is released at high levels in the airways of asthmatic subjects challenged with allergen (22). ATP primes lung DCs to promote T21 to the innocuous experimental antigen ovalbumin (22) and acts at P2Y receptors on respiratory epithelial cells to induce the release of IL-33, a T21-promoting alarmin (4). Less is understood about the roles of UTP and UDP (23), the latter of which preferentially binds P2Y1 receptors (24). P2Y1 receptors are expressed by bronchial epithelial cells during pathologic mucosal inflammation (25, 26) and are also expressed strongly and constitutively by monocytes, macrophages, and DCs (27) (http://rstats.immgen.org/Skyline/skyline.html). Because P2Y1 receptors exhibit the highest affinity (nM range) (28) of all P2Y receptors for their cognate ligand, these cells may be highly sensitive to changes in the local levels of UDP. P2Y1 receptor signaling facilitates proinflammatory chemokine and cytokine production through both autocrine and paracrine circuits in several cell types ex vivo (26, 29, 30), but its role in T21 is unclear. Selective P2Y1 receptor blockade decreases allergen-induced lung eosinophil counts and type 2 cytokine production in immunologically sensitized mice (26). In contrast, induced deletion of P2Y1 receptors markedly enhances pulmonary T21 responses of mice to repetitive inhalation challenges with an extract (Df) of the clinically important HDM Dermatophagoides farinae (31), and a selective P2Y1 receptor agonist prevented the development of airway hyperresponsiveness and inflammation in a mouse asthma model (32). These studies suggest a complex, context-specific function of UDP and P2Y1 receptors that may either protect from or promote T21 in the lung.

In this study, we sought to identify key steps and cell types responsible for the protective effect of P2Y1 receptors and their preferred ligand, UDP, in pulmonary T21; to resolve the potentially contradictory effects of P2Y1 receptor signaling in different models of allergen-induced pulmonary inflammation; and to determine the potential consequences of P2Y1 receptor blockade by CysLT1R antagonists in vivo. We uncover a major role of UDP/P2Y1 receptor signaling in controlling a previously unrecognized protective function of alveolar macrophages (AMs). This function was mediated by IL-12–dependent activation of an innate, NK cell–dependent IFN response that dampened subsequent Df–induced T21. We also demonstrate that the protective P2Y1 response is blocked by antagonists of CysLT1R in vivo and counterbalances the established pro-T21 role of CysLT1R signaling at sensitization (33).

**Results**

**Deletion of P2Y1 at sensitization, but not at challenge, amplifies pulmonary type 2 responses to Df.** P2ry6fl/fl/Cre/+ mice, which were generated from P2ry6fl/fl and Rosa26-CreER<sup>22</sup> mice, display increased T21 and associated eosinophilic pulmonary inflammation compared with P2ry6fl/fl/+ (+/+ ) controls when subjected to repetitive administration of low-dose (3 μg) Df over a 3-week period (31). To distinguish the potential contributions of UDP and P2Y1 receptors to the respective sensitization and effector steps of the Df–induced T21, we modified the protocol to incorporate distinct sensitization and challenge phases (Figure 1A). Both +/+ and P2ry6fl/fl/Cre/+ mice were treated with tamoxifen daily i.p. for 5 days, beginning 14 days before the first Df exposure, to delete the P2ry6 allele prior to sensitization. Mice received sensitizing i.n. doses (20 μg) of Df on successive days (days 0 and 1), followed by challenges with low-dose Df (3 μg) on days 14 and 15, or they received equal volumes of NaCl. BAL fluid and tissues were collected 16 hours after the last challenge dose. Tamoxifen induced 80%–90% deletion of the P2ry6 transcript in the lung, spleen, BM, pararachetal lymph nodes, and BAL fluid cells of the P2ry6fl/fl/Cre/+ mice (Supplemental Figure 1A; supplemental material available online with this article; https://doi.org/10.1172/JCI129761DS1). Quantitative PCR (qPCR) of AMs (CD11c<sup>-</sup> BAL fluid cells) sorted from P2ry6fl/fl/Cre/+ mice demonstrated an 84% decrease in expression of the floxed P2Y1 (exon 3) allele, indicating that more than 80% recombination had occurred (Supplemental Figure 1B). BM-derived macrophages (BMMs) from +/+ mice showed marked dose-dependent increases in intracellular calcium levels in response to stimulation with UDP that were attenuated in macrophages derived from the BMs of P2ry6fl/fl/Cre/+ mice (Supplemental Figure 1C). Both BMMs and DCs and from +/+ mice displayed rapid decreases (3 hours) in P2ry6 mRNA expression in response to stimulation with Df (Supplemental Figure 1D).

To determine whether the deletion of P2Y1 receptors prior to sensitization enhanced Df–induced T21 in the modified protocol, we examined BAL fluid for cell counts and mediators on day 16 (16 hours after the second challenge dose of Df) and for type 2 cytokines. Whereas Df–sensitized and -challenged +/+ mice displayed minimal to negligible BAL fluid eosinophilia and neutrophilia, P2ry6fl/fl/Cre/+ mice showed significant increases in BAL fluid total cell, eosinophil, neutrophil, and AM/lymphocyte counts (Figure 1B). The level of BAL fluid eosinophilia observed in +/+ mice in this Df protocol (2 sensitizing doses followed by 2 challenges 14 days later) was lower than that induced by the repetitive Df protocol used in the previous study (6 i.n. doses over 18 days) (31). H&E-stained lung sections revealed negligible bronchovascular cellular infiltrates in Df–treated +/+ controls but substantial infiltrates in P2ry6fl/fl/Cre/+ mice (Figure 1C). Levels of IL-4, IL-5, IL-13, eosinophil peroxidase (EPO), and myeloperoxidase (MPO) were higher in BAL fluid of Df–sensitized and -challenged P2ry6fl/fl/Cre/+ mice than +/+ controls (Figure 1D). Whole-lung Il33 mRNA expression in response to stimulation with Df (Supplemental Figure 1D).
To determine whether P2Y6 receptor signaling protected against T2I-mediated inflammation when P2ry6 deletion was restricted to the challenge phase, we delayed administration of tamoxifen to the mice to 24 hours after the second sensitization. BAL fluids and higher levels of the corresponding transcripts in the T cell compartment (CD3+CD90+CD45+) of the lungs. In contrast, there were no differences between strains in the cytokines generated by ILCs (Lin CD90+CD45+) (Supplemental Figure 3, A–C).
were significantly lower in the lungs of 
P2ry6 fl/fl/Cre/+ mice than +/+ controls when 
P2ry6 deletion was restricted to the 
challenge phase (Supplemental Figure 5A). In contrast, these 
transcripts did not differ between strains when 
P2ry6 was 
deleted before sensitization (Supplemental Figure 5B). These 
data demonstrated that P2Y6 signaling during the allergen 
sensitization phase was critical for its ability to decrease T2I-
mediated inflammation at the challenge phase.

P2Y6 expression by hematopoietic cells protects against Df-induced 
T2I. To identify specific cell types potentially responsible for the 
protective effect of P2Y6 receptor signaling during sensitization, 
we applied the sensitization and challenge protocol to 2 additional 
strains of conditional null mice: P2ry6 fl/fl 
Scgb1a1-Cre/+ mice, in 
which deletion of 
P2ry6 is driven by an epithelial (Clara) cell–specific 
ing dose of Df (Figure 1F). We verified deletion of P2ry6 in the 
lung and spleen by qPCR in this protocol (Supplemental Figure 
4A). In marked contrast to the deletion of 
P2ry6 prior to sensi-
tization, deletion after the sensitization significantly decreased 
Df-induced accumulation of eosinophils in BAL fluid (Figure 1G).
Additionally, levels of IL-5 and quantities of 
Il5, Il13, and Il33 transcripts were significantly diminished in the BAL and lungs of 
P2ry6 fl/fl/Cre/+ mice compared with +/+ controls after delayed 
tamoxifen administration (Figure 1H and Supplemental Figure 
4B). Total IgE, IgG2c, and IgM levels were similar in the 2 
genotypes in this protocol and not significantly increased by 
Df challenge (Figure 1I and Supplemental Figure 4C). Levels 
of mRNA encoding 
Cxcl1 and Il6 (each of which is generated 
by airway epithelial cells activated by P2Y6 signaling) (26)
were significantly lower in the lungs of P2ry6 fl/fl/Cre/+ mice 
than +/+ controls when P2ry6 deletion was restricted to the 
challenge phase (Supplemental Figure 5A). In contrast, these 
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**Figure 2.** P2Y6 receptors on hematopoietic cells are involved in protection from Df-induced allergic inflammation. (A) BAL fluid cell counts of NaCl- or 
Df-treated +/+ (blue bars; n = 8, n = 11) and P2ry6 fl/fl Scgb1a1-Cre/+ (red bars; n = 3, n = 14) mice on day 16. (B) BAL fluid cell counts of NaCl- or Df-treated 
+/- (n = 6, n = 13) and P2ry6 fl/fl Cd4-Cre/+ (n = 5, n = 15) mice on day 16. (C) BAL fluid cell counts of chimeric mice. Lethally irradiated mice of the indicated 
genotypes received i.v. marrow from mice of the indicated genotypes (10–14 mice/group). P2ry6 transcripts in BM and lung tissue and Epx transcript in 
lung tissue from chimeric mice on day 16, as measured by qPCR. Data are expressed as ratio relative to Gapdh. Values are mean ± SEM. All data are from 2 
experiments. *P < 0.05, ****P < 0.0001 vs. indicated mice receiving +/+ BM by Mann-Whitney U test, parametric t test, or Welch’s t test (C).
The engrafted mice confirmed the repopulation of recipient BM with the donor genotype. Additionally, transferred BM accounted for approximately 80% of the \textit{P2ry6} transcripts in the lung of \textit{Df}-sensitized and -challenged mice, indicating replenishment of \textit{P2Y6} receptors expressed by radiosensitive hematopoietic cell populations resident in the lung (Figure 2C).

\textit{Epx} transcript levels in the lungs of recipients of \textit{P2ry6fl/fl/Cre/+} marrow were increased compared with those in recipients of +/+ BM (Figure 2C). We concluded from these experiments that the regulatory effect of \textit{P2Y6} signaling was mediated by a hematopoietic non–T cell.

Figure 3. \textit{P2Y6} receptors regulate early alveolar IL-12 production. (A) Schematic of BAL and serum collection during the sensitization phase to assess levels of IL-12p40 and UDP and BAL fluid cell counts. (B) Time course of IL-12p40 production in BAL fluid from \textit{Df}-treated WT mice (\textit{n} = 3). (C) IL-12p40 production in BAL fluid and serum from NaCl- or \textit{Df}-treated +/+ (blue bars) and \textit{P2ry6fl/fl/Cre/+} (red bars) mice; and \textit{Il12b} transcript in the lung of NaCl- or \textit{Df}-treated +/+ and \textit{P2ry6fl/fl/Cre/+} mice on day 2. (D) \textit{P2Y6} activity (UDP concentrations) on days 0 (3 hours), 2, and 7 in BAL fluid was quantified by bioassay using 1321N1 cells stably expressing mouse \textit{P2Y6}. (E) BAL fluid cell counts of NaCl- or \textit{Df}-treated +/+ and \textit{P2ry6fl/fl/Cre/+} mice on day 2. Values are mean ± SEM. Data are from at least 2 experiments, except for the time course experiment on IL-12p40 (B) and \textit{P2Y6} activity on day 7 (D, right panel). *\textit{P} < 0.05, **\textit{P} < 0.01, ***\textit{P} < 0.001 vs. NaCl treatment; ****\textit{P} < 0.0001 vs. \textit{NaCl} treatment; **\textit{P} < 0.01, ***\textit{P} < 0.001 vs. +/+ mice; by 1-way (B) or 2-way (C and E) ANOVA followed by Tukey’s multiple-comparisons test or nonparametric Kruskal-Wallis followed by Dunn’s multiple-comparisons test (D).

Promoter; and \textit{P2ry6fl/+ Cd4-Cre/+} mice, in which \textit{P2ry6} is deleted selectively on T cells at the double-positive thymocyte stage; along with respective control strains. Compared with the corresponding +/+ controls, neither conditional null strain showed significant differences in total BAL fluid cell counts or any specific cell subset (Figure 2, A and B). To determine whether hematopoietic or tissue-resident expression of \textit{P2Y6} mediated the protective effect of the receptor during sensitization, we transferred BM from \textit{P2ry6fl/+Cd4-Cre/+} mice into lethally irradiated +/+ controls and vice versa, and subjected the engrafted mice to the \textit{Df} sensitization and challenge protocol (Figure 2C). Both +/+ and \textit{P2ry6fl/+Cd4-Cre/+} recipients of BM from \textit{P2ry6fl/+Cd4-Cre/+} mice showed higher BAL fluid total cell and eosinophil counts than did recipients of +/+ BM (Figure 2C). qPCR analysis of the engrafted mice confirmed the repopulation of recipient BM with the donor genotype. Additionally, transferred BM accounted for approximately 80% of the \textit{P2ry6} transcripts in the lung of \textit{Df}-sensitized and -challenged mice, indicating replenishment of \textit{P2Y6} receptors expressed by radiosensitive hematopoietic cell populations resident in the lung (Figure 2C). \textit{Epx} transcript levels in the lungs of recipients of \textit{P2ry6fl/+Cd4-Cre/+} marrow were increased compared with those in recipients of +/+ BM (Figure 2C). We concluded from these experiments that the regulatory effect of \textit{P2Y6} signaling was mediated by a hematopoietic non-T cell.

Early production of IL-12 depends on \textit{P2Y6} receptor signaling. We next sought to further characterize the early \textit{P2Y6} receptor-dependent events responsible for the dampened type 2 immune
responses during sensitization. We examined BAL fluid from P2ry6fl/fl/Cre/+ mice and +/+ controls for the presence of cytokines produced by resident hematopoietic cells during innate immune responses that regulate subsequent adaptive responses after administration of tamoxifen and a sensitizing dose of Df (Figure 3A). We measured IL-12p40, a subunit of both the Th1-promoting factor IL-12 and the Th17-promoting factor IL-23, during the sensitization phase. We also examined the BAL fluid for IL-23 (using an ELISA with an IL-12p19 capture antibody and an IL-12p40 detection antibody); IL-18, which can facilitate type 1 cytokine production (34); and IL-10, which is immunoregulatory. IL-12p40 levels in the BAL fluid tended to increase as early as 3 hours after the first sensitizing dose of Df in +/+ but not P2ry6fl/fl/Cre/+ mice, reaching a peak on day 2 (16 hours after the second sensitizing dose) and declining thereafter (Figure 3, B and C). Such an increase in BAL fluid IL-12p40 levels did not occur in P2ry6fl/fl/Cre/+ mice (Figure 3C). Serum levels of IL-12p40 were equivalent in the 2 strains and not altered by sensitization to Df (Figure 3A), and whole-lung Il12b transcripts were modestly reduced in the P2ry6fl/fl/Cre/+ mice relative to +/+ controls. IL-12p40 levels were below the limit of detection in the BAL fluid (data not shown). IL-12p40 levels in lung homogenates were similarly low in Df- and NaCl-treated mice, and did not differ between strains (Supplemental Figure 6A). Df treatment induced modest and equivalent increases in BAL fluid levels of IL-18 in both strains on day 2, without significant changes in serum IL-18 levels or Il18 mRNA expression in the lungs (Supplemental Figure 6B).

To quantify the levels of extracellular UDP in the alveolar space, we developed a P2Y6 receptor–dependent calcium flux bioassay in stably transfected mouse P2Y6-expressing 1321N1 cells capable of detecting UDP in BAL fluid at levels as low as 0.2 nM. Both UDP and BAL fluid induced calcium fluxes in P2Y6-transfected cells, but not parental 1321N1 cells, that were prevented by treatment with apyrase (Supplemental Figure 7, A–C). Among the 4 nucleotides, UTP and ADP showed more than 30 and 300 times lower affinity, respectively, to mouse P2Y6 than UDP (Supplemental Figure 7D). In mice of both genotypes, the first sensitizing dose of Df increased UDP activity in BAL fluid collected compared with the NaCl-challenged controls as early as 3 hours (Figure 3D). These increases persisted on day 2, then declined by day 7 to basal levels (Figure 3D). BAL fluid UDP levels in the Df-treated P2ry6fl/fl/Cre/+ mice were indistinguishable from those in +/+ controls at 3 hours and day 2, but were higher in the P2ry6fl/fl/Cre/+ mice on day 16 in both the before-sensitization and delayed-tamoxifen protocols (Supplemental Figure 7E). Total cell counts and neutrophils in the BAL fluid on day 2 increased in +/+ but not P2ry6fl/fl/Cre/+ mice (Figure 3E). We concluded from these experiments that the release of IL-12p40 during the sensitization
phase coincided with the release of UDP, as well as a small influx of neutrophils, and that IL-12p40 release and neutrophil influx both depended on P2Y6 signaling.

To determine whether the P2Y6 receptor–dependent differences in BAL fluid IL-12p40 levels persisted at the challenge phase, we measured IL-12p40 levels in samples from NaCl- and Df-challenged mice on day 16. Compared with NaCl-challenged controls, BAL fluid from Df-challenged mice showed markedly increased levels of IL-12p40, but with no significant differences between strains (Supplemental Figure 8). Thus, P2Y6 receptor
signaling was essential for the early production of IL-12p40 but not for its production after sensitization and when airway inflammation was established.

We next asked whether IL-12 production at the sensitization phase conveyed the protective effect of P2Y6 receptor signaling.

![Image of Figure 6](image_url)

**Figure 6.** Pulmonary +/+ macrophage transplantation restores Df-induced IL-12p40 production and neutrophil infiltration during sensitization of P2ry6fl/fl/Cre/+ mice. (A) BM macrophages from +/+ or P2ry6fl/fl/Cre/+ mice were administered (i.t.) into +/+ and P2ry6fl/fl/Cre/+ mice: +/+ macrophages into +/+ mice (+/+→+/+, n = 11); P2ry6fl/fl/Cre/+→+/+ (n = 4); P2ry6fl/fl/Cre/+→P2ry6fl/fl/Cre/+ (n = 7); and +/+→P2ry6fl/fl/Cre/+ (n = 7). (B) P2ry6 transcript in BAL fluid cells of pulmonary macrophage transplantation–treated (PMT-treated) +/+ and P2ry6fl/fl/Cre/+ mice on day 2 (17 days after transplantation). ***P < 0.01 by parametric t test. (C) BAL fluid IL-12p40 from PMT-treated +/+ and P2ry6fl/fl/Cre/+ mice on day 2. (D) Diff-Quick–stained BAL fluid cells from the indicated mice. Red arrows indicate neutrophils. Scale bar: 50 µm. (E) Percentages of neutrophils in BAL fluid of PMT-treated +/+ and P2ry6fl/fl/Cre/+ mice. (F) MPO levels in BAL fluid of PMT-treated +/+ and P2ry6fl/fl/Cre/+ mice on day 2. (G) BAL cell Mpo transcript levels of PMT-treated mice on day 2. Values are mean ± SEM. Data are from 2 experiments. *P < 0.05, **P < 0.01 vs. +/+→+/+ mice; *P < 0.05, ***P < 0.001 vs. P2ry6fl/fl/Cre/+→P2ry6fl/fl/Cre/+ mice by 1-way ANOVA followed by Tukey’s multiple-comparisons test (C–G).
Df-challenged P2ry6fl/fl/Cre/+ mice were sharply reduced compared with those of their counterparts that did not receive IL-12p70 (Figure 4C). Furthermore, IL-12p70 administration tended to increase serum levels of IgG2c in the Df-challenged P2ry6fl/fl/Cre/+ mice to levels comparable to those observed in +/+ controls (Figure 4D). From these experiments, we concluded that IL-12 generated at sensitization was necessary for P2Y6 receptor–dependent control of type 2 immunopathology and IgG2c production at challenge.

Figure 7. P2Y6-dependent IL-12 from AMs induces a protective IFN-γ signature. (A) Ifng transcripts in BAL fluid cells and lungs from NaCl- or Df-treated +/+ and P2ry6fl/fl/Cre/+ mice on day 2. **P < 0.01 vs. +/+ mice. (B) Ifng transcripts in BAL fluid cells from mice engrafted with +/+ or P2ry6fl/fl/Cre/+ macrophages on day 2. *P < 0.05 vs. +/+ recipient mice with +/+ macrophages (i.t.); ***P < 0.001 vs. P2ry6fl/fl/Cre/+ recipient with P2ry6fl/fl/Cre/+ macrophages (i.t.). (C) Ifng transcripts in CD11c+ and CD11c- BAL fluid cells of P2ry6fl/fl/Cre/+ mice that received vehicle or recombinant IL-12p70. ****P < 0.0001 vs. vehicle treatment. (D) BAL fluid cell counts of Df-treated WT and Ifng−/− mice on day 16. (E) Representative lung sections from Df-treated WT and Ifng−/− mice were stained with H&E on day 16. Scale bar: 300 µm. (F) Total serum IgG2c of Df-treated WT and Ifng−/− mice quantified on day 16. (G) Levels of IL-4, IL-5, IL-13, and of Epx transcript in BAL fluid cells of NaCl- or Df-treated WT and Ifng−/− mice. Values are mean ± SEM. *P < 0.05, **P < 0.01, ***P < 0.001 vs. WT mice by 1-way (B–G) or 2-way (A) ANOVA followed by Tukey’s multiple-comparisons test.

Similar to those detected in Df-sensitized WT mice (217 ± 23 pg/mL) (Figure 4A). On day 16, IL-12p70–treated P2ry6fl/fl/Cre/+ mice showed decreased BAL fluid total cell and eosinophil, but not neutrophil, numbers compared with sensitized and challenged P2ry6fl/fl/Cre/+ mice that did not receive IL-12p70 (Figure 4B). Exogenous IL-12p70 did not alter total or differential BAL fluid cell counts in Df-challenged +/+ mice compared with control mice not receiving IL-12p70. Epx transcript levels in the lungs of IL-12p70–treated
AMs are the dominant source of P2Y₆-regulated IL-12. To identify the cellular source(s) of UDP/P2Y₆ receptor-dependent IL-12p40 production, we analyzed cells from both BAL fluid and enzymatically dispersed lungs for intracellular IL-12p40 protein by flow cytometry. Samples were collected from +/+ and P2ry6⁻⁻/⁻/Cre/⁻ mice before and 4 and 8 hours after the first and second sensitizations with Df. The cells were permeabilized and stained with mAbs against specific lineage markers and IL-12p40, respectively. Before sensitization, approximately 95% of BAL fluid cells in both +/+ and P2ry6⁻⁻/⁻/Cre/⁻ mice were CD68⁺ macro-
Figure 9. CysLT1R antagonists inhibit early P2Y6 receptor–dependent signaling that counterbalances CysLT1R-induced T2I priming function. (A) CysLT1R antagonists blocked UDP elicited calcium flux induced in 1321N1 cells stably expressing mouse P2ry6 (montelukast, zafirlukast, and pranlukast) and the macrophage cell line RAW 264.7 (MK-571). Values are mean (n = 4). (B) The CysLT1R antagonists all inhibited the UDP elicited (10 μM) potentiation of Df-induced IL-12p40 generation in BM-derived macrophages (n = 8). (C) Effect of zafirlukast (i.p.) or MK-571 (i.n.) at sensitization on the BAL concentration of IL-12p40 and expression of Ifng transcripts by BAL fluid cells. The correlation between BAL concentration of IL-12p40 and Ifng transcripts is shown. (D) BAL fluid cell counts of NaCl- or Df-treated +/+ and P2ry6fl/fl/Cre/+ mice administered zafirlukast (i.p.) at sensitization (+/+ Veh: n = 12; +/+ zafirlukast: n = 12; P2ry6fl/fl/Cre/+ Veh: n = 17; and P2ry6fl/fl/Cre/+ zafirlukast: n = 16). ***P < 0.001, ****P < 0.0001, *P < 0.05, **P < 0.01, ***P < 0.001, ****P < 0.0001 by 1-way ANOVA followed by Tukey’s multiple-comparisons test. Values are mean ± SEM. Data are from 2 experiments, except for the calcium flux experiments.
phages, of which approximately 30% expressed IL-12p40 (Figure 5A). The percentages of CD68+ macrophages expressing IL-12p40 in +/+ mice increased by approximately 60% by 4 hours after the second sensitizing dose of Df and remained higher than in NaCl-treated controls to day 2 (24 hours after the second sensitizing dose) (Supplemental Figure 9A and Figure 5A), while the percentages of IL-12p40+ CD68+ cells decreased. The percentage of CD68+ cells expressing IL-12p40+ did not increase significantly in P2ry6fl/fl/Cre/+ BMMs as compared to +/+ mice in response to Df (Figure 5A, top row). All of the IL-12p40+CD68+ cells expressed CD11c (Figure 5A, middle row), consistent with their identity as AMs. At 4 hours after the second sensitizing dose, Df caused an influx of Ly6G+ neutrophils and Ly6G+ lymphocyte-sized cells into the BAL fluid of +/+ but not P2ry6fl/fl/Cre/+ mice, both of which were negative for IL-12p40 (Figure 5A, bottom row). Single-cell suspensions of dispersed lungs contained interstitial macrophages (CD45+CD24−CD11b−CD11c+) that were IL-12p40 negative (Figure 5B). The eosinophils (CD45+CD24+Siglec F−) and neutrophils (CD45+CD24+Siglec F CD11c+DD11b+) present in small numbers in the lungs after Df sensitization were autofluorescent, and the IL-12p40+ positive plasmacytoid DCs (CD45+CD24+Siglec F CD11c+) did not differ between +/+ and P2ry6fl/fl/Cre/+ mice (Figure 5B). CCR2+ inflammatory monocytes (CD45+CCR2+Ly6C+), which showed higher levels of P2ry6 transcript than BAL fluid cells, infiltrated into the lung after the second sensitizing dose of Df, but they were negative for IL-12p40 (Supplemental Figure 9, B and C).

To confirm that macrophage-intrinsic UDP/P2Y6 receptor signaling mediated the early Df-induced production of IL-12p40, we modified an AM transplantation protocol that does not involve myeloablation (35). BMMs were derived in vitro from the BMs of +/+ and P2ry6fl/fl/Cre/+ mice (Figure 6A, middle row), consistent with their identity as AMs. At 4 hours after the second sensitizing dose, Df caused an influx of Ly6G+ neutrophils and Ly6G+ lymphocyte-sized cells into the BAL fluid of +/+ but not P2ry6fl/fl/Cre/+ mice, both of which were negative for IL-12p40 (Figure 5A, top row). All of the IL-12p40+CD68+ cells expressed CD11c (Figure 5A, middle row), consistent with their identity as AMs. At 4 hours after the second sensitizing dose, Df caused an influx of Ly6G+ neutrophils and Ly6G+ lymphocyte-sized cells into the BAL fluid of +/+ but not P2ry6fl/fl/Cre/+ mice, both of which were negative for IL-12p40 (Figure 5A, bottom row). Single-cell suspensions of dispersed lungs contained interstitial macrophages (CD45+CD24−CD11b−CD11c+) that were IL-12p40 negative (Figure 5B). The eosinophils (CD45+CD24+Siglec F−) and neutrophils (CD45+CD24+Siglec F CD11c+DD11b+) present in small numbers in the lungs after Df sensitization were autofluorescent, and the IL-12p40+ positive plasmacytoid DCs (CD45+CD24+Siglec F CD11c+) did not differ between +/+ and P2ry6fl/fl/Cre/+ mice (Figure 5B). CCR2+ inflammatory monocytes (CD45+CCR2+Ly6C+), which showed higher levels of P2ry6 transcript than BAL fluid cells, infiltrated into the lung after the second sensitizing dose of Df, but they were negative for IL-12p40 (Supplemental Figure 9, B and C).

To confirm that macrophage-intrinsic UDP/P2Y6 receptor signaling mediated the early Df-induced production of IL-12p40, we modified an AM transplantation protocol that does not involve myeloablation (35). BMMs were derived in vitro from the BMs of +/+ and P2ry6fl/fl/Cre/+ mice in the presence of M-CSF and GM-CSF. Two million BMMs were delivered intratracheally to cohorts of recipient P2ry6fl/fl/Cre/+ mice. Fifteen days later, the engrafted mice were sensitized with Df, and percentages of IL-12p40 were measured on day 2 (Figure 6A). qPCR of BAL fluid cells confirmed the presence of the P2ry6 transcript in recipients of +/+ macrophages, albeit at lower levels than in +/+ mice (Figure 6B). Df-treated P2ry6fl/fl/Cre/+ recipient mice engrafted with +/+ BMMs showed markedly increased levels of IL-12p40 in the BAL fluid compared with P2ry6fl/fl/Cre/+ recipient mice engrafted with P2ry6fl/fl/Cre/+ BMMs, approximating the levels observed in Df-treated +/+ controls (Figure 6C). In contrast, P2ry6fl/fl/Cre/+ mice receiving P2ry6fl/fl/Cre/+ BMMs showed much lower levels of BAL fluid IL-12p40. Additionally, recipients of +/+ BMMs, but not recipients of P2ry6fl/fl/Cre/+ BMMs, showed increased percentages of neutrophils (Figure 6, D and E), increased levels of MPO in BAL fluid (Figure 6F), and increased levels of Mpo mRNA in BAL fluid cells (Figure 6G) after 2 sensitizing doses of Df.

P2Y6 receptor signaling potentiates dectin-2-dependent IL-12p40 generation by macrophages in response to Df. To determine whether UDP/P2Y6 receptor signaling directly altered IL-12p40 generation by macrophages, we treated BMMs from +/+ and P2ry6fl/fl/Cre/+ mice with UDP and Df. UDP alone was unable to increase the secretion of IL-12p40 from BMMs (Supplemental Figure 10A). However, Df-induced secretion of IL-12p40 by +/+ BMMs was amplified after UDP priming for 6 hours. This UDP priming effect was abolished in BMMs from P2ry6fl/fl/Cre/+ mice.
significantly increased \( \text{Ifng} \) expression by BAL fluid cells in both treatments. Only administration of IL-23 after a single sensitizing dose of \( Df \) induced \( Il17a \) gene expression (Supplemental Figure 12B). In contrast, treatment with a neutralizing anti-IL-12R\( \beta 2 \) mAb directed against a specific subunit of the IL-12p70 receptor significantly decreased recombinant IL-12p70–induced (ex vivo) and \( Df \)-induced (in vivo) \( \text{Ifng} \) transcript levels in BAL fluid cells (Supplemental Figure 12C). To determine whether IFN-\( \gamma \) production at sensitization was essential for \( P2y_6 \)/IL-12–dependent suppression of T2I on day 16, we sensitized \( \text{Ifng}^{-/-} \) mice and WT controls with \( Df \) and compared their responses with subsequent \( Df \) challenges. Compared with WT controls, \( \text{Ifng}^{-/-} \) mice showed sharply increased numbers of BAL fluid total cells, eosinophils, and neutrophils on day 16 (Figure 7D). Histological examination confirmed that bronchovascular inflammation in \( Df \)-sensitized and -challenged \( \text{Ifng}^{-/-} \) mice was more extensive than in WT controls (Figure 7E). Serum IgG2c was nearly undetectable in \( \text{Ifng}^{-/-} \) mice (Figure 7F). Levels of IL-4, IL-5, and IL-13 in BAL fluid and BAL fluid cell expression of \( Il4 \) and \( Il13 \) were significantly higher in \( \text{Ifng}^{-/-} \) mice than in WT controls (Figure 7G and Supplemental Figure 13).

To identify the major cell source(s) of protective IFN-\( \gamma \) present at sensitization, we focused on the Ly6c\(^{-} \) lymphocyte population (CD11c\(^{+} \) population) that was recruited to the alveolar spaces of sensitized +/+ mice and that was largely absent in \( P2ry6^{+/+ Cre^{-/+}} \) mice (Figure 5A, bottom panels). We identified 4 populations within these lymphocytes based on staining for CD3 and NK1.1, with the 2 most abundant groups being CD3 NK1.1\(^{+} \) and CD3\(^{+} \)NK1.1\(^{-} \), respectively (Figure 8A). Both the CD3 NK1.1\(^{+} \) and CD3\(^{+} \)NK1.1\(^{-} \) subsets strongly expressed \( \text{Ifng} \) (relative to \( \text{Gapdh} \)), but the CD3 NK1.1\(^{+} \) subset was the dominant source of the transcript based on the differences in cell numbers (Figure 8A). The CD3 NK1.1\(^{+} \) subset also showed greater mRNA expression of IL-12R\( \beta 2 \) (Supplemental Figure 14A). Intranasal administration of an anti-NK1.1 mAb, which recognizes NK and NKT cells, substantially decreased the quantity of \( \text{Ifng} \) mRNAs transcripts detected in BAL fluid cells (Figure 8B). BAL fluid cells from \( Cd1d^{-/-} \) mice lacking NKT cells showed no decrement in \( Df \)-induced \( \text{Ifng} \) expression compared with WT controls (Figure 8C). Anti-NK1.1 treatment selectively depleted CD3 CD49\(^{+} \) and CD3 NKp46\(^{+} \) lymphocyte populations from the BAL fluid, and sharply decreased expression of \( Nkg2d \), without altering the numbers of CD3\(^{+} \) cells or expression of \( Il2 \) (Supplemental Figure 14, B and C). Compared with \( Df \)-sensitized +/+ mice, BAL fluid from \( P2ry6^{+/+ Cre^{-/+}} \) mice contained markedly decreased numbers of CD3 NK1.1\(^{+} \) and CD3 NKp46\(^{+} \) cells, as well as fewer CD3\(^{-} \) cells (Figure 8D). Treatment of WT mice with i.n. anti-NK1.1 during sensitization sharply increased BAL fluid total cell and eosinophil counts, bronchovascular inflammation, and type 2 cytokine expression after \( Df \) challenge compared with WT mice receiving the isotype control (Figure 8E and Supplemental Figure 14D). In contrast, anti-NK1.1 at sensitization did not alter the expression of \( \text{Ifng} \) or \( Nkg2d \) in BAL fluid cells after \( Df \) challenge on day 16. \( Df \)-challenged NKT cell–deficient \( Cd1d^{-/-} \) and WT mice showed similar numbers of BAL fluid eosinophils on day 16 (Supplemental Figure 14E). We concluded from these experiments that IFN-\( \gamma \)-producing NK cells, and not NKT cells, were the major downstream targets of the \( P2y_6 \)-dependent, AM-derived IL-12 involved in the suppression of inappropriate type 2 responses.

CysLT\(_R1\) antagonists suppress early \( P2y_6 \) receptor–dependent signaling and IL-12p40 generation that counterbalances CysLT\(_R1\)-induced T2I priming function. \( P2y_6 \) receptors are structural homologs of the CysLT-specific GPCRs CysLT\(_R1\), CysLT\(_R2\), and CysLT\(_R3\) (18, 20). CysLT\(_R1\) is central to Dectin-2–mediated DC priming for T2I (33), and clinically active antagonists of this receptor can block \( P2y_6 \) receptor signaling in transfect ed cells (21). We sought to determine whether CysLT\(_R1\) antagonists block natively expressed \( P2y_6 \) receptors, and if so whether this blockade modifies protective \( P2y_6 \) receptor signaling at the initiation of T2I in vivo. The CysLT\(_R1\) antagonists montelukast, zafirlukast, pranlukast, and MK-571 all inhibited UDP-elicited calcium flux in 132.1N1 cells stably expressing mouse \( P2y6 \) (montelukast, zafirlukast, and pranlukast) or the macrophage cell line RAW 264.7 (MK-571) (Figure 9A). All 4 antagonists also blocked the UDP-elicited potentiation of \( Df \)-induced IL-12p40 generation by BMMs (Figure 9B). Montelukast also reduced IL-12p40 generation in response to \( Df \) alone at the highest dose tested. Exogenous LTC\(_4\), LTD\(_4\), and LTE\(_4\), did not potentiate IL-12p40 production by BMMs, and UDP did not induce cysLT generation by BMMs (data not shown). Administration of either zafirlukast (i.p.) or MK-571 (i.n.) to WT mice during \( Df \) sensitization significantly reduced the concentration of IL-12p40 in BAL fluid and decreased expression of \( \text{Ifng} \) by BAL fluid cells (Figure 9C). IL-12p40 levels and \( \text{Ifng} \) transcripts were tightly correlated (Figure 9C). Administration of zafirlukast at sensitization markedly decreased the numbers of BAL fluid total cells, eosinophils, and neutrophils, bronchovascular inflammation, and type 2 cytokine and Epx expression induced by \( Df \) on day 16 in \( P2ry6^{+/+ Cre^{-/+}} \) mice, but did not affect the modest levels seen in +/+ controls (Figure 9D and Supplemental Figure 15, A and B).

**Discussion**

Innate responses of tissue-resident cells to allergens and pathogens instruct subsequent adaptive immune responses, and determine whether subsequent encounters with the same antigens and pathogens will elicit T2I. The broad distribution of P2 receptors (36) — and the availability of their ligands resulting from cellular damage, stress, or activation — suggests the potential for P2 receptor signaling in innate immunity. While the role of ATP in the initial sensitization stage of T2I is well recognized (4, 22), less is known about the role of UDP and its cognate receptor. Extracellular concentrations of uracil nucleotides are lower than those of ATP (37), but the high affinity of P2Y\(_6\) receptors and their expression by both structural and hematopoietic cells of the lung suggest that these cells are highly sensitive to extracellular UDP. While pharmacologic blockade of P2Y\(_6\) receptors blunts T2I in sensitized mice (26), genetic deletion of P2Y\(_6\) receptors sharply enhances T2I and associated eosinophilic inflammation in response to repetitive, low-dose inhalation challenges with \( Df \), an allergen strongly linked to asthma in humans (31). Moreover, selective agonism of P2Y\(_6\) receptors suppresses inflammation and airway reactivity in mice (32). We therefore undertook this study to understand the mechanisms and cellular targets that account for the apparent protective effects of P2Y\(_6\) receptor signaling in the lung, and to explain the contradictory results. Moreover, we sought to determine the potential consequences of off-target P2Y\(_6\) blockade by clinically available CysLT\(_R1\) antagonists in vivo,
particularly in a context where endogenous cysteine leukotrienes play a major role in sensitization for T21.

We modified our protocol to incorporate temporally distinct sensitization and challenge phases to account for the possibility that P2Y6 receptors may serve very different functions during the initial and subsequent responses to antigen exposure (Figure 1A). Deletion of P2ry6 prior to the sensitization phase in this modified model mirrored the enhanced lung T2I in the P2ry6fl/fl/Cre/+ mice that we had observed in our prior studies using a protocol without distinct sensitization and challenge phases (ref. 31 and Figure 1). In contrast, deletion of P2ry6 after sensitization sharply reduced eosinophilia and type 2 cytokine production (Figure 1). Levels of mRNA encoding Il6 and the CXCR2 ligand Cxcl1 (both of which were previously identified as major P2Y6 receptor–driven products of lung epithelium [ref. 26]) were significantly reduced in the lungs of P2ry6fl/fl/Cre/+ mice subjected to gene deletion after, but not before, sensitization (Supplemental Figure 5). Combined with the observed increase in BAL fluid UDP occurring within 3 hours of the initial sensitizing dose of Df (Figure 3D), these observations support an early (and likely innate) cellular target of P2Y6 receptor activation as a prime modifier of the subsequent immune response. Our bioassay showed that P2Y6–induced calcium flux induced by BAL fluids completely disappeared after pretreatment with the nucleotide-hydrolyzing enzyme apyrase (Supplemental Figure 7B). Although we cannot exclude the possibility that nucleotides other than UDP activate the P2Y6 receptor in vivo, UTP (which is rapidly converted to UDP in vivo) and ADP showed more than 30 and 300 times lower affinity, respectively, than UDP for mouse P2Y6 (Supplemental Figure 7D). It is thus likely that UDP is the most relevant ligand driving the activation of P2Y6 receptors. While the effector function of P2Y6 receptors in the challenge phase may well reflect control of epithelial cell–derived proinflammatory factors, as suggested by previous studies (26), the basis of its protective functions during sensitization became the focus of our subsequent experiments.

We used a combination of conditionally gene-deleted mice and cell transfers to identify the target cells and mechanisms responsible for the protective effect of P2Y6 receptors in the sensitization phase. Sgrbla1–driven P2ry6 deletion failed to potentiate inflammation, indicating that the protective effect of P2Y6 receptor signaling was not due to Clara cell–intrinsic signaling (Figure 2A). Although we had previously demonstrated that P2Y6 receptors were inducibly expressed by CD4+ and CD8+ T cells after challenge with Df (31), constitutive Cd4–driven P2ry6 deletion (which eliminates gene expression at the double-positive state of thymic T cell development) did not recapitulate the effect of global P2ry6 deletion proximal to sensitization (Figure 2B). Results of experiments involving adoptive transfer of BM into lethally irradiated mice not only supported the involvement of P2Y6 receptors expressed by hematopoietic cells in the protective response (Figure 2C), but also revealed that hematopoietic cells accounted for approximately 80% of the P2ry6 transcript expressed by the lung on day 16. We therefore focused our attention on the potential role of P2Y6 receptors on lung-resident hematopoietic cells and on whether P2Y6 receptor signaling was critical to their generation of products that regulate immune responses during the initial exposure to antigens.

Macrophages and CD8+ DCs, the innate hematopoietic cells most strongly expressing P2Y6 receptors (http://rstats.imgen. org/Skyline/skyline.html), generate cytokines on initial encounters with antigens that can inform adaptive responses. IL-12 generated at sensitization has a known protective effect against inappropriate T2I (38), although the cell sources and regulatory mechanisms are not fully understood. Our study indicates that UDP/P2Y6 signaling facilitates IL-12 generation by AMs as a prominent component of P2Y6–dependent protection from T2I. BAL fluid levels of IL-12p40 (a subunit of both IL-12 and IL-23), but not levels of IL-23, IL-10, or IL-18, increased shortly after administration of sensitizing doses of Df in +/- but not P2ry6fl/fl/Cre/+ mice, without corresponding changes in serum IL-12p40 (Figure 3C). This increase occurred concomitantly with increases in UDP (Figure 3D). Restoring physiologic concentrations of IL-12 at sensitization (Figure 4A) decreased BAL fluid total cell and eosinophil counts (Figure 4B) and tissue Epx expression (Figure 4C) in P2ry6fl/fl/Cre/+ mice to levels similar to those observed in +/- controls, and also increased serum IgG2c in P2ry6fl/fl/Cre/+ mice to +/- levels (Figure 4D) (in retrospect, a likely reflection of IFN-γ). Df sensitization increased expression of IL-12p40 protein specifically in CD68+CD11c+ AMs in a P2Y6 receptor–dependent manner (Figure 5A and Supplemental Figure 9), with minimal contributions from interstitial macrophages, DCs, and granulocytes (Figure 5B).

Finally, transfer of macrophages derived from the BM of +/-, but not P2ry6fl/fl/Cre/+, mice restored early Df–induced IL-12p40 (Figure 6), along with Df–induced neutrophil recruitment. Although exogenous UDP was insufficient alone to induce IL-12p40 generation, it substantially primed macrophages ex vivo and in vivo (Supplemental Figure 10) for Df–induced IL-12p40 generation. The absence of Df–induced IL-12p40 in AMs of Clec4n–/– mice (Supplemental Figure 11) supports the argument that branched mannans associated with Df act as direct ligands for Dectin-2 on macrophages (3) as they do on DCs (3). Notably, other major “Th2-sensitizing” allergens, such as those associated with Dermatophagoides farinae and Aspergillus fumigatus, require Dectin-2 signaling on DCs to elicit sensitization of Th2 cells and subsequent T2I (33). Our studies suggest that P2Y6 signaling facilitates Dectin-2–dependent AM activation, countering Th2 sensitization and its consequences through early generation of IL-12p40. Moreover, macrophage- and P2Y6–driven IL-12 is necessary to activate a population of NK cells that provide IFN-γ to counteract subsequent T2I at the challenge phase (Figures 7 and 8).

IL-12p40 can homodimerize to form IL-12p80, or can pair with IL-12p35 or IL-23p19 to form IL-12p70 or IL-23, respectively. Although we did not detect IL-12p70 (a known inducer of IFN-γ responses) in BAL fluid, we found a marked correlation between BAL IL-12p40 and IFN-γ production (Supplemental Figure 12A and Figure 9C). Our data suggest that IL-12p80 and IL-23 do not contribute to the innate Th1-like immune response during the sensitization phase (Supplemental Figure 12A). Exogenous IL-23, but not IL-12p70 or IL-12p80, induced IL-17a expression by BAL fluid cells (Supplemental Figure 12B). IL-17a expression was not induced by Df or P2Y6 during the sensitization phase (data not shown). Moreover, neutralizing anti-IL-12Rβ2 (a subunit of the IL-12p70 receptor) mAb inhibits the recombiant IL-12p70–induced (ex vivo) and Df–induced (in vivo) Ifng expression detected in BAL fluid cells (Supplemental Figure 12C). Although these data...
support the functional contribution of IL-12p70 to Ifng expression, we cannot exclude contributions from other P2Y6-regulated pathways. Indeed, levels of Th1/M1 cytokines/chemokines from AMs (including several that act on NK cells) were also decreased in P2ry6fl/fl/Cre/+ mice (Supplemental Table 1). Further investigations are needed to determine the potential contributions from these cytokines and chemokines to the integrated P2Y6-regulated IFN-γ response.

Clinically active CysLT1R antagonists achieve plasma concentrations of 0.5–1.5 μM when administered orally to human subjects, which are within the range reported to block human P2Y6 (and other P2Y receptors) in vitro (21). We confirmed that 4 different CysLT1R antagonists block calcium flux and potentiation of IL-12p40 elicited by mouse P2Y6 receptor signaling ex vivo, albeit at higher concentrations than those reported for the human counterpart (Figure 9, A–C). More importantly, administration of 2 different CysLT1R antagonists significantly impaired the production of IL-12p40 by AMs that leads to IFN-γ production by NK cells to Df sensitization, confirming that these drugs also block P2Y6 receptor signaling in vivo. Several clinically important allergens (Dermatophagoides farinae, Dermatophagoides pteronyssinus, Aspergillus fumigatus) associated with asthma prime DCs for T2I by a Dectin-2/CysLT1R pathway (3, 33). Our results suggest that the effects of this pathway are counteracted by Dectin-2/P2Y6 receptor signaling, and that absence or impairment of the latter pathway magnifies the effects of the former (and hence the ability of CysLT1R antagonists to block T2I). CysLT1R antagonists are FDA approved to treat asthma and rhinitis in children as young as 6 months (39). The incidence of de novo sensitization to inhalant antigens is high in children. Our findings could help to explain the widely variable individual responses to CysLT1R antagonists (15), and suggest that the development of drugs with greater selectivity for CysLT1R (or CysLT1R antagonists with P2Y6 receptor agonism) would confer a more robust and consistent therapeutic effect by protecting susceptible populations from T2I sensitization.

**Methods**

Further information can be found in Supplemental Methods, available online with this article.

**Animals.** P2ry6fl/fl/+ mice and P2ry6fl/fl/Cre/+ mice were derived as previously reported and backcrossed for 10 generations with C57BL/6 mice (31). To delete the P2Y6 receptor ubiquitously, we induced Rosa26 Cre recombinase in 10- to 19-week-old P2ry6fl/fl/Cre/+ mice (as shown in Figure 1C, Figures 3–8, and the supplemental figures) by administering tamoxifen (1 mg; MilliporeSigma) dissolved in a mixture of ethanol/sunflower seed oil (1:4, v/v) by i.p. injections on 5 consecutive days (Figure 1A). C57BL/6-P2ry6tm2450Arte mice were intercrossed with Cd4-Cre mice to generate P2ry6/Δ/Δ Cd4-Cre/+ mice and with Scbg1a1-Cre mice to generate P2ry6/Δ/Δ Scbg1a1-Cre/+ mice (as shown in Figure 2, A and B) at the University Medical Center Freiburg. Ifng−/− (B6.129S7-Ifngtm1Ts/J) and Cdlt−/− [B6.129S6-De1(C3d1d2-Cd1d11)1Bsp/J] mice were obtained from the Jackson Laboratory. Clec4n−/− mice were provided by Yoichiro Iwakura (Tokyo University of Science, Tokyo, Japan). C57BL/6 controls were obtained from the Jackson Laboratory and Charles River.

**Induction of allergic pulmonary inflammation and in vivo injection.** Ten days after the last administration of tamoxifen, allergic pulmonary inflammation was induced in +/+ and P2ry6fl/fl/Cre/+ mice by separate sensitization (20 μg i.n. on days 0 and 1) and challenge (3 μg i.n. on days 14 and 15) with Df (Greer Laboratories). Control groups of +/+ and P2ry6fl/fl/Cre/+ mice received 0.9% NaCl alone. After the Df treatment, mice were euthanized, and blood was collected by cardiac puncture and centrifuged to obtain serum. Lungs were lavaged 2 times with 1.0 mL PBS containing 0.5 mM EDTA. BAL fluid cells were cytocentrifuged onto slides, stained with Diff-Quick (Siemens Healthcare Diagnostics Inc.), and differentially counted. Left lungs were collected at the time of euthanasia, fixed with 4% PFA overnight, and embedded in glycol methacrylate. Sections were stained with H&E as previously reported (40). Light microscopic images were captured using a Leica DM LB2 light microscope with HC PL FLUOTAR 10×/0.30 (H&E stain) and 40×/0.75 (Diff-Quick stain) objective lens (Leica) and Nikon ACT-1 software at 22°C. Pictures were not further adjusted for brightness and contrast.

For restoration of IL-12 protein in the alveolar space, mice received i.n. administered recombinant murine IL-12p70 (500 ng, Thermo Fisher Scientific) on 4 consecutive days during the sensitization phase. For depletion of NK cells in the alveolar space, mice received i.n. administered 10 μg anti-NK1.1 (PK136, Thermo Fisher Scientific) or control mouse IgG2 (Thermo Fisher Scientific) on days 0 and 1. Intranasal injection of UDP (MilliporeSigma; 80 nmol in NaCl) was performed on days 0 and 1, or before (day 0) or after (day 1) a single sensitizing dose of Df (20 μg). To study the role of IL-12p40, we i.n. injected IL-12p70, IL-12p80 (p40 homodimer, 500 ng, BioLegend), and IL-23 (500 ng, BioLegend) on days 0 and 1 or after (day 1) a single sensitizing dose of Df (20 μg). Intranasal injection of neutralizing mouse IL-12Rβ2 antibody (10 μg in PBS, R&D Systems) and control IgG (R&D Systems) was performed on days 0 and 1. Zafirlukast (Cayman Chemical) for i.p. injection (30 mg/kg) was suspended in a 30% propylene glycol (PG) solution (70% NaCl/30% PG [PG/PEG400/Tween-80, 4:2:1]) before use. Control mice received the same volume of 30% PG alone. MK-571 (Cayman Chemical) was dissolved in NaCl (100 nmol/20 μL) and administered i.n.

**BMM and BM-derived DC generation.** For bone BMM generation, BM cells obtained from tamoxifen-treated +/+ and P2ry6fl/fl/Cre/+ mice on day 10 were harvested, and this suspension (4 × 106 cells/10 mL) was cultured in DMEM/F12 medium supplemented with 10% heat-inactivated FBS, 100 U/mL penicillin, 100 μg/mL streptomycin, 2 mM L-glutamine, and 100 ng/mL recombinant mouse M-CSF (R&D Systems). On day 3, 5 mL complete medium with M-CSF was added, and the adherent cells were harvested on day 6 or 7. For IL-12p40 measurement ex vivo, BMMs were plated in 96-well plates in DMEM/F12 medium supplemented with 10% FBS, 100 U/mL penicillin, and 100 μg/mL streptomycin and cultured overnight. BMDCs were generated with GM-CSF, as previously described (3). In brief, BM cells were cultured in RPMI medium with 10% heat-inactivated FBS, 100 U/mL penicillin, 100 μg/mL streptomycin, 2 mM L-glutamine, 50 μM 2-mercaptoethanol, and 20 ng/mL recombinant mouse GM-CSF (Peprotech). On day 3, 10 mL complete medium with GM-CSF was added, and on day 6, 10 mL medium with GM-CSF was exchanged from each plate. The floating cells were harvested on day 7.

**Flow cytometry.** Mouse lungs were transferred into a 6-well dish, and tissue was teased apart with forceps. Then the tissues were digested at 37°C for 30 minutes in RPMI containing 428 U/mL Collagenase IV.
BAL fluid were stained with Zombie Aqua Fixable Viability Kit (BioLegend), Ly-6G–BV510, and CD11c–PE-Cy7 (overnight, 4°C). Lymphocytes in BAL fluid cells, collected BAL cells were washed and immediately fixed/permeabilized with a BD Cytofix/Cytoperm kit. Cells were blocked with 1% anti–mouse CD16/CD32 and stained with permeabilized cell surface markers: CD45-PerCP/Cy5.5, CD24-BV421, CD11c–PE-Cy7, CD11b–APC-Cy7, and Siglec F–PE. Cells were washed and fixed/permeabilized with a BD Cytofix/Cytoperm kit. Cells were blocked (15 minutes, 4°C) with 1% anti–mouse CD16/CD32 and stained with cell surface markers: CD3–NK1.1+, CD3 +NK1.1–, CD3 +NK1.1+, and CD3–NK1.1– (NK1.1–APC, CD3–PE) cells in BAL fluid cells, CCR2 inflammatory monocytes (CCR2-PE+, Ly6C-APC+, CD45-BV421+), T cells (CD45-BV421+, CD3–PE-Cy7 or CD3–APC-Cy7), and CD49b–PE-Cy7. Analyses were performed on a FACSCanto flow cytometer or FACS Fortessa analyzer (BD Biosciences). Data were analyzed with FlowJo (TreeStar). Antibodies used are listed in Supplemental Table 3.

Real-time qPCR of mRNA transcripts. Right lung, spleen, lymph node, BM, and BAL cells were collected at time of euthanasia and snap-frozen. AMs were purified from BAL of Df- and IL-12-treated mice by positive CD11c MACS selection (Miltenyi Biotec), according to the manufacturer’s protocol. AM purity was checked by staining with Diff-Quick and was confirmed to be greater than 95%. Sorting of CD3 NK1.1+, CD3–NK1.1, CD3–NK1.1, and CD3 NK1.1 (NK1.1–APC, CD3–PE) cells in BAL fluid cells, CCR2 inflammatory monocytes (CCR2-PE+, Ly6C-APC+, CD45-BV421+), T cells (CD45-BV421+, CD3–PE+, CD90-APC+), and ILCs (CD45-BV421+, Lin-FITC–, CD90–APC–) in dispersed lungs was performed using a FACSDiva 8.0.1 cell sorter (BD Biosciences). Total RNA was isolated from tissue homogenate, BAL cells, BMDCs, BMMs, and sorted cells with QIAzol reagent (QIAGEN). For a few cells, glycogen (RNA grade, Thermo Fisher Scientific) was used as an inert carrier to increase nucleic acid recovery from alcohol precipitation. Total RNA was converted to cDNA using an RT² First Strand Kit (QIAGEN). The expression of each gene was examined using an RT² SYBR Green qPCR Master Mix (QIAGEN). Expression levels of transcripts were normalized to the expression of Gapdh. All primer sequences are listed in Supplemental Table 4.

UDP bioassy and calcium flux. We transfected 1321N1 cells (gift from Y. Kanaoka, Brigham and Women’s Hospital) with the mouse P2ry6 expression plasmid (pEFI-mouse P2ry6) using Lipofectamine 2000 (Invitrogen). Stably transfected cells were selected with G418 for more than 21 days. Mouse 1321N1 cells stably expressing P2y6, parental cells, BMMs, and the macrophage cell line RAW 264.7 (ATCC) were cultured in DMEM containing 10% heat-inactivated FBS, 100 μg/mL penicillin, 100 μg/mL streptomycin, 2 mM l-glutamine, and 10 ng/mL recombinant mouse GM-CSF and 5 ng/mL recombinant mouse M-CSF (both from R&D System). The next day, non- or weakly adherent cells were transferred to a new petri dish and cultured. After 2 days, 10 mL medium with GM-CSF and M-CSF was exchanged from each plate; and 5 days after seeding, adherent BM-derived macrophages were used for AM transplantation. Two million BM-derived macrophages in a total volume of 50 μL PBS were administered into the lungs of +/+ mice and P2ry6/C−/− mice using a relatively noninvasive endotracheal instillation method described previously (35). The next day, mice were administered with tamoxifen for 5 consecutive days and 10 days after the last tamoxifen treatment, allergic pulmonary inflammation was induced by i.n. instillation of Df, as described above.

Pulmonary macrophage transplantation. AM transplantation was performed as described previously (35). BM cells obtained from tamoxifen-treated +/+ and P2ry6/C−/− mice on day 10 were harvested, and this suspension (-27 x 10⁶ cells/10 mL) was cultured in DMEM medium supplemented with 10% heat-inactivated FBS, 100 U/mL penicillin, 100 μg/mL streptomycin, 2 mM l-glutamine, and 10 ng/mL recombinant mouse GM-CSF and 5 ng/mL recombinant mouse M-CSF (both from R&D System). The next day, non- or weakly adherent cells were transferred to a new petri dish and cultured. After 2 days, 10 mL medium with GM-CSF and M-CSF was exchanged from each plate; and 5 days after seeding, adherent BM-derived macrophages were used for AM transplantation. Two million BM-derived macrophages in a total volume of 50 μL PBS were administered into the lungs of +/+ mice and P2ry6/C−/− mice using a relatively noninvasive endotracheal instillation method described previously (35). The next day, mice were administered with tamoxifen for 5 consecutive days and 10 days after the last tamoxifen treatment, allergic pulmonary inflammation was induced by i.n. instillation of Df, as described above.

Statistics. All values are presented as mean ± SEM. Figures were produced and statistics analyzed using GraphPad Prism 6 (GraphPad Software). When data were not normally distributed, the Mann-Whitney U test was used for comparisons of 2 groups. A 2-tailed unpaired t test for equal variance and Welch’s correction for unequal variance were used when comparing 2 groups. Differences among multiple groups were
assessed using 1-way or 2-way ANOVA with Tukey’s multiple-comparisons test and nonparametric Kruskal-Wallis test followed by Dunn’s post hoc test. P values less than 0.05 were considered statistically significant.

Study approval. All experiments described herein were approved by the Brigham and Women’s Hospital IACUC and the Dana-Farber Cancer Institute Animal Care and Utilization Committee.

Author contributions
JN designed and performed the experiments and analyzed the data. LBF and TK performed experiments. JN developed the UDP assay. BB provided advice on the macrophage-related experiments. HC and JL provided technical assistance. MI, AZ, EYK, PJB provided intellectual and strategic input. JAB oversaw the project and experiments, and composed the manuscript with JN.

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