PD_{n-3} DPA Pathway Regulates Human Monocyte Differentiation and Macrophage Function

Graphical Abstract

Highlights

- PD_{n-3} DPA regulates human monocyte-derived macrophage differentiation and function
- Evidence for the formation and complete stereochemistry of 16S,17S-ePD_{n-3} DPA
- EPHX2 converts 16S,17S-ePD_{n-3} DPA to PD_{2n-3} DPA in human monocytes

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In Brief

Pistorius et al., found that a recently uncovered family of bioactive mediators termed PD_{n-3} DPA controls human macrophage phenotype and function during their differentiation from monocytes. The authors also established the biosynthetic pathway for this family of mediators thereby providing leads into mechanisms that control macrophage responses.

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PD$_n$-3 DPA Pathway Regulates Human Monocyte Differentiation and Macrophage Function

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SUMMARY

Macrophages are central in orchestrating the clearance of apoptotic cells and cellular debris during inflammation, with the mechanism(s) regulating this process remaining of interest. Herein, we found that the n-3 docosapentaenoic acid-derived protectin (PD$_n$-3 DPA) biosynthetic pathway regulated the differentiation of human monocytes, altering macrophage phenotype, efferocytosis, and bacterial phagocytosis. Using lipid mediator profiling, human primary cells, where PD$_n$-3 DPA reduce human neutrophil-endothelial cell interactions, neutrophil chemotaxis, and increased macrophage phagocytosis. The complete stereochemistry of this epoxide was determined using stereoselective conversion of essential fatty acids. The first SPMs identified were the lipoxins (LX) that are produced from arachidonic acid and carry potent leukocyte directed actions (Serhan et al., 1984). The D-series resolvins (RvD) produced from docosahexaenoic acid (DHA) and the E-series resolvins (RvE) produced from eicosapentaenoic acid also display leukocyte-directed actions (Chiang and Serhan, 2017). In addition, RvEs regulate platelet response (Donn et al., 2008) and neutrophil apoptosis (El Kebir et al., 2012), while the RvDs regulate vascular smooth muscle cell migration, a key process in the development of intimal hyperplasia (Mottola et al., 2017). Recently, we found that n-3 docosapentaenoic acid (DPA) is also a substrate for conversion to a new series of pro-resolving mediators, which include the n-3 DPA-derived protectins (PD$_n$-3 DPA) (Dalli et al., 2013a). These mediators were initially identified in self-limited inflammatory exudates, with their production coinciding with monocyte/macrophage trafficking to the site. Human primary macrophages also produce these mediators. Of note, PD$_1$-3 DPA and PD$_2$-3 DPA carry potent leukocyte directed actions, reducing neutrophil recruitment as well as the production of inflammatory cytokines, including monocyte chemotactic protein-1, during sterile inflammation (Dalli et al., 2013a). These actions were also found with human primary cells, where PD$_n$-3 DPA reduce human neutrophil-endothelial cell interactions, neutrophil chemotaxis, and increased macrophage phagocytosis. The complete stereochemistry of the first member of this family of mediators, PD$_1$-3 DPA, was recently established as 10E,17S-dihydroxydocosahexaenoic acid (Aursnes et al., 2014). In humans, PD$_1$-3 DPA is upregulated in colon samples

INTRODUCTION

Macrophages are central in orchestrating host responses to tissue injury and infections (Fredman et al., 2014; Gordon and Mantovani, 2011; Pierdomenico et al., 2015). The biological actions of these cells are associated with their phenotype, where classically activated macrophages are central in the initiation and propagation of the inflammatory response, whereas alternatively activated macrophages orchestrate tissue repair and regeneration (Dalli and Serhan, 2016; Gordon and Mantovani, 2011; Morris et al., 2010). Investigations into the cellular mechanisms underlying these distinct biological functions demonstrate that each macrophage subtype produces a characteristic repertoire of bioactive molecules, including cytokines and lipid mediators (LMs) (Dalli and Serhan, 2016; Gordon and Mantovani, 2011). In this context, recent studies demonstrate that, while classically activated macrophages and alternatively activated macrophages produce both inflammation initiating eicosanoids and pro-resolving mediators, their LM phenotypes differ. Indeed, classically activated macrophages produce higher amounts of pro-inflammatory eicosanoids, whereas alternatively activated macrophages produce higher levels of the recently identified specialized pro-resolving mediators (SPMs) (Dalli and Serhan, 2012). Tissue macrophages arise during either embryonic development or are differentiated in situ from monocyte precursors (Kopf et al., 2015). Mechanisms that dictate monocyte differentiation to different macrophage phenotypes are of relevance in determining whether inflammation is perpetuated or resolved, thus permitting regain of function.

SPMs are produced via the stereoselective conversion of essential fatty acids. The first SPMs identified were the lipoxins (LX) that are produced from arachidonic acid and carry potent leukocyte directed actions (Serhan et al., 1984). The D-series resolvins (RvD) produced from docosahexaenoic acid (DHA) and the E-series resolvins (RvE) produced from eicosapentaenoic acid also display leukocyte-directed actions (Chiang and Serhan, 2017). In addition, RvEs regulate platelet response (Donn et al., 2008) and neutrophil apoptosis (El Kebir et al., 2012), while the RvDs regulate vascular smooth muscle cell migration, a key process in the development of intimal hyperplasia (Mottola et al., 2017). Recently, we found that n-3 docosapentaenoic acid (DPA) is also a substrate for conversion to a new series of pro-resolving mediators, which include the n-3 DPA-derived protectins (PD$_n$-3 DPA) (Dalli et al., 2013a). These mediators were initially identified in self-limited inflammatory exudates, with their production coinciding with monocyte/macrophage trafficking to the site. Human primary macrophages also produce these mediators. Of note, PD$_1$-3 DPA and PD$_2$-3 DPA carry potent leukocyte directed actions, reducing neutrophil recruitment as well as the production of inflammatory cytokines, including monocyte chemotactic protein-1, during sterile inflammation (Dalli et al., 2013a). These actions were also found with human primary cells, where PD$_n$-3 DPA reduce human neutrophil-endothelial cell interactions, neutrophil chemotaxis, and increased macrophage phagocytosis. The complete stereochemistry of the first member of this family of mediators, PD$_1$-3 DPA, was recently established as 10E,17S-dihydroxydocosahexaenoic acid (Aursnes et al., 2014). In humans, PD$_1$-3 DPA is upregulated in colon samples
Figure 1. Inhibiting 15-Lipoxygenase Activity Reduces PD_{1\rightarrow3} DPA Production Dysregulating Macrophage Phenotype and Function

(A) Human monocytes were incubated with M-CSF (20 ng/mL) and either an ALOX15 inhibitor or vehicle (37°C, 5% CO2). On day 7 incubations were quenched, lipid mediators were extracted, identified, and quantified using lipid mediator profiling (see the STAR Methods for details). Results are mean ± SEM. n = 6 donors. *p < 0.05.

(B) Human monocytes were isolated and incubated with GM-CSF (20 ng/mL), IFN-γ (20 ng/mL), and LPS (100 ng/mL) to produce M1 or M-CSF (20 ng/mL) and IL-4 (20 ng/mL) to obtain M2 cells, and the expression of ALOX15 and ALOX15B was evaluated during the differentiation time course using flow cytometry. Results are mean ± SEM n = 4–6 donors per interval.

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from IBD patients and displays potent tissue protective actions in mice during experimental colitis, reducing leukocyte-mediated tissue damage (Gobetti et al., 2017).

The PD$_{n-3}$ DPA biosynthetic pathway in human monocytes and macrophages is proposed to be initiated by 15-lipooxygenases (LOX), which convert n-3 DPA to an intermediate epoxide that in turn is enzymatically hydrolyzed to PD$_{1n-3}$ DPA (Aursnes et al., 2014) and PD$_{2n-3}$ DPA (16,17R-dihydroxy-7Z,10,13,14,19Z-docosapentaenoic acid) (Dalli et al., 2013a). Given the potent biological actions of PD$_{n-3}$ DPA, and the central role of the proposed epoxide intermediate in the formation of these pro-resolving mediators, it was deemed important to (1) obtain evidence for its formation and role in the PD$_{n-3}$ DPA biosynthesis, (2) establish the identity of enzyme(s) involved in the formation of this epoxide in human monocytes/macrophages, and (3) establish the biological actions of the PD$_{n-3}$ DPA pathway in monocyte differentiation and macrophage responses. Using acid alcohol trapping, recombinant enzymes and human primary cells we found that both ALOX15 and ALOX15B catalyze the formation of 16,17S-ePD$_{n-3}$ DPA. Using total organic synthesis we obtained stereochemically pure material, that, when incubated with primary human macrophages, was converted to both PD$_{1n-3}$ DPA and PD$_{2n-3}$ DPA. Furthermore, we found that the PD$_{n-3}$ DPA pathway was important in regulating macrophage phenotype and function.

RESULTS

Inhibition of 15-Lipooxygenase Activity in Human Monocytes Reduced Macrophage PD$_{n-3}$ DPA Metabolome and Altered Macrophage Function

To obtain evidence for the role of the PD$_{n-3}$ DPA biosynthetic pathway in regulating human macrophage function during monocyte differentiation, we incubated monocytes with an ALOX15 inhibitor and assessed the production of components within the PD$_{n-3}$ DPA pathway. In human primary cells, using liquid chromatography-tandem mass spectrometry (LC/MS-MS), we identified 17-HDPA, PD$_{1n-3}$ DPA, and PD$_{2n-3}$ DPA. In these incubations we also identified 10R/S,17S-dihDPA and 16R/S,17S-dihDPA, which correspond to the non-enzymatic products of the proposed epoxide intermediate in the PD$_{n-3}$ DPA pathway (Figure 1A). Of note, incubation of monocytes with an inhibitor of ALOX15 led to a significant reduction in all the components of the PD$_{n-3}$ DPA biosynthetic pathway, including the bioactive mediators PD$_{1n-3}$ DPA and PD$_{2n-3}$ DPA (Figure 1A; Table S1). The role of 15-lipooxygenases in PD$_{n-3}$ DPA biosynthesis was further corroborated using cells from mice deficient in the murine homolog of this enzyme (ALOX15$^{-/-}$), which demonstrated a significant reduction in all the components within the PD$_{n-3}$ DPA pathway (Figure S1; Table S2).

Human leukocytes express two ALOX15 subtypes, therefore we sought to determine the temporal regulation of these enzymes in circulating monocytes and during monocyte-macrophage differentiation. Flow cytometric analysis demonstrated that human circulating monocytes express both ALOX15 and ALOX15B. The expression of both enzyme isoforms was upregulated during monocyte-to-macrophage differentiation. Of note, ALOX15 expression was higher in M2 differentiated cells, whereas the expression of ALOX15B was higher in M1 cells (Figure 1B). We next investigated the contribution of each of these two enzymes to PD$_{n-3}$ DPA biosynthesis in human monocytes. Here we found that transfection of human monocytes with short hairpin RNA (shRNA) to ALOX15 reduced enzyme expression by ~50% (p < 0.05; n = 5 donors), and the concentrations of several components of the PD$_{n-3}$ DPA biosynthetic pathway, including PD$_{1n-3}$ DPA and PD$_{2n-3}$ DPA. Transfection of monocytes with shRNA to ALOX15B significantly reduced both enzyme expression (~50%; p < 0.05; n = 5 donors) and PD$_{n-3}$ DPA concentrations, albeit to a lesser extent than observed with shRNA to ALOX15 (Figure S2).

Given that PD$_{1n-3}$ DPA and PD$_{2n-3}$ DPA carry potent biological actions, we next questioned whether inhibition of this pathway during monocyte-to-macrophage differentiation influenced macrophage phenotype and function. We therefore assessed the expression of phagocytic receptors and adhesion molecules, which are linked with macrophage phenotype. Incubation of cells with ALOX15 inhibitor during monocyte differentiation led to the downregulation of several lineage markers, including CD206, CD163, and CD64, and a shift in macrophage phenotype (Figure 1C). This downregulation in phagocytic receptors was of functional consequence since inhibition of the PD$_{n-3}$ DPA biosynthetic pathway also significantly downregulated the ability of human macrophages to uptake apoptotic cells (Figure 1D), a key pro-resolving action (Chiang and Serhan, 2017; Dalli and Serhan, 2016). This alteration in macrophage phenotype and function was also observed in tissue-resident macrophages from ALOX15-deficient animals. In these mice the expression of lineage markers on splenic macrophages, small peritoneal macrophages, and large peritoneal macrophages was altered with a downregulation in the expression of several markers including transforming growth factor β and T-cell immunoglobulin and mucin domain containing protein (TIM)-4 (Figure 1E). This was associated with a decreased ability of peritoneal macrophages to uptake apoptotic cells in vivo (Figure 1F; Table S3).

Having established a role of the PD$_{n-3}$ DPA metabolome in regulating human macrophage function we next sought to obtain further evidence for the PD$_{n-3}$ DPA biosynthetic pathway. Given the central role that the proposed 16,17S-ePD$_{n-3}$ DPA plays in the biosynthesis of PD$_{n-3}$ DPA, we sought to gain evidence for the formation of the proposed epoxide in human monocytes. For

(C and D) Human monocytes were incubated with vehicle or ALOX15 inhibitor and then with M-CSF (20 ng/mL) for 7 days, and (C) expression of lineage markers was determined using fluorescently labeled antibodies and flow cytometry on day 7, and interrogated using OPLS-DA. n = 6 donors. (D) Phagocytosis of fluorescently labeled apoptotic cells investigated. Results for are mean ± SEM. n = 6 donors. *p < 0.05. (E) Peritoneal macrophages were harvested from wild-type (WT) and ALOX15$^{-/-}$ mice, and the expression of lineage markers on CD64$^+$ cells was determined using flow cytometry. Results were interrogated using OPLS-DA and are representative of n = 7 mice. (F) Fluorescently labeled apoptotic cells were administered to WT and ALOX15$^{-/-}$ mice via intraperitoneal injection. After 1 hr peritoneal cells were harvested, and phagocytosis of apoptotic cells by CD64$^+$ cells was evaluated using flow cytometry. Results are mean ± SEM. n = 7 mice per group. *p < 0.05.

Related to Figures S1 and S2 and Tables S1–S3.
this purpose we assessed the formation of the epoxide using acid alcohol trapping, given that allylic epoxides are known to be unstable in aqueous solutions (Radmark et al., 1980). Using LC/MS-MS we identified four peaks, the first two gave a retention time (TR) and MS-MS fragmentation spectra corresponding with 10-methoxy,17S-hydroxy-docosapentaenoic acid (Figure 2A), whereas, the TR and MS-MS spectra for peaks III and IV corresponded to 16-methoxy,17S-hydroxy-docosapentaenoic acid (Figure 2A).

Having obtained evidence for the formation of an epoxide intermediate by human monocytes we sought to determine whether ALOX15 enzymes were responsible for the formation of this intermediate. For this purpose we incubated human recombinant (hr) ALOX15 and hr-ALOX15B with n-3 DPA and assessed the acid alcohol trapping products using LC/MS-MS. This also gave four peaks that displayed essentially identical TR and MS-MS fragmentation spectra to the products obtained with primary human macrophages (Figures 2B and 2C). Thus, these results establish the role of human ALOX15 enzymes in the formation of 16,17S-ePDn-3 DPA.

Establishing the Complete Stereochemistry of 16S,17S-ePDn-3 DPA

Having obtained evidence for the formation of 16,17S-ePDn-3 DPA by human monocytes and the role of ALOX15 enzymes in its production, we next sought evidence for its role in the biosynthesis of PD1n-3 DPA and PD2n-3 DPA. For this purpose we obtained stereochimically pure material using total organic synthesis. The methyl ester of 16S,17S-ePDn-3 DPA was constructed via two key precursors (Figure 3A). These precursors were obtained using the following stereoselective reactions: a Katsuki-Sharpless epoxidation protocol (Kumar and Meshram, 2013) and one Z- and two E-selective Wittig reactions (Maryanoff and Reitz, 1989). The detailed description of the synthetic route...
is reported in Primdahl et al. (2017). The chemical purity and stereochemical integrity was validated using LC/MS-MS and nuclear magnetic resonance (NMR). The unambiguous assignment of the geometrical configurations of the olefins constituting the $E, E, Z$-triene moiety was performed using a $1H-1H$ COSY-45 NMR experiment (Figures 3B and S3). The following chemical shifts and coupling constants ($J$ values) were recorded for the epoxy methyl ester of 16$S$,17$S$-ePD$_{n-3}$ DPA: 12: 6.54 ppm, $J = 14.7$, 11.4 Hz; H14: 6.39 ppm, $J = 15.3$, 10.8 Hz; H13: 6.11 ppm, $J = 14.8$, 10.9 Hz; H11: 6.04 ppm, $J = 11.2$ Hz; H10 and H15: 5.48–5.32 ppm. Notably, H11 has a $3J$-coupling constant to H10 with a value of 11.2 Hz, which is consistent with the assigned $Z$-rather than an $E$-configured C10-C11 double bond. The UV chromophore of this compound was also consistent with that of an allylic epoxide conjugated to a triene double bond system, giving a $\lambda_{max}^{MeOH}$ of 280 nm with shoulders at 271 and 298 nm. These results confirm the stereochemistry of the synthetic epoxide as 16$S$,17$S$-epoxy-$7Z,10Z,12E,14E,19Z$-docosapentaenoic acid methyl ester.

We next assessed whether the physical properties of the synthetic material obtained after lenient hydrolysis of the epoxy methyl ester matched those of the biological material. Incubation of the synthetic material in acid alcohol gave four distinct peaks with products displaying essentially identical $T_R$ and MS-MS fragmentation spectra as those obtained with hr-ALOX15 enzymes and human monocytes (Figures 2 and S4).

We next investigated whether the synthetic material was substrate for conversion to PD$_{1n-3}$ DPA and PD$_{2n-3}$ DPA. To this end, human macrophages were incubated with 16$S$,17$S$-ePD$_{n-3}$ DPA and products were assessed using LC/MS-MS. In multiple reaction monitoring (MRM) we obtained two peaks. The first peak eluted with a $T_R$ of 13.8 min and gave characteristic fragments in the MS-MS that are consistent with PD$_{1n-3}$ DPA. These fragments included: $m/z$ 361, $m/z$ 183, and $m/z$ 155 (Figures 4A and 4B). The second peak eluted with a $T_R$ of 15.2 min and gave ions in the MS-MS that were characteristic of PD$_{2n-3}$ DPA including: $m/z$ 361, $m/z$ 263, and $m/z$ 233 (Figures 4A and 4C). Of note, incubation of 16$S$,17$S$-ePD$_{n-3}$ DPA in PBS only or with cells that had been previously kept at 100°C did not yield notable levels of these mediators (Figure 4D), indicating that the conversion of the epoxide to PD$_{1n-3}$ DPA and PD$_{2n-3}$ DPA was reliant on enzyme-mediated catalysis. In addition, these results establish the stereochemistry of the allylic epoxide intermediate as 16$S$,17$S$-epoxy-$7Z,10Z,12E,14E,19Z$-docosapentaenoic acid.

Epoxide Hydrolases Convert 16$S$,17$S$-ePD$_{n-3}$ DPA to PD$_{1n-3}$ DPA and PD$_{2n-3}$ DPA

Having established a role for enzymes in catalyzing the conversion of 16$S$,17$S$-ePD$_{n-3}$ DPA to PD$_{1n-3}$ DPA and PD$_{2n-3}$ DPA we next sought to establish the identity of these enzymes. Given the role that epoxide hydrolases play in converting allylic epoxides to bioactive mediators we assessed whether
this class of enzymes was also involved in catalyzing the conversion of 16S,17S-ePDn-3 DPA in human monocytes. Incubation of human macrophages with the epoxide hydrolase inhibitor AUDA led to a reduction in the concentrations of both PD1n-3 DPA and PD2n-3 DPA (Figure 5A).
Figure 6. EPHX2 Converts 16S,17S-ePD_{n-3}DPA to PD_{2n-3}DPA in Human Monocytes and Macrophages
(A) Human monocytes were isolated and differentiated using GM-CSF (20 ng/mL), IFN-γ (20 ng/mL), and LPS (100 ng/mL) to produce M1 or M-CSF (20 ng/mL) and IL-4 (20 ng/mL) to obtain M2 cells and the expression of EPHX2 during the differentiation time course was evaluated using flow cytometry. Results are mean ± SEM. n = 4–6 donors per interval.
(B) Human monocytes were transfected with shRNA to EPHX2 or CT shRNA (see the STAR Methods for details), cells were incubated for 10 hr at 37°C, then with E. coli for 45 min, and PD_{2n-3}DPA concentrations evaluated using LM profiling. Results are mean ± SEM. n = 4 donors. *p < 0.05.
(C) 16S,17S-ePD_{n-3}DPA (10 nM) was incubated with hrEPHX2 (0.2 μM; Tris buffer). Incubations were quenched using ice-cold methanol and products identified using lipid mediator profiling. Left panel: MRM chromatogram m/z 361 > 233. Center panel: MS-MS spectrum employed in the identification of PD_{2n-3}DPA. Right panel: PD_{2n-3}DPA concentrations. Results are representative of n = 4 independent experiments. *p < 0.01 versus EPHX2 incubations.

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16S,17S-ePDn-3 DPA gave four peaks that were identified as the non-enzymatic hydrolysis products of 16S,17S-epoxy-PDOn-3 DPA. The TR and MS-MS spectra of peaks I and II corresponded to 10R/S,17S-dihydroxy-7Z,11E,13E,15E,19Z-docosapentaenoic acid, whereas peaks III and IV were identified as 16R/S,17S-dihydroxy-7Z,10Z,12E,14E,19Z-docosapentaenoic acid (Figure 5B). For comparison LM profiling of the products obtained when LT4H was incubated with LTA4 gave a major peak that eluted with a TR of 13.6 min, with the MS-MS spectrum corresponding to LTBA (n = 4 incubations).

Another epoxide hydrolase expressed by human macrophages, which is also involved in the catalysis of allylic epoxides to bioactive mediators, is epoxide hydrolase 2 (EPHX2) (Deng et al., 2014). Thus, we next investigated whether this enzyme may play a role in the PDn-3 DPA biosynthetic pathway. Using flow cytometry we found that EPHX2 is expressed in monocytes and is upregulated during monocyte-to-macrophage differentiation (Figure 6A). Transfection of human monocytes with shRNA to EPHX2 led to a significant downregulation of PD2n-3 DPA and an increase in PD1n-3 DPA Concentrations (Figure 6B). To further validate the role of EPHX2 in PD2n-3 DPA biosynthesis, we incubated hr-EPHX2 with 16S,17S-ePDn-3 DPA, which gave a peak with TR of 15.2 min and an MS-MS spectrum that corresponded to that of PD2n-3 DPA (Figure 6C). Of note, in these incubations we did not identify PD1n-3 DPA, suggesting that this enzyme selectively catalyzed the conversion of 16S,17S-ePDn-3 DPA to PD2n-3 DPA. Having found that hr-EPHX2 converted the epoxide to PD2n-3 DPA, we next co-incubated hr-ALOX15 with hr-EPHX2 to obtain further evidence for the PD2n-3 DPA biosynthetic pathway in human cells. LM profiling of these incubations gave a peak with TR and MS-MS spectrum corresponding to PD2n-3 DPA (Figure 6D). Of note, levels of this mediator in incubations with either hr-ALOX15 or EPHX2 were negligible. Similar findings were also made with incubations of hr-ALOX15B and EPHX2 (n = 3 distinct incubations). To further assess the role of EPHX2 in the biosynthesis of PD2n-3 DPA we incubated this enzyme with increasing concentrations of 16S,17S-ePDn-3 DPA and determined the reaction kinetics. We found that 16S,17S-ePDn-3 DPA was rapidly converted with a Vmax of 3,439 ± 1,539 mmol/min and a Km of 332 ± 188 μM (Figure 6E). Together these findings demonstrate that epoxide hydrolase enzymes are key in the conversion of 16S,17S-ePDn-3 DPA to PD1n-3 DPA and PD2n-3 DPA, and identify EPHX2 as the enzyme responsible for the conversion of 16S,17S-ePDn-3 DPA to PD2n-3 DPA.

**PDn-3 DPA Regulate Human Macrophage Phenotype and Responses**

Having found that inhibition of the PDn-3 DPA pathway alters macrophage phenotype we next sought to determine the role of select members of the PDn-3 DPA pathway during monocyte-to-macrophage differentiation. Co-incubation of human macrophages with an ALOX15 inhibitor and either 16S,17S-ePDn-3 DPA or PD1n-3 DPA upregulated the expression of several lineage markers, including CD163 and CD64, when compared with cells incubated with the inhibitor alone (Figure S5A). We next investigated whether the upregulation of these lineage markers was also associated with a restoration of the ability of human macrophages to clear apoptotic cells. Indeed, incubation of monocyte-derived macrophages with 16S,17S-ePDn-3 DPA or PD1n-3 DPA rectified their ability to uptake apoptotic cells (Figures S5B and S5C). These restorative actions of PDn-3 DPA on macrophage function were also retained in vivo, where administration of PD1n-3 DPA rectified the expression of lineage markers on both splenic and peritoneal macrophages from ALOX15−/− mice (Figures 7A–7C; Table S3). Furthermore PD1n-3 DPA also upregulated the ability of peritoneal macrophages from ALOX15−/− mice to uptake apoptotic cells (Figure 7D) and bacteria (Figure 7E) in vivo. Together, these results demonstrate that the PDn-3 DPA pathway regulates key human macrophage functions in promoting the resolution of inflammation.

**DISCUSSION**

Herein, we established the PDn-3 DPA biosynthetic pathway and the role of this pathway in regulating key macrophage responses during monocyte-to-macrophage differentiation. Results from the present experiments demonstrate that, in human and mouse monocytes and macrophages, 15-lipoxygenases are the initiating enzyme in the PDn-3 DPA pathway catalyzing the formation of 17S-HpDPA and 16S,17S-ePDn-3 DPA. The epoxide is then converted via enzymatic hydrolysis to PD1n-3 DPA and PD2n-3 DPA, whereby the conversion of 16S,17S-ePDn-3 DPA to PD2n-3 DPA is catalyzed by EPHX2. Inhibition of ALOX15 activity led to a significant alteration in the expression of lineage markers as well as the ability of macrophages to uptake apoptotic cells and bacteria, actions that were recovered by either 16S,17S-ePDn-3 DPA or PD1n-3 DPA.

During acute inflammation, monocytes are recruited to the site where under ideal conditions, they differentiate to macrophages with a tissue protective phenotype; a process that is central in the resolution of inflammation as well as tissue repair and regeneration (Bystrom et al., 2008; Dalli and Serhan, 2016). Chronic, unresolved inflammation is associated with dysregulated monocyte differentiation leading to macrophages displaying a pro-inflammatory phenotype (Dalli and Serhan, 2016; Newson et al., 2014; Sica et al., 2000). The mechanisms dictating whether monocytes differentiate into a tissue reparative or pro-inflammatory phenotype remain of interest. In the present study, we found that inhibiting ALOX15 expression and activity using both genetic and pharmacological approaches in mice and human primary cells dysregulated both macrophage phenotype and function (Figure 1). Incubation of monocytes throughout their differentiation into macrophages...
Figure 7. PD1n-3 DPA Rectifies Murine Resident Macrophage Phenotype and Function in ALOX15-Deficient Mice

(A–C) The expression of phenotypic markers was assessed in peritoneal and splenic macrophages from ALOX15−/− mice administered PD1n-3 DPA (10 ng/mouse for 7 days) or vehicle and WT mice using flow cytometry and macrophage phenotype interrogated using PLS-DA in (A) large peritoneal macrophages, (B) small peritoneal macrophages, and (C) splenic macrophages. Results are representative of n = 8 mice per group for (A and B) and n = 3–4 mice per group for (C).
with either 16S,17S-ePDn-3 DPA or PD1n-3 DPA rescued the ability of macrophages to uptake apoptotic cells as well as upregulated the expression of lineage markers, including CD64 and CD163, which were decreased following ALOX15 inhibition. PD1n-3 DPA administration to ALOX15-deficient mice also rectified the expression of lineage markers on tissue-resident macrophages and the ability of peritoneal macrophages to uptake apoptotic cells and bacteria (Figure 7). Of note, ALOX15 is also the initiating enzyme in the RvD and LX biosynthetic pathway. Thus, while these results do not rule out the contribution of other ALOX15-derived pro-resolving mediators in the regulating monocyte-to-macrophage differentiation, they support a role for the PDn-3 DPA metabolite in regulating this process. The present findings also suggest that 16S,17S-ePDn-3 DPA is both a biosynthetic intermediate in the PDn-3 DPA pathway and exerts independent biological actions regulating cellular functions. These results are in line with findings made with other allylic epoxides including LAT4 (Ohishi et al., 1987) and 13S,14S-epoxy-maresin (Dalli et al., 2013c), which are also bioactive, regulating LM biosynthesis and cellular phenotype. LMs are produced via the stereoselective conversion of essential fatty acids by their biosynthetic enzymes. Recent studies demonstrate that the differential regulation of the SPM biosynthetic enzymes may reflect disease state and contribute to the propagation of ongoing inflammation. In this context, for example, Fredman et al. (2014) found that phosphorylation of the ALOX5 changes the product profile of the enzyme, whereby phosphopho-ALOX5 translocates to the nuclear membrane where it couples with phospholipase A2 and LAT4H to produce the potent leukocyte chemoattractant LTB4. In the absence of this phosphorylation the enzyme is found in the cytosol, where it couples with ALOX15, producing the tissue-protective and pro-resolving mediators RvD1 and LXA4 (Fredman et al., 2014). These mechanisms are linked with disease onset/progression whereby, in atherosclerosis, recent studies identified an increased expression of phosphorylated ALOX5 and a decrease in the RvD1 to LTB4 ratio in aortic lesions (Fredman et al., 2016). These results may also shed light on the apparently discordant results obtained with omega-3 supplementation. This is because, if the expression and/or subcellular localization of SPM biosynthetic enzymes is dysregulated, the product profile will be altered and the expected beneficial actions via SPM production may be diminished. Thus, furthering our understanding on the biosynthetic pathways governing SPM biosynthesis is essential in order to gain better insights into disease etiopathology. The present studies focus on establishing the role of the PDn-3 DPA in cells of the monocytic lineage. These findings demonstrate that human ALOX15 and ALOX15B catalyze the first two steps in the PDn-3 DPA biosynthetic pathway yielding an allylic epoxide. Using total organic synthesis, we established the absolute stereochemistry of this epoxide as 16S,17S-epoxy-7Z,10Z,12E,14E,19Z-docosapentaenoic acid. Furthermore, using this synthetic material we found that 16S,17S-ePDn-3 DPA was converted to PD1n-3 DPA and PD2n-3 DPA by different epoxide hydrolase enzymes, where in human cells EPHX2 was found to catalyze the conversion of 16S,17S-ePDn-3 DPA to PD2n-3 DPA (Figure 7). Of note, we also found that LAT4H was not involved in the biosynthesis of PD1n-3 DPA. In summation, the present findings establish the identity of enzymes involved in the PDn-3 DPA biosynthetic pathway, as well as the complete stereochemistry of the key intermediate in this pathway (Figure 7F). They also demonstrate that the PDn-3 DPA pathway plays a role in the differentiation of monocytes to macrophages, where inhibition of 15-lipoxygenase activity during monocyte-to-macrophage differentiation results in impaired monocyte-derived macrophage responses that are rescued by addition of PDn-3 DPA. Thus, these findings establish candidates in understanding the etiopathology of inflammatory diseases as well as targets for patient stratification and essential fatty acid supplementation.

**SIGNIFICANCE**

Macrophages are central players in controlling the body’s response to both sterile injury and infections. During the course of inflammation monocytes are recruited to the site of injury and/or infections, where they differentiate to macrophages. The phenotype displayed by these macrophages dictates whether the cells promote the termination of inflammation or lead to chronicity, with the mechanisms that control the monocyte-to-macrophage differentiation process remaining of interest. In the present study, we found that inhibition of ALOX15 activity during monocyte-to-macrophage differentiation reduced PDn-3 DPA production, altered human macrophage phenotype and their ability to clear dead cells, a key step in the termination of inflammation. Incubation of these cells with components of the PDn-3 DPA biosynthetic pathway rectified these responses. We established the identity and complete stereochemistry of 16S,17S-ePDn-3 DPA, a key intermediate in this pathway, and the role of human 15-lipoxygenases in producing this intermediate. We also provide evidence for a role of epoxide hydrolase enzymes in catalyzing the conversion of this intermediate to the pro-resolving mediators PD1n-3 DPA and PD2n-3 DPA in human macrophages. Here we found that EPHX2 selectively catalyzes the conversion of 16S,17S-ePDn-3 DPA to PD2n-3 DPA. Together these results establish the PDn-3 DPA pathway in human macrophages and the contribution of this pathway in monocyte-to-macrophage differentiation.

(D and E) Mice were treated as in (A–C), and on day 7 administered fluorescently labeled (D) apoptotic cells (6 × 10⁶ cells/mouse) or (E) E. coli (10⁶ CFU/mouse) via an intraperitoneal injection. Peritoneal cells were collected after 1 hr and phagocytosis was assessed in (D) CD64+ large peritoneal macrophages (left panel) and small peritoneal macrophages (right panel), and (E) total CD64+ macrophage population. Results are mean ± SEM. n = 8 mice per group for (D) and n = 4 mice per group for (E). *p < 0.05.

(F) Structures are illustrated in most likely configurations based on biosynthetic evidence. The stereochemistries for PD1n-3 DPA and 16S,17S-PDn-3 DPA are established (Aursnes et al., 2015; Aursnes et al., 2014; Dalli et al., 2013a).

Related to Figures S5.
STAR METHODS

Detailed methods are provided in the online version of this paper and include the following:

- KEY RESOURCES TABLE
- CONTACT FOR REAGENT AND RESOURCE SHARING
- EXPERIMENTAL MODEL AND SUBJECT DETAILS
  - Animal Studies
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- METHOD DETAILS
  - Monocyte Incubations
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  - Lipid Mediator Profiling
  - Total Organic Synthesis of 16S,17S-ePDn-3 DPA
- QUANTIFICATION AND STATISTICAL ANALYSIS
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SUPPLEMENTAL INFORMATION

Supplemental Information includes five figures and three tables and can be found with this article online at https://doi.org/10.1016/j.chembiol.2018.04.017.

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AUTHOR CONTRIBUTIONS

K.P., P.R.S., K.G.P., R.A.C., R.D.M., S.A.-W., R.M.M., F.M., and J.D. carried out the experiments and analyzed the data. A.V., T.V.H., and J.D. designed the experiments. All authors contributed to the preparation of the paper. J.D. designed the overall research plan.

DECLARATION OF INTERESTS

J.D. is an inventor on a patent detailing the structure and actions of mediators described in this manuscript (PCT/US2014/037969). Remaining authors declare no competing interests.

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## STAR METHODS

### KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| Antibodies          |        |            |
| APC/Cy7 anti-human CD 14 | Biolegend | Clone 63D3; Cat # 367107 |
| PE-Cy7 anti-human CD 32 | eBiosciences | Clone 6C4; Cat # 25-0329-41 |
| PE-Cy5 anti-CD 64   | Abcam  | Clone 10.1; Cat # ab192338 |
| Alexa Fluor 405 anti-ICAM-1 | Novus | Clone 1A29; Cat # NBP2-22541 |
| Alexa Fluor anti-human 488 CD 68 | Biolegend | Clone Y1/82A; Cat # 333811 |
| Brilliant Violet anti-human 650 CD 80 | Biolegend | Clone 2D10; Cat # 305227 |
| PerCP-Cy5.5 anti-human CD 206 | Biolegend | Clone 15-2; Cat # 321121 |
| PE-CF 594 mouse anti-human CD 163 | BD Biosciences | Clone GHI/61; Cat # 562670 |
| Monoclonal anti-ALOX15B | Sigma | Clone 4A7 Cat # SAB1402114-100UG |
| EPXH2 antibody [2F2] | Gene Tex | Cat # GTX84567 |
| Alexa Fluor 488 Goat anti-mouse IgG (H + L) | Invitrogen | Cat # A11029 |
| Alexa Fluor 647 15-Lipoxygenase 1 rabbit polyclonal | Biolegend | Cat # bs-6505R-A647 |
| APC-Cy7 anti-mouse CD 64 | Biolegend | Clone X54-5/7.1 Cat # 139303 |
| PE anti-mouse CD 64 (FCγRI) | Biolegend | Clone X54-5/7.1 Cat # 139303 |
| PE-Cy5 anti-mouse/human CD 11b | Biolegend | Clone M1/70 Cat # 101210 |
| Brilliant Violet 650 rat anti-mouse I-A/I-E (MHC II) | Biolegend | Clone M5/114.15.2 Cat # 107641 |
| APC-Cy7 rat anti-mouse F4/80 | Biolegend | Clone BM8 Cat # 123117 |
| BV785 hamster anti-mouse CD 11c | Biolegend | Clone N418 Cat # 117336 |
| PerCP-eFluor710 rat anti-mouse TIM-4 | eBiosciences | Clone 54 (RMT4-54) Cat # 46-5866-28 |
| Alexa Fluor 488 COX 2 | Cell Signaling Technologies | Clone (D5H5) XP ® Cat # 13596S |
| PE-Dazzle 594 rat anti-mouse IL-10 | Biolegend | Clone JESS-16E3 Cat # 505034 |
| Brilliant Violet mouse anti-mouse 421 TGF-β1 | Biolegend | Clone TW7-16B4 Cat # 141407 |
| PE sheep anti-mouse Arginase 1 | R&D | Cat # IC5868P |
| Alexa Fluor 647 rabbit- anti-human iNOS | Novus | Clone 4E5 Cat # NBP2-22119AF647 |
| TruStain Fc-blocking IgG (anti-mouse CD16/32) | Biolegend | Clone 93 Cat # 101310 |

(Continued on next page)
### REAGENT or RESOURCE SOURCE IDENTIFIER

#### Bacterial and Virus Strains

**Escherichia coli**
- **Strain**: O6:K2:H1

#### Biological Samples

**Leukocyte cones**
- **GBS Re NHS Blood/Transplant**: Cat # NC24

#### Chemicals, Peptides, and Recombinant Proteins

| Chemical, Peptide, or Protein | Source | Identifier |
|------------------------------|--------|------------|
| Ethanol, Absolute (200 proof), Mol Biology grade, Dnase, Rnase, Protease-free | Fisher Scientific Uk Ltd | Cat # 10644795 |
| Methanol, Optima(TM) LC/MS grade | Fisher Scientific Uk Ltd | Cat # 10767665 |
| Acetic acid | Fluka | Cat # 07692-1L-F |
| Methyl formate, 98% for spectroscopy | Fisher Scientific Uk Ltd | Cat # 10414315 |
| n-Hexane; For HPLC; 97+%; Acros Organics | Fisher Scientific Uk Ltd | Cat # 11934421 |
| Deuterium-labelled 5S-HETE | Cayman Chemicals | Cat #: 10007276 |
| Deuterium-labelled Leukotriene (LT) B4 | Cayman Chemicals | Cat #: 320110 |
| Deuterium-labelled Lipoxin (LX) A4 | Cayman Chemicals | Cat #: 10007737 |
| Deuterium-labelled Resolvin (Rv) D2 | Cayman Chemicals | Cat #: 11184 |
| Deuterium-labelled prostaglandin (PG) E2 | Cayman Chemicals | Cat #: 314010 |
| PGD2 | Cayman Chemicals | Cat #: 10007202 |
| PGE2 | Cayman Chemicals | Cat #: 10007211 |
| PGF2α | Cayman Chemicals | Cat #: 16010 |
| Thromboxane B2 | Cayman Chemicals | Cat #: 10007237 |
| LTB4 | Cayman Chemicals | Cat #: 10007240 |
| 20-OH-LTB4 | Cayman Chemicals | Cat #: 20190 |
| LXA4 | Cayman Chemicals | Cat #: 10007271 |
| LXB4 | Cayman Chemicals | Cat #: 90420 |
| 5S,12S-diHETE | In-house biogenic synthesis (J Dalli) | (Borgeat et al., 1982) |
| 5S,15S-diHETE | Cayman Chemicals | Cat #: 35280 |
| 15-epi-LXA4 | Cayman Chemicals | Cat #: 90415 |
| 15-epi-LXB4 | Custom Synthesis (Dr Charles Serhan, Harvard Medical School) | (Claria et al., 1996) |
| RvE1 | Cayman Chemicals | Cat #: 10007848 |
| RvE2 | Custom synthesis | (Serhan et al., 2000) |
| RvE3 | Custom synthesis | (Isobe et al., 2012) |
| RvD1 | Cayman Chemicals | Cat #: 10012554 |
| RvD2 | Cayman Chemicals | Cat #: 10007279 |
| RvD3 | Cayman Chemicals | Cat #: 13834 |
| RvD4 | Custom Synthesis (Dr Charles Serhan, Harvard Medical School) | (Winkler et al., 2016) |
| RvD5 | Cayman Chemicals | Cat #: 10007280 |
| RvD6 | In-house biogenic synthesis (J Dalli) | N/A |
| 17R-RvD1 | Cayman Chemicals | Cat #: 13096 |
| 17R-RvD3 | Custom Synthesis (Dr Charles Serhan, Harvard Medical School) | Dalli et al., 2013b |
| Maresin (MaR) 1 | Cayman Chemicals | Cat #: 10878 |
| MaR2 | Cayman Chemicals | Cat #: 16369 |
| 4S,14S-diHDHA | In-house biogenic synthesis (J Dalli) | (Serhan et al., 2009) |
| 7S,14S-diHDHA | In-house biogenic synthesis (J Dalli) | (Serhan et al., 2009) |
| 22-OH-MaR1 | In-house biogenic synthesis (J Dalli) | (Colas et al., 2016) |
| 14-oxo-MaR1 | In-house biogenic synthesis (J Dalli) | (Colas et al., 2016) |
| Protectin (PD)1 | Custom Synthesis (Dr Charles Serhan, Harvard Medical School) | (Petasis et al., 2012) |

(Continued on next page)
| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| 10S,17S-diHDHA      | Custom Synthesis [Dr Charles Serhan, Harvard Medical School] | (Petasis et al., 2012) |
| 22-OH-PD1           | Custom Synthesis [Dr Trond V. Hansen, University of Oslo] | (Tungen et al., 2014b) |
| n-3 DPA             | Custom Synthesis [Dr Trond V. Hansen, University of Oslo] | Item N° 21907 |
| RvD1,h-3 DPA        | In-house biogenic synthesis (J Dalli) | Dalli et al. (2013a) |
| RvD2,h-3 DPA        | In-house biogenic synthesis (J Dalli) | Dalli et al. (2013a) |
| RvD5,h-3 DPA        | In-house biogenic synthesis (J Dalli) | Dalli et al. (2013a) |
| MaR1,h-3 DPA        | Custom Synthesis [Dr Trond V. Hansen, University of Oslo] | (Tungen et al., 2014a) |
| 7S, 14S-diHDP A      | In-house biogenic synthesis (J Dalli) | Dalli et al. (2013a) |
| PD1,h-3 DPA         | Custom Synthesis [Dr Trond V. Hansen, University of Oslo] | Aursnes et al. (2014) |
| PD2,h-3 DPA         | In-house biogenic synthesis (J Dalli) | Dalli et al. (2013a) |
| 16S, 17S-ePDh-3 DPA  | Custom Synthesis [Dr Trond V. Hansen, University of Oslo] | This paper |
| 10S,17S-diHDP A      | In-house biogenic synthesis (J Dalli) | Dalli et al. (2013a) |
| Δ15trans-PD1h-3 DPA | In-house biogenic synthesis (J Dalli) | This paper |
| 10epi-Δ15trans-PD1h-3 DPA | In-house biogenic synthesis (J Dalli) | This paper |
| 17R-PD1             | Custom Synthesis [Dr Charles Serhan, Harvard Medical School] | Petasis et al. (2012) |
| RvT1                | In-house biogenic synthesis (J Dalli) | (Dalli et al., 2015) |
| RvT2                | In-house biogenic synthesis (J Dalli) | Dalli et al. (2015) |
| RvT3                | In-house biogenic synthesis (J Dalli) | Dalli et al. (2015) |
| RvT4                | In-house biogenic synthesis (J Dalli) | Dalli et al. (2015) |
| Histopaque – 1077   | Sigma | Cat # 10771-100ML |
| Dulbecco’s phosphate buffer saline with MgCl₂ and CaCl₂ (PBS+/+) | Sigma | Cat # D8662 |
| Dulbecco’s phosphate buffer saline without MgCl₂ and CaCl₂ (PBS-) | Sigma | Cat # D8537 |
| RPMI-1640           | Sigma | Cat # R8758 |
| RPMI 1640 Medium, no glutamine, no phenol red | Gibco/Life Technologies | Cat # 32404014 |
| Recombinant Human M-CSF | R&D | Cat # 216-MC-025 |
| PD146176            | Cambridge Bioscience | Cat # 10010518 |
| Human serum type AB (male) | Sigma | Cat # H4522-100ML |
| Penicillin-Streptomycin | Sigma | Cat # P4333 |
| Fetal bovine serum (FBS) | Gibco/Life Technologies | Cat # 105000-64 |
| DMSO, cell culture reagent | ChemCruz | Cat # sc-358801 |
| Bovine serum albumin (BSA) | Sigma | Cat # A9418 |
| EDTA                | Invitrogen | Cat # 15575-038 |
| PKH26 Red Fluorescent Cell Linker kit | Sigma | Cat # PKH26GL-1KT |
| PKH67 Green Fluorescent Cell Linker kit | Sigma | Cat # PKH67GL-1KT |
| Trypan Blue         | Sigma | Cat # T8154 |
| Polybrene Transfection Reagent | Millipore, UK Ltd | Cat # TR-1003-G |
| Recombinant Human GM-CSF | Biolegend | Cat # 572902 |
| Human recombinant INF-γ | Biolegend | Cat # 570206 |
| Human recombinant IL-4 | Biolegend | Cat # 574002 |
| Lipopolysaccharides | Sigma | Cat # L2630 |

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Continued

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| ALOX15 Human, 4 unique 29mer shRNA constructs in lentiviral GFP vector (Gene ID = 246). 5μg purified plasmid DNA per construct | OriGENE | TL314822 |
| MISSION shRNA Bacterial Clone ALXO15B | Sigma | Cat #s SHCLNG - TRCN0000432221, TRCN0000056583, TRCN0000056584, TRCN0000056585, TRCN0000056586 |
| MISSION shRNA Bacterial Glycerol Stock EPHX2 | Sigma | Cat #s SHCLNG - TRCN0000050553, TRCN0000050554, TRCN0000050555, TRCN0000050556, TRCN0000050557 |
| MISSION® pLKO.1-puro Empty Vector Control Plasmid DNA | Sigma | Cat # SHC001 |
| Trizma hydrochloride | Sigma | Cat # T3253 |
| hr-ALOX15 | Novus Biologicals | Cat # H00000246-P01 |
| hr-ALOX15B | Cayman Chemicals | Cat # 10011263-100ug-CAY |
| hr-EPHX2 | Cayman Chemicals | Cat # 10011669 |
| AUDA | Sigma | Cat # SML0177 |
| 4% PFA solution | Affymetrix | Cat # 19943 |
| Permeabilization buffer (10X) | eBiosciences | Cat # 00-8333-56 |
| Fixation/Permeabilization Concentrate | eBiosciences | Cat # 00-5123-43 |
| Fixation/Permeabilization Diluent | eBiosciences | Cat # 00-5223-56 |
| Hanks Balanced Salt solution | Sigma | Cat # H6648 |
| γ-Globulins from human blood (FC block) | Sigma | Cat # G4386-5G |
| LIVE/DEAD BacLight Bacterial Viability dye | Molecular Probes, Life Technologies | Cat # B35000 |

Critical Commercial Assays

| Name | Source | Cat # |
|------|--------|-------|
| APC Annexin V Apoptosis Detection Kit with PI | Biolegend | 640932 |
| EasySep™ Human Monocyte Isolation Kit | StemCell Technologies | 19319 |

Experimental Models: Organisms/Strains

| Name | Source | Cat # |
|------|--------|-------|
| Mus musculus, NCBI Taxonomy ID:10090, C57BL/6/+/FVB | Charles River | JAX™ C57BL/6J |
| Mus musculus, NCBI Taxonomy ID:10090, C57BL/6/+/FVB 12/15-LOX KO | The Jackson Laboratory | B6.129S2-ALox15tm1Fun/J Stock # 002778 |

Experimental Models: Cell lines

| Name | Source | Cat # |
|------|--------|-------|
| HL-60 | ATCC | CCL-240 |

Software and Algorithms

| Name | Source | URL | Notes |
|------|--------|-----|-------|
| FlowJo software | Tree Star, Ashland, OR | https://www.flowjo.com/solutions/flowjo/downloads | |
| GraphPad Prism 6.0f | GraphPad Software, CA | https://www.graphpad.com/support/prism-6-updates/ | |
| SIMCA 14.1 software | Umetrics, Umea, Sweden | https://umetrics.com/kb/simca-141 | |
| IDEAS® (Image Data Exploration and Analysis Software, Version 6.0) | Amnis®, EMD Millipore | http://www.merckmillipore.com/GB/en/life-science-research/cell-analysis-flow-cytometry/amnis-imaging-flow-cytometers/analysis-acquisition-software/ideas-software/qe2b.qB.Oq8AAAFLQM8J.x34R,n | |

Other

| Name | Source | Cat # |
|------|--------|-------|
| Isolute 500 mg / 3ml C18 SPE column | Biotage, Sweden | 220-0050-B |
| Poroshell 120 EC-18 4.6 mm 100 mm 2.7 μm reversed phase column | Agilent, USA | N/A |
| LSR Fortessa cell analyser | BD Biosciences, UK | N/A |
| Extra-Hera | Biotage, Sweden | N/A |

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CONTACT FOR REAGENT AND RESOURCE SHARING

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Jesmond Dalli (j.dalli@qmul.ac.uk).

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Animal Studies

Healthy 6-11 week old male C57/Black6 wild type mice (Charles River) and ALOX15−/− (The Jackson Laboratory) mice were used in the reported studies. The experiments strictly adhered to UK Home Office regulations (Guidance on the Operation of Animals, Scientific Procedures Act, 1986) and Laboratory Animal Science Association (LASA) Guidelines (Guiding Principles on Good Practice for Animal Welfare and Ethical Review Bodies, 3rd Edition, 2015). Animals were kept on a 12 h light dark cycle, with lights turned on at 7:00 h and lights turned off at 19:00h under specific pathogen free housing and had access to food and water ad libitum. Sample size was based on the statistical analysis of previous experiments and no mice were excluded. Animals were randomly assigned to control and experimental groups. The investigators were not blinded to group assignments.

Human Primary Cells

Healthy human peripheral blood mononuclear cells (PBMCs) were isolated from leukocyte cones obtained from the NHS Blood and Transplant bank and experiments were conducted in accordance with a protocol approved by Queen Mary Research Ethics Committee (QMREC 2014:61) and in accordance with the Helsinki declaration. Informed consent was obtained from all volunteers. Given that cones were obtained from the NHS Blood and Transplant bank and volunteers were unidentified, no information was available on sex, age, their involvement in previous experiments or if they were drug or test naïve. Primary cells were incubated at 37°C for 5 % CO2. The HL60 cell line was established in a female donor.

METHOD DETAILS

Monocyte Incubations

Human PBMCs were isolated from healthy human volunteers purchased from the NHS Blood and Transplant bank and experiments conducted in Queen Mary Research Ethics Committee approval (QMREC 2014:61) and the Helsinki declaration. Here blood cones were used and PBMCs were isolated by density centrifugation where cells were layered on to Histopaque -1077 (Sigma) and centrifuged for 30 minutes at 400 x g at room temperature (RT). Macrophages were prepared using published protocols (Dalli and Serhan, 2012) where PBMCs were plated into 10 cm tissue culture plates and incubated at 37°C for 30 min in PBS+. Non-adherent cells were then removed by adding PBS-/- and washed vigorously. Adherent cells (3 × 10⁷ cells/incubation) were then incubated with vehicle (PBS+0.1% ethanol), 1 nM 16S,17S-ePDn-3 DPA or 1 nM PD1n-3 DPA in RPMI 1640 (Sigma) for 30 minutes at 37°C. These were then incubated with either vehicle or ALOX15 inhibitor (10 μM PD146176 (Sendobry et al., 1997); Cambridge Bioscience) in RPMI 1640 containing 10% human serum, 1% Penicillin-Streptomycin and 20 ng/mL M-CSF. The cells were incubated for 3 days at 37°C at 5% CO2 and the media together with mediators and inhibitors were added as above and the cells were incubated a further 4 days. For lipid mediator profiling of cell supernatant, RPMI 1640 Medium (no glutamine) with no phenol red was used. Cell supernatants were collected and placed in two volumes of methanol containing deuterium labelled internal standards (500 pg of d₄-Prostaglandin (PG)E₂, d₄-LXA₄, d₄-RvD₂, d₄-Leukotriene (LT)B₄ and d₈-5S-Hydroxy-eicosatetraenoic acid; Cayman Chemicals). These were stored at -20°C until extraction (see below). In select experiments to obtain M1 macrophages adherent cells were cultured with GM-CSF (20ng/mL) for 6 days then with LPS (1ng/mL) and interferon-γ (20ng/mL) for 24h. To obtain M2 macrophages adherent cells where incubated with M-CSF (20ng/mL) for 7 days then with IL-4 (20ng/mL) for 48 h. In all cases the purity of the cell preparations ranged between 90-95%.

For human monocyte incubations we isolated monocytes from PBMCs using the EasySep™ Human Monocyte Isolation Kit, following manufacturer’s instruction where the purity of the resultant cell population was of ~95%. Human monocytes (1 × 10⁸ cells/mL) were incubated with n-3 DPA (10μM) and Escherichia coli (5 × 10⁹ CFU/mL) in PBS at 37°C. Incubations were quenched

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| TurboVap LV         | Biotage, Sweden | N/A        |
| Qtrap 5500/6500     | AB Scix | N/A        |
| Shimadzu SIL-20AC auto-injector | Shimadzu Corp. | N/A        |
| LC-20AD Binary pump | Shimadzu Corp. | N/A        |
| FLUOstar Omega microplate reader | BMG Labtech | N/A        |
| ImageStream X MK2 Imaging Flow Cytometer | Amnis®, EMD Millipore | N/A        |
using excess acidified methanol (apparent pH ~3) containing deuterium labelled \( \text{d}_3-\text{LTB}_4 \) and products were extracted as detailed below.

In select experiments monocytes (1x10^6 cells/ml) were incubated with vehicle (PBS + 0.01% DMSO) or 12-[[tricyclo[3.3.1.1\text{3,7}]dec-1-ylamino(carbonyl)amino]-dodecanoic acid (AUDA; 25\( \mu \)M) for 20 min at RT. Cells were then incubated with either vehicle (PBS + 0.1% EtOH) or 16S,17S-e-PDn-3 DPA (10nM) for 15 min. Incubations were then quenched and products identified and profiled using LM profiling as detailed below.

Monocytes (1.5x10^6 cells/ml) were suspended in phenol red free RPMI 1640 containing 10% human serum, 8\( \mu \)g/ml of polybrene and 50\( \mu \)g/ml of shRNA to ALOX15 or CT shRNA. The cell suspension was then centrifuged at 1000 x g for 90 min at 4°C. Cells were then plated and incubated for a further 12h at 37°C. Monocytes were then detached after 5 minutes incubation in 2mM EDTA in PBS−/−, and the expression of ALOX15 was evaluated using flow cytometry. Cells were also suspended in PBS containing 0.1% human serum and incubated with E. coli (1:50 monocytes to bacteria) for 45 min at 37°C. Incubations were quenched using two volumes of ice-cold methanol and products profiled using lipid mediator profiling.

In separate experiments monocytes were plated in 6 well plates and incubated with phenol red free RPMI 1640 containing 10% human serum, 8\( \mu \)g/ml of polybrene and 2\( \mu \)g/ml of shRNA to ALOX15, EPHX2 or CT shRNA. The cell incubations were gently mixed for the first 5 hours, media was then removed and cells incubated with 15% DMSO in Hanks’ Buffered Saline Solution for 4 mins at 37°C. The solution was then removed, cells were rinsed with PBS and then incubated for a further 7h in phenol red free RPMI containing 10% human serum and 1% Penicillin-Streptomycin. Cells were then detached after 5 minutes incubation in 2mM EDTA in PBS−/−, and expression of ALOX15 and EPHX2 was evaluated using flow cytometry. Cells were also suspended in PBS containing 0.1% human serum and incubated with E. coli as detailed above.

Mouse monocytes were isolated by adhesion of bone marrow cell suspension from WT and ALOX15-deficient mice to tissue culture dishes. After 45 min cell were detached and cell populations determined to be ~80-90% monocytes. Monocytes (2 x 10^6 cells/ml) were incubated in phenol red free RPMI containing 0.1% PBS and E. coli (1 x 10^8 CFU/mL) at 37°C for 45 min. Incubations were quenched using 2 volumes of ice-cold methanol containing deuterium labelled internal standards and lipid mediators identified and quantified using lipid mediator profiling.

**Enzyme Incubations**

Human recombimant (hr)-ALOX15 (0.2\( \mu \)M; Novus Biologicals) or (hr)-ALOX15B (0.2\( \mu \)M; Cayman Chemicals) was incubated with n-3 DPA (10 nM) in Tris buffer (pH = 8.0) at RT for 2 min. Products were quenched using excess acidified methanol (apparent pH ~3) containing deuterium labelled \( \text{d}_3-\text{LTB}_4 \) and extracted as detailed in the lipid mediator profiling section below. In select incubations hr-AOX15 or hr-ALOX15B were incubated with EPHX2 (0.2\( \mu \)M; Cayman Chemicals) and n-3 DPA (10\( \mu \)M) in Tris buffer (pH = 8.0) at RT for 15 min. Incubations were quenched using ice-cold methanol and products extracted, identified and quantified as detailed below.

In select incubations EPHX2 (0.2\( \mu \)M) was incubated with 16S,17S-ePDn-3 DPA (10nM) in Tris buffer (pH = 8.0) at RT for 10 min; incubations were then quenched, and products identified and quantified as detailed below.

**Flow Cytometry**

Flow cytometry was used to determine the phenotypic lineage of the monocyte-derived macrophages using fluorescently conjugated antibodies. Cells were fixed in 1% paraformaldehyde (4% paraformaldehyde in PBS−/−) for 10 minutes at room temperature. Cells were incubated with the following antibodies in a 1:100 dilution for 30 minutes at 4°C: APC/Cy7 anti-human CD14 (clone 63D3, Biolegend), PE-Cyanine7 anti-human CD32 (clone 6C4, eBiosciences), Alexa Fluor (AF) 405 anti-ICAM-1 (clone 1A29, Novus), PE/Cy5 anti-CD64 (clone 10.1, Abcam), AF 488 anti-human CD68 (clone Y1/82A, Biolegend), Brilliant Violet (BV) 650 anti-human CD80 (clone 2D10, Biolegend), PerCP/Cy5.5 anti-human CD206 (clone 15-2, Biolegend) and PE-CF594 anti-human CD163 (clone GHI/61, BD Biosciences). LSR Fortessa cell analyser (BD Biosciences) was used to perform multivariate parameter analysis, followed by analysis using FlowJo (Tree Star Inc., V10).

To determine the expression of PDn-3 DPA biosynthetic enzymes, monocytes and macrophages were incubated with M-CSF and IL-4 or GM-CSF, IFNγ and LPS for 7 days as detailed above. At the indicated intervals cells were collected, permeabilized using eBioscences Fixation/Permeabilization Solution Kit following manufacturer’s instructions, non-specific binding was quenched using non-specific IgG (16\( \mu \)g/mL) and cells were then incubated with rabbit anti-human 15-LOX type 1, mouse anti-human 15-LOX type 2, and mouse anti-human EH2 for 30 min at 4°C. To determine the expression of 15-LOX type 2 and EH2, cells were then incubated with AF 488 goat anti-mouse IgG for 30 min at 4°C and staining evaluated using LSR Fortessa cell analyser (BD Biosciences) was used to perform multivariate parameter data acquisition, followed by analysis using FlowJo (Tree Star Inc., V10).

In select experiments cells were collected from the peritoneum of WT and ALOX1-deficient mice by lavaging the peritoneum with PBS. Splenic cells were collected using the staining solution with PBS and ALOX15-deficient mice using a 70 \( \mu \)m filter. The cells were incubated with the following antibodies in a 1:100 dilution for 30 minutes at 4°C in staining solution (1:1 PBS with 0.02 % BSA and TrueStain FC Blocking IgG): PE-Cy5 anti-mouse/human CD 11b (Clone M1/70, Biolegend), BV 650 rat anti-mouse I-A/I-E (MHC II) (Clone M5/114.15.2, Biolegend), APC-Cy7 rat anti-mouse F4/80 (Clone BM8, Biolegend), BV785 hamster anti-mouse CD 11c (Clone N418, Biolegend) and PerCP-eFluor710 rat anti-mouse TIM-4 (Clone 54 (RM4T-54), eBiosciences). For intracellular staining, cells were permeabilized using eBioscences Fixation/Permeabilization Solution Kit following manufacturer’s instructions for 20 min at RT and then with fluoroscenically labelled antibodies in a 1:50 dilution for 30 minutes at 4°C in staining solution.
In Vitro Macrophage Incubations

(1:10 Permeabilization buffer (10X) (eBiosciences) and PBS with 0.02 % BSA); AF 488 COX 2 (Clone (D5H5) XP ®, Cell Signaling Technologies), PE-Dazzle 594 rat anti-mouse IL-10 (Clone JESS-16E3, Biolegend), BV 421 mouse anti-mouse TGF-β1 (Clone TW7-16B4, Biolegend), PE sheep anti-mouse Arginase 1 (Cat # IC5868P, R&D) and AF 647 rabbit- anti-human iNOS (Clone 4E5, Novus). Non-specific binding was quenched using TruStain Fc-blocking IgG (anti-mouse CD16/32) and the staining was evaluated as above. For cells obtained from both in vivo efferocytosis and phagocytosis experiments non-specific binding was quenched as detailed above. Cells were then incubated in PBS containing 0.02% BSA and either APC-Cy anti-mouse CD64 or PE anti-mouse CD64 at 1:100 dilution for 30 min at 4°C and staining evaluated.

Macrophage Incubations

In Vitro Efferocytosis

Macrophages, prepared as described above, were seeded into 96 well plates at 5 x 10^4 cells per well in RPMI 1640 medium with 10% human serum at 37°C. Apoptotic cells were prepared by incubating the human promyelocytic leukemic cell line HL60 at 70°C for 2h yielding ~60% Annexin V positive / propidium iodide negative cells and 35% Annexin V positive / propidium iodide positive cells using APC Annexin V Apoptotic Detection Kit with PI (Biolegend). These were then stained using PKH26 Red Fluorescent Cell Linker kit (Sigma) following manufacturer’s instructions and efferocytosis was assessed as in (Dalli and Serhan, 2012). In brief, fluorescently labelled apoptotic cells (1:3, macrophages to apoptotic cells) were added to the macrophages and incubated at 37°C for 45 minutes. Cells were washed with PBS and extracellular fluorescence was quenched using trypan blue (1:15 in PBS). Fluorescence was then measured using a FLUOStar Omega microplate reader (BMG Labtech). In select experiments apoptotic cell were labelled using PKH67 and incubated with macrophages as detailed above. Macrophage efferocytosis was then evaluated using an ImageStream X MK2 and analysis was performed using IDEAS® (Image Data Exploration and Analysis Software, Version 6.0).

In Vivo Efferocytosis

Mice were administered either vehicle or PD1n-3 DPA (10ng/mouse) via i.p. injection for 7 days. Apoptotic cells were prepared and labelled as detailed above using PKH67 dye. These (6x10^6 cells per mouse) were injected via intraperitoneal injection to WT and ALOX15-deficient mice. After 1h cells were collected by peritoneal lavages and incubated with an APC-Cy7 labelled anti-mouse CD64 antibody as detailed above and efferocytosis in CD64 positive macrophages was evaluated using Flow cytometry.

E. coli Incubations

Macrophages (2x10^6 cells) were incubated with E. coli (1x10^7 cells) for 30 minutes at 37°C. Incubations were quenched using two volumes of ice-cold methanol containing deuterium labelled internal standards (500 pg each of d2-PGE2, d3-LXA4, d4-RvD2, d4-LTB4 and d8-5S-HETE, Cayman Chemicals). Samples were then stored at -20°C prior to extraction and lipid mediator profiling.

In Vivo Phagocytosis

Mice were administered either vehicle or PD1n-3 DPA (10ng/mouse) via i.p. injection for 7 days. E. coli were labelled using LIVE/DEAD BacLight Bacterial Viability dye following manufacturer’s instructions. Fluorescently labelled bacteria were then injected via intraperitoneal injection to WT and ALOX15-deficient mice and peritoneal lavages were collected after 1h. Cells were stained using fluorescently labelled PE anti-mouse CD64 antibody and bacterial phagocytosis was evaluated using flow cytometry as detailed above.

Lipid Mediator Profiling

Proteins were allowed to precipitate by keeping samples at -20°C for 30 minutes. Deuterium-labelled standards, d2-PGE2, d3-LXA4, d4-RvD2, d4-LTB4 and d8-5S-HETE were used to aid in identification and quantification of lipid mediators (Dalli and Serhan, 2012). Following protein precipitation, samples were then extracted using an ExtraHera (Biotage) with ISOLUTE C18 columns (500 mg, 3 mL; Biotage). This involved the conditioning of the Solid-phase ISOLUTE C18 500 mg/3 mL columns with methanol for 60 seconds using 2.5 bar positive pressure. Samples, brought to 10 mL with pH 3.5 water, were loaded onto the columns for 90 seconds using 2.5 bar positive pressure. The acid in the C18 columns was neutralised by washing the columns with 2 mL pH 7 water for 45 seconds using 2.5 bar positive pressure, followed by the hexane wash to elute hydrophobic molecules. This was done four times with 3 mL hexane using 2.5 bar positive pressure for 60 seconds for each wash. Mediators were then eluted into collection tubes with the addition of 5 mL methyl formate at 1.5 bar positive pressure for 120 seconds. Products were brought to dryness using a gentle nitrogen stream and TurboVap LV (Biotage), these were then suspended in phase containing methanol and water, 1:1 (vol/vol).

An LC-MS-MS system, comprising of a Qtrap 5500 (AB Sciex) or Qtrap 6500 plus (AB Sciex), Shimadzu SIL-20AC autoinjector, LC-20AD binary pump (Shimadzu Corp.) and Agilent C18 Poroshell column (150 mm x 4.6 mm x 2.7 μm) was used to profile lipid mediators. The gradient was initiated at 20:80:0.01 (vol/vol/vol) methanol/water/acetic acid for 0.2 mins this was ramped to 50:50:0.01 (vol/vol/vol) over 12 seconds, maintained for 2 minutes, then ramped to 80:20:0.01 (vol/vol/vol) over 9 minutes, and maintained for 3.5 minutes. The ratio was then ramped to 98:2:0.01 (vol/vol/vol) for 5.5 minutes. The flow rate was kept at 0.5 mL/minute throughout.

Mediator concentrations were determined using multiple reaction monitoring (MRM) using signature parent ion (Q1) and characteristic daughter ion (Q3) pairs. A minimum of six diagnostic ions were used to confirm their identities, using published criteria (Dalli and Serhan, 2012). The peak area of the MRM transition and linear calibration curves with an r² value of 0.98 to 0.99 were used to quantify each of the molecules. The detection limit was ~0.1 pg.
Total Organic Synthesis of 16S,17S-EPDn-3 DPA

Unless stated otherwise, all commercially available reagents and solvents were used in the form they were supplied without any further purification. The stated yields are based on isolated material. All reactions were performed under an argon atmosphere using Schlenk techniques. Reaction flasks were covered with aluminium foil during reactions and storage to minimize exposure to sunlight. Thin layer chromatography was performed on silica gel 60 F254 aluminium-backed plates fabricated by Merck. Flash column chromatography was performed on silica gel 60 (40–63 μm) produced by Merck. NMR spectra were recorded on a Bruker AV600, Bruker AVII400 or a Bruker DPX300 spectrometer at 600 MHz, 400 MHz or 300 MHz respectively for 1H NMR and at 150 MHz, 100 MHz or 75 MHz respectively for 13C NMR. Coupling constants (J) are reported in hertz and chemical shifts are reported in parts per million (δ) relative to the central residual protium solvent resonance in 1H NMR (CDCl3 = δ = 7.26, DMSO-d6 = δ = 2.50, benzene-d6 = δ = 7.16 and MeOD-d4 = δ = 3.31 ppm) and the central carbon solvent resonance in 13C NMR (CDCl3 = δ = 77.00, DMSO-d6 = δ = 128.06 and MeOD-d4 = δ = 49.00 ppm). Mass spectra were recorded at 70 eV on Micromass Prospec Q or Micromas QTOF 2W spectrometer using EI, ES or CI as the methods of ionization. High resolution mass spectra were recorded on Micromass Prospec Q or Micromas QTOF 2W spectrometer using EI or ES as the methods of ionization. Optical rotations were measured using a 0.7 mL cell with a 1.0 dm path length on an Anton Paar MCP 100 polarimeter. HPLC analyses were performed on an Agilent Technologies 1200 Series instrument with diode array detector set at 254 nm and equipped with a C18 stationary phase (Eclipse XDB-C18 5 μm, 4.6 × 150 mm), applying the conditions stated. GLC analyses were performed on an Agilent 7820A with a FID detector, HP-5 capillary column, with helium as the carrier gas and by applying the conditions stated.

Synthesis of methyl (7Z,10Z,12E,14E)-15-(2S,3S)-3-(Z-pent-2-en-1-yl)oxiran-2-yl)pentadeca-7,10,12,14-tetraenoate (methyl EPD1n-3 DPA)

This reaction was performed as earlier reported (Aursnes et al., 2014). In brief, 1.3 equiv. of the Wittig salt methyl (Z)-10-(iodotriphenyl-λ5-phosphanyl)dec-7-enoate was dissolved in THF and then HMPA was added. This solution was cooled to ~78°C. Then NaHMDS (0.60 M in toluene, 1.3 equiv.) was added drop-wise. The reaction mixture was stirred for 30 min at this temperature. The epoxy aldehyde (2E,4E)-5-(2S,3S)-3-(Z-pent-2-en-1-yl)oxiran-2-yl)penta-2,4-dienal was prepared from commercially available 2-(triphényl-λ5-phosphanylidene)acetaldheyde and (2R,3S)-3-(Z-pent-2-en-1-yl)oxyiran-2-carbaldehyde as described (Aursnes et al., 2015). The Wittig-salt methyl (Z)-10-(iodotriphenyl-λ5-phosphanyl)dec-7-enoate was made according to literature protocols (Primidahl et al., 2017).

Z-Selective Wittig Reaction

This reaction was performed as earlier reported (Aursnes et al., 2014). In brief, 1.3 equiv. of the Wittig salt methyl (Z)-10-(iodotriphenyl-λ5-phosphanyl)dec-7-enoate was dissolved in THF and then HMPA was added. This solution was cooled to ~78°C. Then NaHMDS (0.60 M in toluene, 1.3 equiv.) was added drop-wise. The reaction mixture was stirred for 30 min at this temperature. The epoxy aldehyde (2E,4E)-5-(2S,3S)-3-(Z-pent-2-en-1-yl)oxiran-2-yl)penta-2,4-dienal was prepared from commercially available 2-(triphényl-λ5-phosphanylidene)acetaldheyde and (2R,3S)-3-(Z-pent-2-en-1-yl)oxyiran-2-carbaldehyde as described (Aursnes et al., 2015). The Wittig-salt methyl (Z)-10-(iodotriphenyl-λ5-phosphanyl)dec-7-enoate was made according to literature protocols (Primidahl et al., 2017).

QUANTIFICATION AND STATISTICAL ANALYSIS

Results are represented as mean ± SEM. Differences between groups were assessed using one-sample t test (normalized data), Student’s t test (2 groups), 1-way ANOVA (multiple groups) followed by post hoc Dunnett’s test using GraphPad Prism 6 software. Investigators were not blinded to group allocation or outcome assessment. The criterion for statistical significance was P ≤ 0.05.
Sample sizes for each experiment were determined on the variability observed in prior experiments (Dalli and Serhan, 2016) and preliminary experiments. Partial least squares-discrimination analysis (PLS-DA) and principal component analysis (PCA) (Janes and Yaffe, 2006) were performed using SIMCA 14.1 software (Umetrics, Umea, Sweden) following mean centering and unit variance scaling of LM levels. PLS-DA is based on a linear multivariate model that identifies variables that contribute to class separation of observations on the basis of their variables (LM levels). During classification, observations were projected onto their respective class model. The score plot illustrates the systematic clusters among the observations (closer plots presenting higher similarity in the data matrix). Loading plot interpretation identified the variables with the best discriminatory power (Variable Importance in Projection greater than 1) that were associated with the distinct intervals and contributed to the tight clusters observed in the Score plot.

DATA AND SOFTWARE AVAILABILITY

Data are available upon request to the Lead contact.