Polymorphonuclear neutrophil (PMN) activation is pivotal in acute inflammation and injury from reperfusion. To elucidate components controlling PMNs in vivo, we prepared novel transgenic mice with the human leukotriene (LT) B₄ receptor (BLTR) for functional characterization. Overexpression of BLTR in leukocytes dramatically increased PMN trafficking to skin microabscesses and lungs after ischemia-reperfusion, whereas mice deficient in 5-lipoxygenase (5-LO) showed diminished PMN accumulation in reperfused lungs. Hence, both BLTR expression and LT biosynthesis are critical for PMN infiltration in reperfusion-initiated second-organ injury. Also, in BLTR transgenic mice, 5-LO expression and product formation were selectively increased in exudates, demonstrating that receptor overexpression amplifies proinflammatory circuits. Endogenous lipoxin (LX) A₄ was produced in ischemic lungs and elevated by reperfusion. Because LXA₄ and aspirin-triggered 15-epimeric LXA₄ (ATL) selectively regulate leukocyte responses, they were tested in BLTR transgenic mice. Despite excessive PMN recruitment in BLTR transgenic mice, intravenous injection of ATL sharply diminished reperfusion-initiated PMN trafficking to remote organs, and topical application of LX was protective in acute dermal inflammation. These results demonstrate a direct role for BLTR with positive feedback, involving BLTR and 5-LO signaling in controlling PMNs. Moreover, LXA₄ and ATL counter BLTR-amplified networks, revealing a novel protective role for LX and ATL in stress responses that has applications in perioperative medicine.
Polymorphonuclear neutrophil (PMN) recruitment and sequestration to sites of inflammation and second-organ injury is initiated by proinflammatory mediators, among which leukotriene (LT) B4 is considered to be very important (1, 2). Recently, the LTB4 receptor (BLTR) was also reported to serve as a coreceptor for HIV-1 entry (3), which further emphasizes the crucial role of this system in host defense. Aspirin is widely used for its anti-inflammatory and analgesic properties and in several newly identified therapeutic actions, including preventing cardiovascular diseases and decreasing the incidence of lung, colon, and breast cancers (4), which raises the importance of obtaining complete knowledge of aspirin’s mechanism of actions. In addition to inhibiting prostanoid biosynthesis, administration of aspirin also triggers the endogenous transcellular production of 15-epimeric or 15R-LXA4, termed aspirin-triggered LXA4 (ATL), which appears to partly mediate some of aspirin’s therapeutic impact (5) and which is generated in vivo (6). LXA4 controls leukocyte responses via its own specific G protein–coupled receptor, denoted ALXR (5, 7). As with other local mediators, lipoxins (LXs) are rapidly generated, evoke responses, and are regulated by further metabolism (5). Thus, to explore potential anti-inflammatory actions of LXs and regulation of BLTR in vivo, LXA4 and ATL stable analogues were designed that resist rapid metabolic conversion and are ALXR agonists (5, 7). These designer compounds, modeled in view of the biochemical events initiated by aspirin, appear to act through similar mechanisms as endogenous LX to regulate leukocytes without aspirin’s unwanted side effects.

Recent cloning of BLTR in humans (8) and mice (9, 10) enabled the elucidation of this signaling pathway at the receptor and gene levels. Each of these BLTRs shows homology with ALXR and the chemoattractant

Leukotriene B4 receptor transgenic mice reveal novel protective roles for lipoxins and aspirin-triggered lipoxins in reperfusion

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peptide and chemokine family of receptors (~30%) exemplified by FMLP, C5a, and IL-8 receptors (11), but does not show extensive homology with the prostanoid receptors, providing further evidence that the origin of receptors for LX and LT is distinct from that for prostanoids. Along these lines, we developed transgenic (TG) mice with human BLTR (hBLTR). By using these novel hBLTR TG mice in conjunction with mice deficient in 5-lipoxygenase (5-LO) (12), we demonstrate that LXA4 and ATL regulate PMN-initiated second-organ injury in reperfusion; we also uncover a novel positive circuit in 5-LO signaling.

**Methods**

**Cloning and functional expression of hBLTR.** Total RNA was isolated from retinoic acid differentiated HL-60 cells (13) using Trizol reagent (GIBCO BRL, Grand Island, New York, USA), and was reverse transcribed for 30 minutes at 50°C followed by 40 cycles of PCR using Vent DNA polymerase (New England Biolabs Inc., Beverly, Massachusetts, USA) (98°C for 1 minute, 60°C for 1 minute, and 72°C for 2 minutes) with specific primers for hBLTR (BLTR-N: 5¢-GCCGGATCCATGAACACTACATCTTCTGCA-3¢; and BLTR-C: 5¢-GCCCTCGAGCTAGTTCAGTTCGTTTAACTT-3¢) (Figure 1a). The hBLTR cDNA was inserted into pGL vector (Promega Corp., Madison, Wisconsin, USA) at the HindIII-PpuMI site downstream of human CD11b promoter (Figure 1a), which was subcloned from a CD11b genomic clone (a gift from D.G. Tenen, Beth Israel Deaconess Medical Center, Boston, Massachusetts, USA), as described (14). This CD11b promoter fragment extended from bp –1704 to +83, and therefore included 83 bp of the untranslated region extending up to the ATG start codon. Human embryonic kidney (HEK293) cells were transfected with this pGL-CD11b-hBLTR construct using SuperFect reagent (QIAGEN Inc., Valencia, California, USA). Ligand binding and intracellular calcium release were carried out as described (refs. 8 and 15, respectively). For stable transfection, hBLTR cDNA was inserted into pcDNA3 vector (Invitrogen Corp., Carlsbad, California, USA) carrying a neomycin-resistant gene. Human ALXR cDNA was inserted into pcDNA6 (Invitrogen Corp.) carrying a different selection marker (blasticidin-resistant gene). HEK293 cells were transfected with the pGL-CD11b-hBLTR construct using SuperFect reagent (QIAGEN Inc., Valencia, California, USA). Ligand binding and intracellular calcium release were carried out as described (refs. 8 and 15, respectively). For stable transfection, hBLTR cDNA was inserted into pcDNA3 vector (Invitrogen Corp., Carlsbad, California, USA) carrying a neomycin-resistant gene. Human ALXR cDNA was inserted into pcDNA6 (Invitrogen Corp.) carrying a different selection marker (blasticidin-resistant gene). HEK293 cells were transfected with both constructs and selected with both neomycin and blasticidin. Cytosensor microphysiometry analysis of extracellular acidification rate (EAR) was carried out as described (16).

**Preparation and identification of hBLTR TG mice.** An approximately 3.4-kb hBLTR transgene was obtained by digestion of the plasmid (pGL-CD11b-hBLTR) with NotI and BamHI (Figure 1a) and was purified by Elutip (Schleicher & Schuell, Keene, New Hampshire, USA). Purified transgene construct (2 ng/µL) was injected into the male pronucleus of FVB homozygote oocytes that had been collected after hyperovulatory stimulation of donor females. The injected embryos were implanted in pseudopregnant Swiss females. Potential TG founder mice were screened by PCR analysis using genomic DNA isolated from mouse whole blood. Briefly, 15 µL of blood was collected from each mouse and resuspended in 250 µL of 10 mM Tris-HCl (pH 7.8) containing 1 mM EDTA and 0.5% NP-40. Cellular materials were obtained by centrifugation at approximately 9,500 g for 5 minutes and were resuspended in 40 µL of 0.1% Triton X-100 and 10 µL of 0.4 N NaOH to partially denature cell membranes and proteins. They were then heated at 95°C for 5 minutes, cooled on ice, and neutralized with 10 µL of 1 M Tris (pH 7.5), and each sample (1 µL) was used for 25 µL PCR. Hot-start PCR (98°C for 5 minutes before adding the Vent polymerase) was performed with 40 cycles of amplification (98°C for 1 minute, 60°C for 1 minute, and 72°C for 2 minutes) using primer sets (a) specific for hBLTR: BLTR-N and BLTR-C, which amplify the entire coding region of hBLTR; or (b) CD11b5’ (5’-GGCACCCTTTGGATAGTGTTTGTA-3’) and
CD11b3'-BLTR (5'-GTAGTGTTCATGATGTAGCGGTAGGTG-3'), which amplify 330 bp of the human CD11b promoter extending up to the bp +11 of hBLTR coding region (see Figure 1a). The positive founder obtained was bred with mice of the same strain. Potential positive litters (F1) from the TG founder were screened by PCR analysis as already described here. To verify BLTR and 5-LO expression, peritoneal leukocytes from mice with casein-induced peritonitis (2% casein for 4 hours) were collected as described (6), and total RNA was isolated. RT-PCR was performed using essentially the same conditions as described here and with primers that amplify both human and mouse BLTR: BLTR-III: 5'-TACGCCAGCGTCCTGCTT-3'; and BLTR-VII: 5'-GCTGCTCAGGAAGGCGAG-3' (Figure 2a). For amplifying 5-LO, mouse-specific sense (5'-ATCAGGACGTTCACGGCAGG-3') and antisense (5'-CCAGGAAAGCAGCTCAGGT-3') primers were used. Relative intensities of RT-PCR products were quantitated and normalized by GAPDH message levels using the NIH Image program (National Institutes of Health, Bethesda, Maryland, USA).

Ear skin inflammation model. hBLTR TG and nontransgenic (nTG) littermates were anesthetized, and 16-phenoxy-LXA4 [15(R)-16-phenoxy-LXA4 methyl ester] was prepared by total synthesis by N.A. Petasis and colleagues (Department of Chemistry, University of Southern California, Los Angeles, California, USA) following procedures used in ref. 7. Each LX (~26 nmol/10 μL acetone) was applied to the inner side of right ears. Acetone alone was applied to the left ear for a vehicle control. Five minutes later, LTB4 (5S,12R-dihydroxy-6,8,10,14-eicosatetraenoic acid; 1 μg) in 10 μL acetone was applied to both ears. Ear skin punch biopsies were collected for leukocyte myeloperoxidase (MPO) activity (7).

Hindlimb ischemia-reperfusion–induced second-organ injury. Mice were anesthetized, and ATL analogue 15-epi-16-(para-fluoro)-phenoxy-LXA4 [15(S)-16-(para-fluoro)-phenoxy-LXA4 methyl ester] as in ref. 7 or vehicle was administered intravenously to the TG mice. Approximately 5 minutes later, a tourniquet was placed proximally around each hindlimb and secured with a metal clamp (17). Vascular occlusion was verified by engorgement and discoloration of the feet. After 3 hours of ischemia, the tourniquets were removed and followed by 3 hours of reperfusion. The 5-LO+/+ and 5-LO–/– mice (The Jackson Laboratory, Bar Harbor, Maine, USA) (18), as well as hBLTR TG mice, were then euthanized with an overdose of pentobarbital by intraperitoneal injection, in accordance with the Harvard Medical Area Standing Committee on Animals (protocol no. 02570-R98). The left lungs were harvested, homogenized in 1.5 mL of potassium phosphate buffer (pH 7.4), and then centrifuged at approximately 15,000 g (5 minutes). The precipitates were extracted for leukocyte MPO analysis (7), and the supernatants were stored (~80°C) for eicosanoid analysis.

Liquid chromatography-tandem mass spectrometry. Murine peritoneal exudates were obtained from casein-induced peritonitis and incubated with A23187 (5 μM) at 37°C for 30 minutes. The samples were prepared by solid-phase extraction (7), with 10 ng of d4-LTB4 (6,7,14,15-tetradecuterium [d4]-LTB4) added to each as internal standard to calculate recovery. Liquid chromatography-tandem mass spectrometry (LC/MS/MS) was performed with an LCQ (Finnigan Corp., San Jose, California, USA) ion trap mass spectrometer system.

Figure 2

Overexpression of hBLTR selectively enhanced PMN infiltration in hBLTR TG mice. (a) BLTR-III (sense; right-facing arrow) and BLTR-VII (antisense; left-facing arrow) primers used for amplifying human and mouse BLTR messages are indicated. (b) BLTR (lanes 2 and 4) and GAPDH (lanes 5 and 6) expression in nTG (lanes 2 and 5) and hBLTR TG (lanes 4 and 6) mice was analyzed by RT-PCR. RT-PCR controls (minus RT) are shown in lane 1 (nTG) and lane 3 (TG). The expected molecular size for BLTR at approximately 0.5 kb is indicated by an arrow. The relative intensities of BLTR messages normalized by GAPDH (BLTR/GAPDH) are reported. (c) LTB4 (1 μg), arachidonic acid (10 μg), or PMA (100 ng) was applied topically to the right ears of nTG and TG mice. Left ears received vehicle alone (acetone). Ear skin biopsy samples were taken for MPO analysis. Data were expressed as ratio (TG/nTG) of dermal leukocyte infiltration and (inset) fold increase above vehicle control (*P < 0.01; n = 7 for nTG and n = 10 for TG-hBLTR).
Topical application of LXA4 inhibits PMN infiltration in hBLTR TG mice. The LXA4 stable analogue (10 μg, 26 nmol) 16-phenoxy-LXA4 (inset: LXA4 analogue template R2= phenoxy) was applied topically to right ears of nTG and hBLTR TG mice (hatched bars). Left ears received vehicle alone (filled bars). LTB4 (1 μg) was then applied to both ears of each mouse 5 minutes later. Skin biopsies were obtained as just described. Values represent total PMN infiltration into ear skin (representative data from 3 experiments).

Results
TG mice were prepared for these experiments with hBLTR cloned in our laboratory from retinoic acid differentiated HL-60 cells that display LTB4-specific binding (13), using reported sequences (8). This BLTR transgene was placed in control of a human CD11b promoter (Figure 1a) capable of driving high levels of transgene expression in mature murine myeloid cells (14). To verify our clone, HEK293 cells were transfected with hBLTR cDNA (pGL-CD11b-hBLTR). These cells displayed both specific [3H]LTB4 binding (data not shown) and ligand-stimulated mobilization of intracellular calcium (Figure 1b), which together qualified its use for preparing hBLTR TG mice. Genomic DNA from founder mice was isolated and screened by PCR using primer sets (a) denoted as BLTR-N and BLTR-C (Figure 1a), which were specific to the hBLTR; or (b) CD11b5¢ and CD11b3¢-BLTR, which amplified the nucleotide segment corresponding to the CD11b promoter-hBLTR junction that is unique to our transgene. Both sets of primers yielded PCR products with genomic DNA from 1 potential founder of 12, whereas no apparent bands were observed with genomic DNA from nTG mice of the same strain (Figure 1c). Primers that amplified the entire coding region of hBLTR (e.g., BLTR-N and BLTR-C) yielded more prominent PCR signals, and thus were selected for future screening of the F1 generations. Twenty-four positive F1 mice were obtained (out of 36), with representative results shown in Figure 1d.

Peritoneal leukocyte exudates (from casein-induced peritonitis) from positive hBLTR TG mice were collected, and displayed an approximately 7-fold increase in BLTR message levels compared with nTG littermates (Figure 2b, lanes 2 and 4) that were age and strain matched (relative intensities of BLTR messages were ~0.09 for nTG and ~0.60 for TG mice, normalized by GAPDH as shown in Figure 2b). Primers BLTR-III and BLTR-VII (Figure 2a), which amplified both human and mouse BLTR, were used in RT-PCR to compare the message levels of endogenous mouse BLTR with those of exogenous hBLTR transgene. No apparent bands were observed in RT-PCR controls (minus RT; Figure 2b, lanes 1 and 3), indicating no genomic DNA contamination in our total RNA preparations. Thus, our results showed that the hBLTR transgene was indeed expressed and total BLTR expression was dramatically increased.

Of interest, these hBLTR TG mice appeared healthy and without apparent gross pathological findings in the absence of a specific challenge (see below).

LTB4 applied topically to mouse ear skin induces PMN influx and microabscess in an acute dermal inflammatory response (7). Our TG mice showed a pronounced increase (7.4 ± 0.9-fold) in PMN infiltration into skin (Figure 2c, inset), whereas the nTG mice response was increased 3.2 ± 0.5-fold compared with the vehicle control. In sharp contrast, PMN infiltration into ear skin with either the native precursor of LTB4, namely arachidonic acid (C20:4), or the protein kinase C direct agonist PMA was not augmented in hBLTR TG mice (Figure 2c). Arachidonic acid (10 μg/ear) produced 3.0- and 2.5-fold increases in PMN infiltration in nTG and TG mice, respectively. Application of PMA (100 ng/ear) evoked 12.0- and 11.5-fold increases in nTG and TG mice, respectively. It should be noted that
both C20:4 and PMA were added at levels that did not destroy tissue architecture of the ear skin. These findings indicate that hBLTR was functionally expressed and directly correlated with profound amplification of LTB4-initiated PMN microabscess formation in skin.

To determine whether LXA₄ directly interacts with hBLTR. To address this, we examined specific binding and signaling in HEK293 cells expressing BLTR or BLTR together with ALXR as stable transfectants. LXA₄ did not compete with [³H]LTB₄ binding (Figure 4a), nor did LXA₄ or its analogues affect the LTB₄-evoked EAR (Figure 4b). Human BLTR couples to pertussis toxin–sensitive G proteins in HL-60 cells (8). Along these lines, Martin et al. (10) recently reported that transfected mouse BLTR functionally couples with Gₛ in *Xenopus laevis* and in Chinese hamster ovary cells. These signal transduction components of BLTR demonstrated in in vitro systems may also be involved in vivo, and are likely to mediate LTB₄'s proinflammatory actions. It is thus likely that ALXR activation inhibits signal transduction components that occur downstream from LTB₄-BLTR recognition.

Ischemia-reperfusion is an event with major clinical importance. In humans, surgically based clamping procedures are well known to lead to aberrant PMN activation, giving rise to second-organ injury that contributes to longer hospitalization (19). To evaluate the contribution of both BLTR and LT formation in reperfusion injury, we used a hindlimb tourniquet model of second-organ injury (17) with both 5-LO–deficient and hBLTR TG mice (Figure 5, a and b).

5-LO+/+ mice demonstrated twice as much leukocyte accumulation in lungs compared with 5-LO–/– mice (10.5 × 10⁵ vs. 5.4 × 10⁵ PMNs greater than that of control animals) (Figure 5a). These 5-LO–deficient mice are unable to synthesize LTs (18). Thus, our data indicate that 5-LO–derived products (e.g., LTB₄; see Figure 7, inset) are major mediators of PMN recruitment into lungs after hindlimb ischemia-reperfusion. Pharmacological evidence also implicated LTB₄ as a key component in ischemia-reperfusion–induced second-organ injury in mice (17) and rats (20). Results with PMN recruitment in 5LO⁺/⁺ mice also point to the contribution of other mediators, such as C5a, IL-8, and platelet-activating factor (17), as PMN recruitment occurs in the absence of endogenous LTB₄ formation (Figure 5a). The hBLTR TG mice also showed dramatic increases in PMN infiltration into lungs after reperfusion, compared with nTG mice (Figure 5b). When taken together, results from these genetically manipulated mice (namely, 5-LO–deficient and hBLTR TG mice) demonstrate that application of the LXA₄ stable analogue was clearly able to block PMN infiltration (Figure 3). The percent inhibition in hBLTR TG versus nTG mice (~45% vs. ~37% inhibition) did not prove to be significantly different. But in magnitude, the LXA₄ stable analogue inhibited infiltration of approximately 3 times as many PMNs in hBLTR TG mice as in nTG mice (Figure 3), which was evident when leukocyte MPO activities from these punch biopsies were calibrated and converted to total number of PMNs. No apparent increases in ALXR message levels were observed (data not shown). Hence, LXA₄ and ALXR constitute a potent system for BLTR regulation in vivo.

Figure 4
Recombinant BLTR recognition and signaling. (a) Displacement of specific [³H]LTB₄ binding by LTB₄ (filled squares; structure shown), LXA₄ (open diamonds), or LXB₄ (open circles) was determined in HEK293 cells stably transfected with BLTR. (b) EARs in response to LXA₄ and LTB₄ were analyzed by cytosensor microphysiometry in HEK293 cells with BLTR and ALXR (stable transfectants). Cells were exposed to LXA₄ (1 μM; open circles) or media alone (filled circles) 20 minutes before exposure to LTB₄ (1 μM). Values are expressed as EAR (µV/s) normalized to baseline (100%), and ligand additions are indicated by arrows.
both the enhanced appearance of BLTR and the LT formation are important events in ischemia-reperfusion–induced second-organ injury, and therefore are potential therapeutic targets for further consideration in perioperative treatment.

In view of the ability of LXA₄ to modulate PMN-directed responses (5), it was of interest to determine whether LXA₄ is generated during ischemia-reperfusion and whether it has an impact in the TG amplified BLTR signaling. Hindlimb ischemic mice showed increased amounts of endogenous LXA₄ in lungs, and tourniquet release (reperfusion) further increased these levels (Figure 6a). Therefore, an analogue of ATL (Figure 6b, inset) was administered via intravenous injection in hBLTR TG mice before hindlimb ischemia-reperfusion. The administration of ATL significantly

Figure 5
S-LO pathway and BLTR are major determinants of PMN infiltration into lungs after hindlimb ischemia-reperfusion. S-LO−/− and S-LO+/+ (a) and hBLTR TG and nTG (b) mice were subjected to hindlimb ischemia-reperfusion. The left lungs were collected. MPO activities were determined, and data are expressed as (a) increase in lung PMN infiltrates (×10⁵) compared with control animals (ischemia alone) from 2 separate experiments, and (b) total PMN infiltration into lungs after ischemia-reperfusion. Total PMN numbers obtained from hBLTR TG and nTG mice were significantly different (*P = 0.02; n = 3).

Figure 6
ATL inhibits PMN infiltration into lungs after hindlimb ischemia-reperfusion. (a) BALB/c mice were subjected to ischemia-reperfusion (~0.8 ng; filled bar) or ischemia alone (~0.5 ng; hatched bar). After removal of hindlimb tourniquet, left lungs were collected with or without reperfusion. LXA₄ present within each lung tissue was quantitated by ELISA (Neogen Corp., Lexington, Kentucky, USA) or LC/MS/MS. (b) hBLTR TG mice were injected with vehicle (hatched bar) or the ATL analogue (10 µg; filled bar) [inset: ATL analogue template R₂ = (para-fluoro)-phenoxy]. Leukocyte MPO values were obtained from left lung of each mouse after ischemia-reperfusion and were expressed as percent inhibition of lung PMN infiltration. Values from mice that received ATL analogue (~7 × 10⁵ PMN) versus vehicle alone (~12 × 10⁵ PMN) were significantly different (*P = 0.04; n = 3).
diminished PMN infiltration into lungs when compared with hBLTR TG mice injected with vehicle alone (Figure 6b). Thus, LXA₄ formation within ischemic tissue, and its elevation by reperfusion, might represent an endogenous compensatory or protective role to limit PMN trafficking and PMN-mediated damage. This is supported by results showing that intravascular injection of the ALXR agonist (e.g., ATL stable analogues that display increased biostability compared with the endogenous LXA₄; compare refs. 5 and 7) before ischemia-reperfusion attenuates BLTR-driven PMN infiltration. In addition, because ATL inhibited PMN infiltration, which is approximately the reciprocal level (40–60% change) observed in the S-LO⁻/⁻ and BLTR TG mice (Figure 5, a and b; Figure 6b), it emphasizes the importance of the LXA₄-LTB₄ regulatory system in these genetically defined mice.

Using LC/MS/MS–based analysis, we found that in hBLTR TG mice, 5S-HETE (Figure 7a) (a product of 5-LO; Figure 7b, inset) was selectively and dramatically elevated in peritoneal inflammatory exudates compared with nTG mice (Figure 7b). In contrast, significant differences were not observed in the amount of other mono-HETEs from the major lipoxygenases, including 15S-HETE, 12S-HETE (Figure 7b), or LTB₄ (data not shown). It is noteworthy that 5S-HETE is also a PMN chemoattractant, albeit less potent than LTB₄ (21). Expression of 5-LO transcript in peritoneal leukocytes was approximately 3.5-fold higher in hBLTR TG mice than in nTG mice. Relative intensities of 5-LO messages were approximately 0.26 and approximately 0.91 for nTG and hBLTR TG mice, respectively (normalized by GAPDH message levels; Figure 7c). The observed increase in 5S-HETE, rather than LTB₄, in hBLTR TG mice may reflect enhanced LTB₄ utilization and clearance (i.e., further metabolism) in this model. We noted no apparent increases in cycloxygenase-2 message levels or striking elevations in cyclooxygenase products (e.g., prostaglandin E₂) (data not shown). Thus, the overexpression of BLTR specifically enhances a proinflammatory network by acting at the level of 5-LO gene transcription that is evident in vivo (Figure 7c), suggesting that this positive feedback within the 5-LO pathway could be operative in a wide range of disease states.

Discussion
To our knowledge, our findings provide the first demonstration that overexpression of BLTR profoundly amplifies PMN recruitment, function, and 5-LO signaling in murine models of acute skin inflammation, peritonitis, and reperfusion-initiated second-organ injury. In addition, this is the first direct evidence, to our knowledge, for an in vivo role of BLTR expression in regulating PMN activation and upregulation of the 5-LO pathway. Together, these results emphasize the impact of receptor expression as an important regulatory step in host responses, as well as in ischemia-reperfusion, and are consistent with the regulation of ALXR (22) and BLTR (9) expression by cytokines that are linked to the control of human immune functions. Moreover, these results open new avenues for the utility of counter-regulatory signals, namely ALXR agonists and metabolically stable ATL mimetics, to selectively regulate inflammatory diseases and reperfusion-associated injury events that are characterized and exacerbated by excessive PMN infiltration and activation.
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