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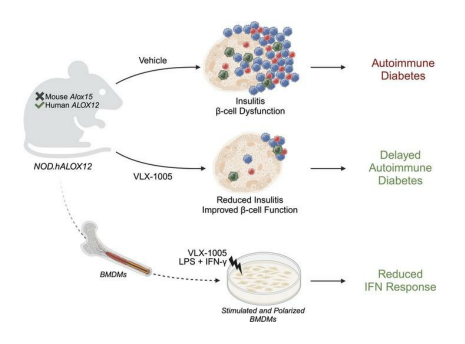
## 12-Lipoxygenase inhibition delays onset of autoimmune diabetes in human gene replacement mice

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## **Graphical abstract**





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## 12-Lipoxygenase inhibition delays onset of autoimmune diabetes in human gene replacement mice

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Type 1 diabetes (T1D) is characterized by the autoimmune destruction of insulin-producing  $\beta$  cells and involves an interplay between  $\beta$  cells and cells of the innate and adaptive immune systems. We investigated the therapeutic potential of targeting 12-lipoxygenase (12-LOX), an enzyme implicated in inflammatory pathways in  $\beta$  cells and macrophages, using a mouse model in which the endogenous mouse Alox15 gene is replaced by the human ALOX12 gene. Our finding demonstrated that VLX-1005, a potent 12-LOX inhibitor, effectively delayed the onset of autoimmune diabetes in human gene replacement non-obese diabetic mice. By spatial proteomics analysis, VLX-1005 treatment resulted in marked reductions in infiltrating T and B cells and macrophages, with accompanying increases in immune checkpoint molecule PD-L1, suggesting a shift toward an immunosuppressive microenvironment. RNA sequencing analysis of isolated islets and polarized proinflammatory macrophages revealed significant alteration of cytokine-responsive pathways and a reduction in IFN response after VLX-1005 treatment. Our studies demonstrated that the ALOX12 human replacement gene mouse provides a platform for the preclinical evaluation of LOX inhibitors and supports VLX-1005 as an inhibitor of human 12-LOX that engages the enzymatic target and alters the inflammatory phenotypes of islets and macrophages to promote the delay of autoimmune diabetes.

#### Introduction

The pathogenesis of type 1 diabetes (T1D) involves a complex interplay between multiple cell types within the pancreatic islet, including innate immune cells (macrophages, dendritic cells), insulin-producing cells ( $\beta$ cells), and adaptive immune cells (T cells, B cells) (1). Although the disease has traditionally been viewed as arising from a primary defect in immune tolerance, an emerging perspective posits that environmental factors (such as viruses or other systemic inflammatory disorders) may aggravate an interaction between macrophages and  $\beta$  cells, facilitating oxidative and endoplasmic reticulum (ER) stress pathways in  $\beta$  cells (2–4). These pathways facilitate the generation of  $\beta$  cell neoepitopes that then trigger adaptive autoimmunity (5, 6). Disease-modifying therapies — those that alter disease pathogenesis rather than correcting the underlying disease phenotypes — have largely focused on the adaptive immune system and seen some successes in clinical trials. For example, an anti-CD3 monoclonal antibody (teplizumab) that targets activated T cells has been shown to delay the onset of T1D by up to 2 years in individuals at high risk for the disease (7). Given the increasing appreciation of innate immune cells and  $\beta$  cells in early T1D pathogenesis, the identification of drugs targeting these cell types raises the possibility that combination therapeutic approaches may provide more durable outcomes.

The lipoxygenases (LOXs) encompass a family of enzymes involved in lipid metabolism that facilitates the oxygenation of polyunsaturated fatty acids to form eicosanoids, some of which are proinflammatory in nature (8). In the mouse, 12/15-LOX is encoded by the *Alox15* gene and is the primary active LOX present in macrophages and  $\beta$  cells and produces the proinflammatory eicosanoid 12-hydroxyeicosatetraenoic acid (12-HETE) as a principal product from the substrate arachidonic acid (9). Whole-body deletion of *Alox15* 

**Conflict of interest:** RGM and SAT received an investigator-initiated award from Veralox Therapeutics. RGM serves on the Scientific Advisory Board for Veralox Therapeutics. DJM and MBB are Veralox Therapeutics employees.

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**Reference information:** *JCI Insight*. 2024;9(24):e185299. https://doi.org/10.1172/jci. insight.185299. on the autoimmune non-obese diabetic (NOD) mouse background results in almost complete protection against diabetes (10). Deletion of *Alox15* in either the innate immune myeloid cells (2) or in  $\beta$  cells (11) recapitulates the autoimmune diabetes protection seen in the whole-body deletion, emphasizing both the early role of these cell types in T1D and the importance of the 12/15-LOX pathway in disease pathogenesis. In these cell-specific deletion models, islets exhibit marked reductions in invading pathogenic T cells (insulitis), a finding reflecting the disease-modifying response. The molecular events tied to disease protection ostensibly emanate from reductions in oxidative and ER stress (and the resultant reduction in neoepitope formation and presentation) as well as from enhanced display of PD-L1 (an immunosuppressive checkpoint ligand) on the surface of myeloid cells and  $\beta$  cells (2, 11).

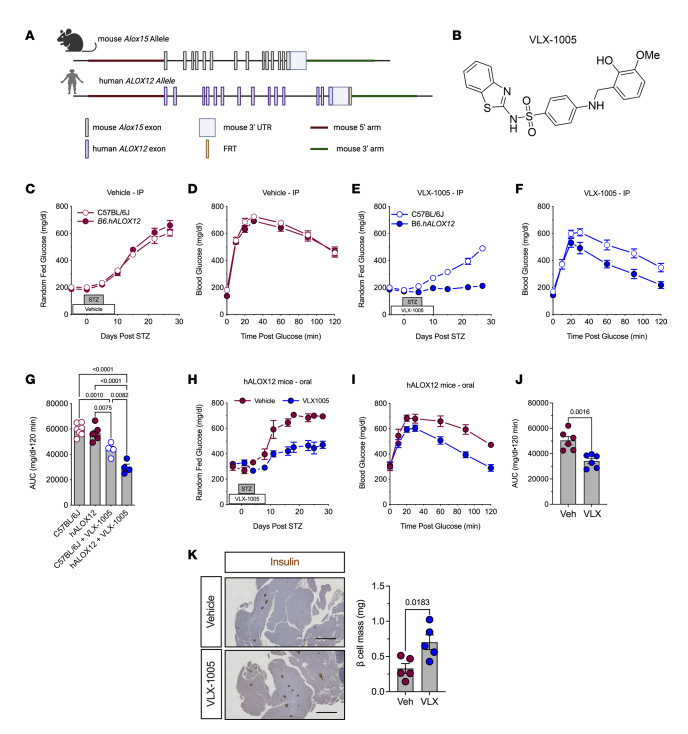
In humans, the relevant LOX enzyme that produces 12-HETE is 12-LOX, encoded by the *ALOX12* gene. Like the mouse 12/15-LOX, human 12-LOX is present in residual insulin-positive cells in donors with T1D or in autoantibody-positive donors at risk for T1D (12) — a finding consistent with a potential role in promoting  $\beta$  cell sensitivity to autoimmunity. A major challenge to using mice as a platform to test inhibitors is that human 12-LOX exhibits structurally distinct characteristics from mouse 12/15-LOX, thereby necessitating the development of different inhibitors that cannot be tested for efficacy in mice (13–15). Previously, VLX-1005 (also known as ML355) was described as a potent and selective inhibitor of human 12-LOX, while also displaying a favorable half-maximal inhibitory concentration (IC<sub>50</sub>) and pharmacokinetic properties (16). VLX-1005 was shown to protect human islets in vitro against dysfunction caused by proinflammatory cytokines (17), but the lack of appropriate in vivo model systems has made it challenging to pharmacologically validate VLX-1005 as a therapeutic target in autoimmune diabetes. To address this challenge, we developed new mouse strains in which the mouse *Alox15* gene is replaced by the human *ALOX12* gene while retaining the mouse gene's upstream control elements. This human gene replacement platform was leveraged to test whether and how pharmacologic inhibition of human 12-LOX with VLX-1005 modifies disease progression in autoimmune T1D.

#### Results

Generation and validation of the hALOX12 gene replacement mouse model. To establish a platform to test potential inhibitors of human 12-LOX in vivo, we generated a mouse model in which the endogenous mouse *Alox15* gene is replaced by the human *ALOX12* gene (Figure 1A). This model leaves the mouse upstream regulatory region intact to ensure that the expression of *ALOX12* recapitulates the expression of *Alox15*. These mice (henceforth referred to as *hALOX12* mice) were introgressed onto the C57BL/6J (also known as B6) mouse background using a speed congenics approach and bred to homozygosity. Microsatellite genotyping showed that the mice were 100% congenic on the C57BL/6J background (Supplemental Table 1; supplemental material available online with this article; https://doi.org/10.1172/jci.insight.185299DS1). To confirm the successful deletion of mouse *Alox15* and replacement with human *ALOX12*, we performed standard genotyping (Supplemental Figure 1A). Additionally, we isolated tissues (kidney, spleen, lung, fat, liver, islets, peritoneal macrophages, and bone marrow–derived macrophages [BMDMs]) from WT C57BL/6J and *B6.hALOX12* mice and subjected them to gene expression analysis for *Alox15* and *ALOX12*. As anticipated, WT tissues expressed mouse *Alox15* and did not express human *ALOX12*; conversely, *B6. hALOX12* mouse tissues expressed *ALOX12* but not *Alox15* (Table 1).

Because lipoxygenases are known to affect metabolic function, we next performed metabolic characterization to determine whether/how the replacement of *Alox15* with *ALOX12* altered metabolic phenotypes. We found no significant differences in body weight, lean mass, fat mass, random-fed blood glucose levels, or glucose tolerance between WT C57BL/6J and *B6.hALOX12* mice (Supplemental Figure 1, B–F). Moreover, islet ultrastructure (relative immunostaining patterns of  $\alpha$  cells and  $\beta$  cells) and composition ( $\alpha$  cell mass and  $\beta$  cell mass) were indistinguishable between 10-week-old WT and *B6.hALOX12* mice (Supplemental Figure 1, G–I). Taken together, these data suggest that the successful replacement of *Alox15* with human *ALOX12* did not alter gross metabolic or islet phenotypes.

Effects of VLX-1005 against STZ-induced diabetes are specific to B6.hALOX12 mice. Prior studies demonstrated that whole-body deletion of mouse Alox15 protects against diabetes induced by the chemical streptozotocin (STZ) (18). To test whether the human 12-LOX inhibitor VLX-1005 (14) (Figure 1B) phenocopies deletion of the enzyme in our human gene replacement mice, we employed a similar STZ diabetes induction protocol. STZ is a  $\beta$  cell toxin that induces low-grade inflammation, macrophage influx into islets, and eventual diabetes in mice after 5 daily low-dose i.p. injections (55 mg/kg) (19). Eight-week-old male WT C57BL/6J and B6.hALOX12 mice were injected i.p. daily with vehicle or



**Figure 1. 12-LOX inhibition protects against streptozotocin-induced diabetes.** C57BL/6J and *B6.hALOX12* male mice (*n* = 4–7 per group as indicated) were treated with 30 mg/kg i.p. or p.o. VLX-1005 and multiple low-dose streptozotocin (STZ). (**A**) Schematic of the generation of *hALOX12* mice by replacing mouse *Alox15* with human *ALOX12*. (**B**) Chemical structure of VLX-1005. (**C**) Random-fed blood glucose values in vehicle-treated male C57BL/6J and *B6.hALOX12* mice on day 4 after STZ treatment. (**E**) Random-fed blood glucose values in VLX-1005–treated male C57BL/6J and *B6.hALOX12* mice after STZ. (**D**) GTT of vehicle-treated male C57BL/6J and *B6.hALOX12* mice on day 4 after STZ treatment. (**E**) Random-fed blood glucose values in VLX-1005–treated male C57BL/6J and *B6.hALOX12* mice after STZ. (**F**) GTT of VLX-1005–treated male C57BL/6J and *B6.hALOX12* mice after STZ. (**F**) GTT of VLX-1005–treated male C57BL/6J and *B6.hALOX12* mice after STZ. (**F**) GTT of VLX-1005–treated male C57BL/6J and *B6.hALOX12* mice after STZ. (**F**) GTT of VLX-1005–treated male C57BL/6J and *B6.hALOX12* mice after STZ. (**F**) GTT of VLX-1005–treated (p.o.) *B6.hALOX12* mice after STZ. (**I**) GTT of male vehicle- or VLX-1005–treated (p.o.) *B6.hALOX12* mice after STZ treatment. (**I**) AUC of *B6.hALOX12* mice after STZ treatment. (**K**) Pancreata stained for insulin (left panel) and β cell mass measurement (right panel) from male *B6.hALOX12* mice on day 26 after STZ treatment. Scale bars: 500 µm. Data are presented as mean ± SEM and statistical significance was determined by a 2-tailed Student's t test (all except **G**) or 1-way ANOVA with Tukey's post hoc test (**G**).

Mouse strain	Tissues	ALOX12 (ACt)	<i>Alox15</i> (∆Ct)
C57BL/6J	Islets	ND	7.66 ± 0.32
	Spleen	ND	18.41 ± 1.31
	BMDMs	ND	17.47 ± 0.03
	Peritoneal macrophages	ND	3.62 ± 0.26
	Liver	ND	12.2 ± 1.77
	Kidney	ND	15.02 ± 0.09
	Lung	ND	9.65 ±.063
	Fat	ND	10.84 ± 0.55
B6.hALOX12	Islets	7.91 ± 0.18	ND
	Spleen	14.11 ± 0.26	ND
	BMDMs	14.48 ± 0.47	ND
	Peritoneal macrophages	6.06 ± 0.16	ND
	Liver	13.88 ± 0.32	ND
	Kidney	10.12 ± 0.10	ND
	Lung	13.68 ± 0.32	ND
	Fat	8.05 ± 0.42	ND

## Table 1. RNA expression levels of human ALOX12 and mouse Alox15 normalized to mouse Actb from various isolated tissues of C57BL/6J and B6.hALOX12 mice

30 mg/kg VLX-1005 in the peri-STZ treatment period (for the 5 days before, during, and after STZ). STZ-injected C57BL/6J and *B6.hALOX12* mice receiving vehicle became overtly hyperglycemic within 10 days of starting STZ treatment and displayed equivalent glucose intolerance by glucose tolerance test (GTT) (Figure 1, C and D). Upon receiving VLX-1005, however, *B6.hALOX12* mice showed complete protection from STZ-induced diabetes, whereas WT C57BL/6J mice became overtly hyperglycemic (Figure 1E); GTTs at the end of the study confirmed improved glucose tolerance in *B6.hALOX12* mice compared with WT C57BL/6J mice (Figure 1, F and G). These data indicate a specific effect of the drug in preventing hyperglycemia in *B6.hALOX12* mice and support the effectiveness of the *hALOX12* platform for interrogating VLX-1005 action.

Pharmacokinetics of oral VLX-1005 and its effects on STZ-induced diabetes. Given that the oral route is the preferred route for systemic drug delivery in humans, we next asked whether oral administration (p.o.) of VLX-1005 provides adequate exposure in mice. We performed pharmacokinetic analysis following a single p.o. administration (as a suspension in 0.5% methylcellulose) of VLX-1005 spray-dried dispersion at a dose of 30 mg/kg in C57BL/6J mice, followed by serial analysis of VLX-1005 levels by liquid chromatographytandem mass spectrometry (LC-MS/MS). The pharmacokinetic profile of p.o. administered VLX-1005 in mice showed a mean half-life  $(t_{1/2})$  of 3.24 ± 0.07 hours and a consistent  $t_{max}$  of 0.250 hours across all mice. The  $C_{max}$  was 13,300 ± 624 ng/mL, with moderate variability in AUC (15,029 ± 3177 h•ng/mL). These parameters, particularly the low variability in  $t_{max}$  and  $C_{max}$ , support the feasibility of once-daily dosing for maintaining therapeutic levels over a 24-hour period (Table 2). We next tested the effects of p.o. administration of VLX-1005 on the low-dose STZ model, with VLX-1005 (at 30 mg/kg) given 3 days prior to the start of STZ, during STZ, and for 3 days following STZ treatment. Similar to i.p. delivery, p.o. administration of VLX-1005 in B6.hALOX12 mice resulted in lower random-fed blood glucose levels (Figure 1H) and significantly improved glucose tolerance (Figure 1, I and J) compared with vehicle — although this effect was not as robust as with i.p. delivery of the drug. Consistent with improved glucose homeostasis, p.o. VLX-1005-treated mice exhibited greater  $\beta$  cell mass at the end of the study compared with vehicle-treated mice (Figure 1K). Collectively, these data suggest that a single daily p.o. delivery of VLX-1005 (at 30 mg/ kg) achieves plasma levels with therapeutic efficacy.

VLX-1005 treatment reduces  $\beta$  cell inflammation in NOD.hALOX12 mice. The NOD mouse model is a model of T1D that recapitulates many of the immune and  $\beta$  cell features of the disease (20). We, therefore, asked whether pharmacologic inhibition of 12-LOX using p.o. administered VLX-1005 protects against spontaneous diabetes development in the NOD mouse model. To address this question, we introgressed humanized hALOX12 mice onto the NOD background using a speed congenics approach.

	C57BL/6J		NOD.ShiLt/J	
Pharmacokinetic parameter	Mean ± SD	CV (%)	Mean ± SD	CV (%)
t <sub>1/2</sub> (h)	3.24 ± 0.07	2.17	2.53 ± 0.41	16.1
$t_{\rm max}$ (h)	0.250 ± 0.00	0.00	0.250 ± 0.00	0.000
C <sub>max</sub> (ng/mL)	13,300 ± 624	4.70	14,253 ± 5474	38.4
AUC <sub>last</sub> (h•ng/mL)	15,029 ± 3177	21.1	13,211 ± 1631	12.3
AUC <sub>inf</sub> (h•ng/mL)	15,083 ± 3206	21.3	13,225 ± 1625	12.3
AUC <sub>%extrap_obs</sub> (%)	0.342 ± 0.133	38.9	0.115 ± 0.099	85.6
MRT <sub>inf obs</sub> (h)	2.57 ± 0.38	14.8	3.81 ± 0.71	18.7
AUC <sub>last</sub> /D (h•ng/mL)	501 ± 106	21.1	440 ± 54	12.3

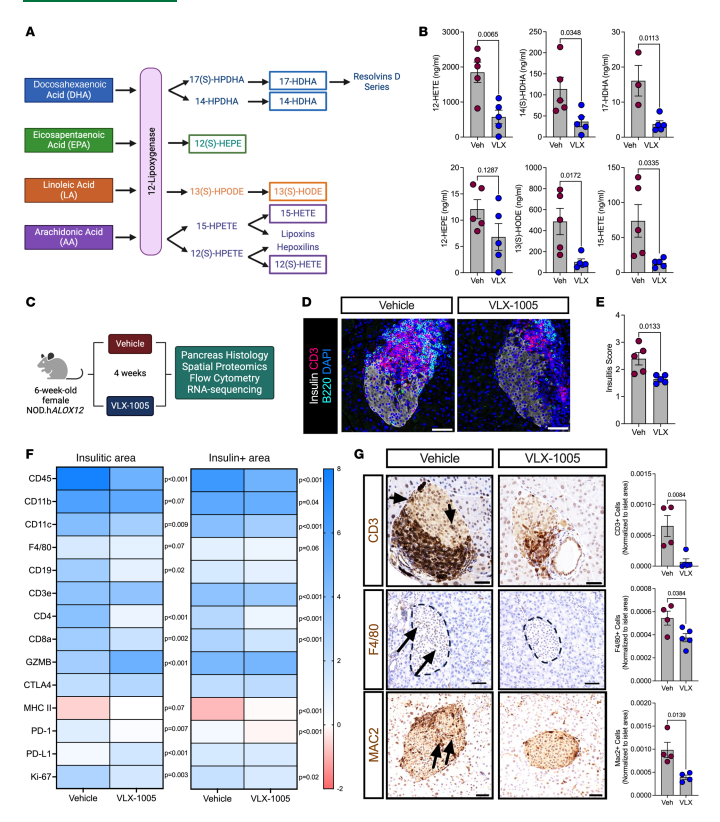
#### Table 2. Plasma concentration vs. time profile for VLX-1005 after 30 mg/kg p.o. in C57BL/6J and NOD.ShiLt/J mice.

 $t_{_{1/2}}$ , half-life;  $t_{_{max}}$ , time to maximum drug concentration;  $C_{_{max}}$ , maximum drug concentration; AUC<sub>\_{last</sub>, AUC from the time of dosing to the last measurable concentration; AUC<sub>\_{int</sub>, AUC from the time of dosing extrapolated to infinity; AUC<sub>\_{sectrap\_obs</sub></sub>, AUC from the time of dosing extrapolated to last observed concentration; MRT<sub>\_{inf\_obs</sub></sub>, mean residence time from the time of dosing extrapolated to infinity; AUC<sub>\_{sectrap\_obs</sub></sub>, D, dose-normalized AUC to time of last measurable concentration.

Genome scanning of microsatellites was performed to confirm that mice were 100% congenic on the NOD mouse background (*NOD.hALOX12* mice) (Supplemental Table 1). We next measured human 12-LOX protein levels in the *NOD.hALOX12* mice. Similar to the gene profile we observed in *B6.hALOX12* mice compared with WT C57BL/6J mice, WT NOD tissues robustly expressed mouse 12/15-LOX (the protein encoded by mouse *Alox15*) and little or no human *ALOX12*; conversely, *NOD.hALOX12* mice tissues robustly expressed human 12-LOX and minimal levels of 12/15-LOX (Supplemental Figure 1, J and K). Consistent with their congenic nature, female *NOD.hALOX12* mice exhibited islet pathology similar to that of NOD mice at the (prediabetic) age of 10 weeks, with evidence of T and B cell infiltration of islets (Supplemental Figure 1L) and indistinguishable insulitis score (Supplemental Figure 1M), suggesting that replacement of *Alox15* with human *ALOX12* did not alter the islet pathology of the disease. Pharmacokinetics of p.o. administered VLX-1005 (30 mg/kg) on the NOD background were similar to those seen in C57BL/6J mice (Table 2), suggesting that the NOD background does not affect drug absorption or clearance.

To assess the effect of VLX-1005 administration on products of 12-LOX activity in *NOD.hALOX12* mice, we administered 30 mg/kg VLX-1005 (or vehicle) p.o. to female *NOD.hALOX12* mice for 1 week during the prediabetic phase (8 weeks of age) and harvested serum. Lipidomics analysis (by LC-MS/MS) was performed for a series of 12-LOX products resulting from different fatty acid substrates (Figure 2A). Notably, levels of 12-HETE (from arachidonic acid), 13-HODE (from linoleic acid), and 14-HDHA and 17-HDHA (from docosahexaenoic acid) were all significantly reduced (Figure 2B). Levels of 12-HEPE (from eicosapentaenoic acid) were not significantly changed (Figure 2B), suggesting minimal involvement of eicosapentaenoic acid metabolism in NOD mice. Lipids within the pathway that are processed by enzymes other than 12-LOX were not statistically significantly altered (Supplemental Table 2). These data are collectively consistent with the expected 12-LOX engagement by VLX-1005.

To assess the effect of VLX-1005 administration on immune cell phenotypes in *NOD.hALOX12* mice, we administered 30 mg/kg VLX-1005 (or vehicle) p.o. to female *NOD.hALOX12* mice for 4 weeks during the prediabetic phase (6–10 weeks of age) and harvested pancreas, pancreatic lymph nodes (pLNs), and spleen (Figure 2C). Pancreas pathology showed reduced T and B cell infiltration and that the extent of insulitis (by insulitis scoring) was significantly reduced in VLX-1005–treated *NOD.hALOX12* mice compared with vehicle-treated mice (Figure 2, D and E). To specifically interrogate the nature of immune cells within the insulitic region, we performed spatial tissue-based proteomics (NanoString). We used insulin immunostaining and nuclei staining to identify  $\beta$  cells and the surrounding insulitic regions. Prevalidated antibodies in the GeoMx mouse immune panel were used to probe for immune cell subtypes in the peri-islet insulitic region and within the islet. *NOD.hALOX12* mice exhibited a notable reduction in myeloid population subtypes in both insulitic and islet areas, including macrophages (F4/80<sup>+</sup>CD11b<sup>+</sup>) and dendritic cells (CD11c<sup>+</sup>) (Figure 2F and Supplemental Figure 2, A and B). This reduction in myeloid cell populations was accompanied by a decrease in T and B cells populations, including CD4<sup>+</sup>, CD3<sup>+</sup>, CD8<sup>+</sup>, and CD19<sup>+</sup> cells (Figure 2F and Supplemental Figure 2, A and B). Immunohistochemistry of pancreas sections confirmed the

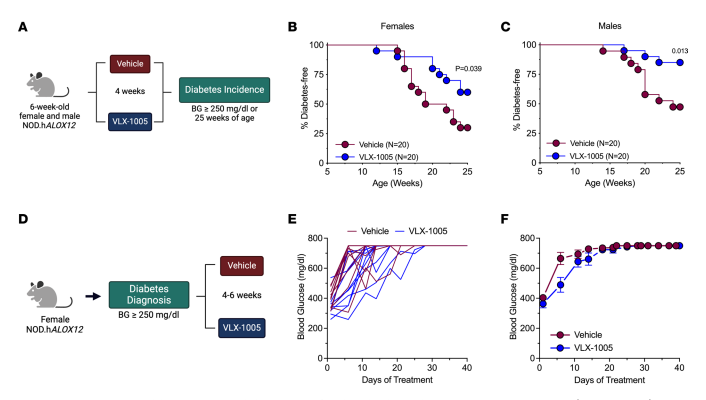


**Figure 2. VLX-1005 decreased islet inflammation in NOD.hALOX12 female mice.** Six-week-old female prediabetic NOD.hALOX12 mice were treated p.o. with 30 mg/kg VLX-1005 for 4 weeks prior to tissue analysis. (**A**) Schematic representation of 12-lipoxygenase products. (**B**) Serum lipidomics results of 12-lipoxygenase products as indicated (*n* = 4–5). (**C**) Schematic representation of mouse treatment paradigm. (**D**) Pancreata from mice stained for CD3 (magenta), B220 (teal), insulin (white), and nuclei (blue). Scale bars: 50 μm. (**E**) Average insulitis score; each dot represents an individual mouse (*n* = 4–5). (**F**) Heatmap of identified proteins in the insulitic area (left) and insulin-positive area (right). (**G**) Pancreata of mice stained and quantified for CD3 (brown, top panels: arrows indicate positive CD3 staining within the islet), F4/80 (brown, middle panels: arrows indicate positive F4/80 staining within the islet), or MAC2 (brown, bottom panels: arrows indicate positive MAC2 staining within the islet) and nuclei (blue). Each dot represents an individual mouse (*n* = 4–5). Scale bars: 50 μm. Data are presented as mean ± SEM and statistical significance was determined by a 2-tailed Student's *t* test in all cases.

reductions in both T cells (CD3<sup>+</sup>), macrophages (F4/80<sup>+</sup>), and activated macrophages (Mac2<sup>+</sup>) following p.o. VLX-1005 treatment (Figure 2G). A notable observation in spatial proteomics was the increased levels of the immune checkpoint ligand PD-L1 (Figure 2F). Enhanced PD-L1/PD-1 interactions shift T cells to less aggressive, more regulatory phenotypes (21). To interrogate this possibility, we performed immune profiling by flow cytometry of pLNs from mice treated with p.o. VLX-1005 or vehicle. pLNs are key sites in the initial priming of autoreactive T cells in NOD mice (22). Treatment with p.o. VLX-1005 led to an increase in CD4<sup>+</sup>Foxp3<sup>+</sup> regulatory T cells (Tregs) in the pLNs (Supplemental Figure 2C). This effect on Tregs was specific for the pLNs since no changes in Tregs were observed in the spleen after VLX-1005 treatment (Supplemental Figure 2D).

Orally administered VLX-1005 reduces autoimmune diabetes incidence in female and male NOD.hALOX12 mice. Because 4 weeks of p.o. VLX-1005 dosing led to improvements in insulitis and reductions in infiltrating T and B cells, we next asked whether these changes lead to prevention or delay of subsequent diabetes development in NOD.hALOX12 mice. Both female and male mice were administered VLX-1005 via daily oral gavage (30 mg/kg) or vehicle for 4 weeks during the prediabetic phase (6-10 weeks of age). Mice were followed for diabetes development (blood glucose  $\geq 250 \text{ mg/dL}$  on 2 consecutive days) until 25 weeks of age (Figure 3A). At 25 weeks of age, 60% of female mice and 75% of male mice receiving VLX-1005 were protected from diabetes development compared with 25% of female and 50% of male mice receiving vehicle (Figure 3, B and C). Whereas the preceding studies demonstrate that 12-LOX inhibition with p.o. VLX-1005 delays the development of diabetes, they do not address whether administration of the drug might reverse established diabetes or mitigate hyperglycemia. We allowed female NOD.hALOX12 mice to develop diabetes (defined as 2 consecutive random-fed blood glucose measurements  $\geq$ 250 mg/dL), and then administered VLX-1005 or vehicle for up to 6 weeks via daily oral gavage or until the mice exhibited signs of physical deterioration from hyperglycemia (loss in body weight, dishevelment) (Figure 3D). Notably, we did not observe a reversal in diabetes, but did see relative reductions in blood glucose levels in mice treated with VLX-1005 compared with vehicle (Figure 3, E and F).

Orally administered VLX-1005 reduces islet death and oxidative stress in NOD.hALOX12 mice. 12-LOX is primarily present in islets and macrophages, and deletion of the mouse gene (Alox15) in either tissue separately was previously shown to reduce diabetes incidence. We, therefore, first asked how treatment with VLX-1005 affects islet cell phenotypes. We first analyzed isolated islets from female NOD.hALOX12 mice treated with vehicle or VLX-1005 by RNA sequencing to identify how islet gene expression might be altered. Principal component analysis of transcriptomics revealed that islets from vehicle- and VLX-1005-treated NOD.hALOX12 mice clustered separately, suggesting an effect of VLX-1005 treatment on gene expression (Figure 4A). Pairwise comparison of gene expression using a false discovery rate (FDR) of less than 0.05 and fold-change (FC) of greater than 2 yielded only 189 differentially expressed genes. Instead, a P-value cutoff of 0.05 and FC greater than 2 revealed changes in 709 genes between vehicle- and VLX-1005-treated NOD.hALOX12 mice (Figure 4B, volcano plot, and Supplemental Table 3 for full sequencing results). Gene Ontology (GO) pathway analysis showed significantly altered pathways related to DNA replication (e.g., Anp32b, Skp1a, Itfg2, Dmrt1i), inflammation (NF-KB activity; e.g., Elf1, Trim75, Rnase1, Lmo1, Bcl3, Ptgis, Commd1, Lrrc14, Foxp3), and G protein-coupled receptor signaling (e.g., Gpr89, Glp2r), among others (Figure 4C). These pathways suggest responses that may be related to changes in cellular survival in response to VLX-1005. We, therefore, immunostained pancreatic sections for markers of cell death and proliferation in the islet. VLX-1005-treated NOD.hALOX12 mice exhibited decreased islet cell death, as measured by reduced terminal deoxynucleotidyl transferase dUTP nick end labeling (TUNEL) and H2A histone family member X (H2A.X) staining compared with vehicle-treated mice (Figure 4D). Additionally, VLX-1005-treated mice demonstrated decreased  $\beta$  cell proliferation, as measured by proliferating cell nuclear antigen (PCNA) immunostaining (Figure 4D); reduced PCNA immunostaining was also consistent with the reduction in Ki-67 observed in spatial proteomics of the insulin<sup>+</sup> area (Figure 2F and Supplemental Figure 2B). We interpret the reduction in  $\beta$  cell proliferation as a consequence of reduced  $\beta$  cell apoptosis. The change in NF- $\kappa$ B signaling led us to investigate whether markers of oxidative stress were affected since inflammation, oxidative stress, and  $\beta$  cell survival are closely linked (23, 24). We performed immunostaining for the oxidative stress marker, 4-hydroxynonenal (4-HNE), and observed reduced immunostaining in mice treated with VLX-1005 compared with placebo (Figure 4E). Consistent with this observation,  $\beta$  cells from VLX-1005-treated animals also displayed an increase in levels of the antioxidant enzyme GPx1 (Figure 4E). Collectively, these data are consistent with prior observed effects of reduced inflammation, oxidative stress, and  $\beta$  cell death in  $\beta$  cell–specific deletion of mouse *Alox15* in NOD mice (11).



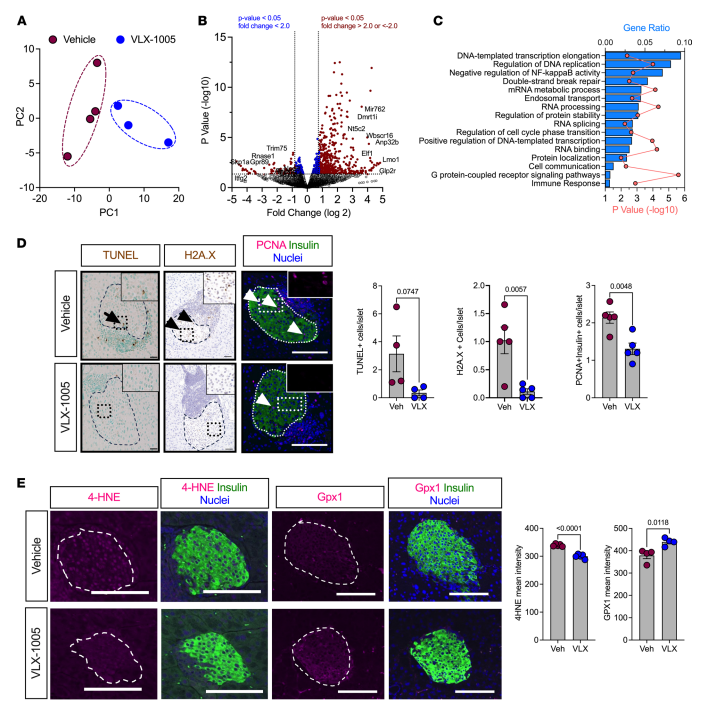
**Figure 3. VLX-1005 treatment delays autoimmune diabetes onset in female and male** *NOD.hALOX12* mice. *NOD.hALOX12* mice (*n* = 20 per group) were treated during the prediabetic stage from 6 to 10 weeks of age or at the time of diabetes development (*n* = 11-12 per group). (**A**) Schematic representation of diabetes prevention experimental design. (**B**) Diabetes incidence in female *NOD.hALOX12* mice. (**C**) Diabetes incidence in male *NOD.hALOX12* mice. (**D**) Schematic representation of diabetes reversal experimental design. (**E**) Random-fed blood glucose levels in each female mouse. (**F**) Average random-fed blood glucose levels of female mice. Data are presented as mean ± SEM and statistical significance was determined by a Mantel-Cox log-rank test.

VLX-1005 alters the proinflammatory macrophage phenotype. Whereas the preceding findings are consistent with improved  $\beta$  cell survival following p.o. VLX-1005 administration, these studies do not rule out the possibility that the drug directly modifies the phenotype of infiltrating macrophages, which could secondarily affect  $\beta$  cells. Because bulk islet transcriptomics analysis does not resolve gene expression events associated with specific cell types, we isolated BMDMs from female NOD.hALOX12 mice and then performed RNA sequencing in the presence or absence of VLX-1005. BMDMs were unpolarized (M0) or polarized to an M1-like state (with lipopolysaccharide and IFN- $\gamma$ ) to mimic the inflammatory state that might be observed during T1D pathogenesis. During polarization, BMDMs were treated with vehicle or VLX-1005 (Figure 5A). Principal component analysis of transcriptomics revealed that M0 macrophages treated with VLX-1005 co-clustered with vehicle-treated M0 macrophages, suggesting a minimal transcriptional effect of the drug on unpolarized cells (Figure 5B). Consistent with this interpretation, only 1% of genes (159 out of 15,888) were significantly altered with VLX-1005 treatment (when using criteria FC > 2 and P < 0.05) (Supplemental Table 4 for full sequencing results). Upon polarization to the M1-like state, a clear rightward shift in the principal component analysis plot was observed with both vehicle- and VLX-1005-treated BMDMs and a notable separation was seen between vehicle and VLX-1005 treatment (Figure 5B); this finding suggests that the impact of 12-LOX inhibition is more prominent upon a shift to a proinflammatory state of macrophages.

We next interrogated the gene expression events associated with the M0 to M1 transition in both vehicle- and VLX-1005–treated BMDMs. Most genes altered (FC > 2 and P < 0.05) in this transition (2467 genes) were common between vehicle and VLX-1005 treatment (Figure 5, C and D, Supplemental Figure 3A, and Supplemental Table 4). These common genes mapped to pathways related to cytokine-mediated signaling, T cell activation, and antigen processing and presentation (Supplemental Figure 3B). An additional 507 genes were significantly altered in vehicle-treated cells, and 459 additional genes were significantly altered with VLX-1005 treatment (Supplemental Figure 3A). The 507 genes altered with vehicle treatment mapped to GO pathways related to the M1 polarization phenotype (myeloid cell differentiation, immune response, response to cytokines) and pathways related to oxidative stress (cell redox homeostasis, response to hydrogen peroxides) (Figure 5E). These pathways were not identified in the genes that were

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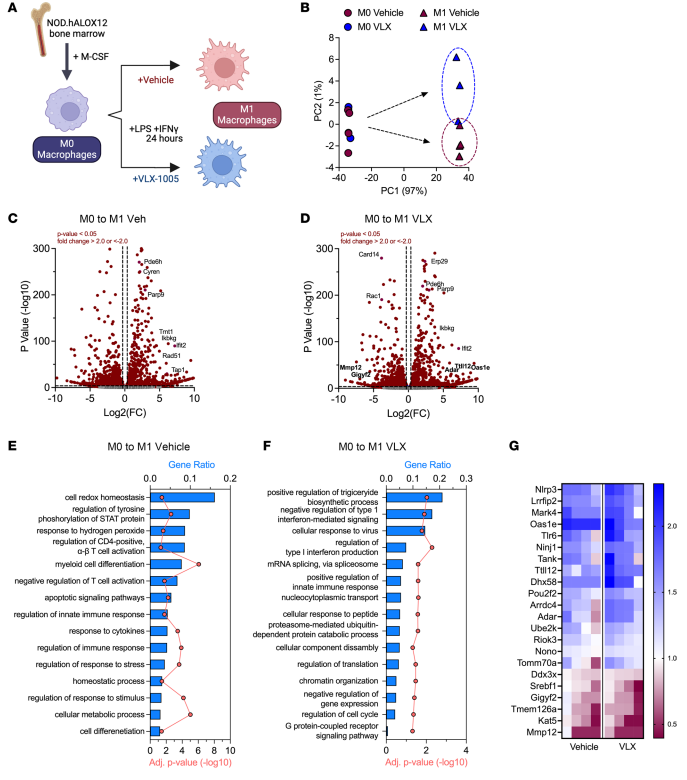
### **JCI** insight



**Figure 4. VLX-1005 decreased** β **cell death, proliferation, and oxidative stress in female NOD.hALOX12 mice.** Pancreata or islets were harvested from 10-week-old prediabetic female *NOD.hALOX12* mice after 4 weeks of treatment with vehicle or VLX-1005 (*n* = 3–4 per group). (**A**) Principal component analysis plot of RNA sequencing results from isolated islets of vehicle- or VLX-1005–treated mice. (**B**) Volcano plot of differentially expressed genes. (**C**) Gene ontology pathway analysis of differentially expressed genes. (**D**) Pancreata from mice stained and quantified for TUNEL (brown, left panels: black arrow indicates positive TUNEL staining within the islet), H2A.X (brown, middle panels: black arrowheads indicate positive H2A.X staining within the islet), or PCNA (magenta, right panels: white arrowheads indicate positive PCNA staining within the islet), insulin (green) and nuclei (blue). Each dot represents an individual mouse (*n* = 4). Scale bars: 50 µm. Data are presented as mean ± SEM and statistical significance was determined by a 2-tailed Student's *t* test.

differentially expressed during VLX-1005 treatment. GO pathway analysis of the 459 genes altered with VLX-1005 showed particularly significant changes in pathways related to modification of the IFN response (Figure 5F). Next, we looked specifically at the genes that mapped to IFN pathways (Figure 5G), which

## **JCI** insight



**Figure 5. RNA sequencing analysis of M1-like BMDMs reveals a reduction in the inflammatory response upon VLX-1005 treatment.** Bone marrowderived macrophages (BMDMs) were isolated and polarized to the M1-like state and treated with vehicle or VLX-1005 (10  $\mu$ M) during polarization. RNA was isolated and sequenced (*n* = 4 per group). (**A**) Schematic of experimental design. (**B**) Principal component analysis plot. (**C**) Volcano plot of differentially expressed genes in M0 and M1-like vehicle-treated macrophages. (**D**) Volcano plot of differentially expressed genes in M0 and M1-like VLX-1005– treated macrophages. (**E**) Gene ontology pathway analysis of differentially expressed genes in M0 vs. M1-like vehicle-treated macrophages. (**F**) Gene ontology pathway analysis of differentially expressed genes in M0 vs. M1-like VLX-1005–treated macrophages. (**G**) Heatmap of significantly altered IFN-related genes. Columns represent sequencing results from each sample (*n* = 4 per group). Numbers on the heatmap scale indicate fold change (FC) compared with M0 macrophages. revealed that VLX-1005 augmented significantly (P < 0.05 by t test) the magnitude of the gene changes compared with vehicle treatment. Notable genes whose directional changes are known to counter the IFN response included *Oas1e* (25), *Ttll12* (26), and *Adar* (27) (all upregulated compared with control-treated M1 macrophages), and *Gigyf2* (28) and *Mmp12* (29) (correspondingly downregulated) (Figure 5G). These data suggest effects of VLX-1005 that may lead to reduced macrophage IFN signaling.

#### Discussion

To date, the adaptive immune system has remained the primary focus for the development of therapeutics aimed at preventing or reversing T1D. Notwithstanding the utility of agents such as anti-CD3 monoclonal antibodies in preserving  $\beta$  cell function (30) or delaying T1D development (7), there has been impetus in the research community to develop therapeutics that target other cell types that contribute to T1D development (31), including innate immune cells and  $\beta$  cells. A multi-targeted approach is expected to aid in better disease modification and result in more durable and broadly applicable therapy (32). In this respect, 12/15-LOX (in mice) is a particularly appealing target since it is active in both macrophages and  $\beta$  cells and contributes to the development of inflammatory disorders, including insulin resistance, atherosclerosis, and T1D (for review, see ref. 9). The deletion of *Alox15*, specifically in either myeloid cells or islet  $\beta$  cells, proved sufficient to delay/prevent T1D in NOD mice (2, 11). Evidence for 12-LOX contributions to T1D pathogenesis identifies this enzyme as an attractive target in human disease. 12-LOX is elevated in  $\beta$  cells of autoantibody-positive, pre-T1D individuals and in residual  $\beta$  cells of individuals with established T1D (12). A proinflammatory product derived from the 12-LOX-mediated (and mouse 12/15-LOX-mediated) metabolism of arachidonic acid is 12-HETE, an eicosanoid that either directly or indirectly (through G protein-coupled receptors; refs. 33-36) augments reactive oxygen species generation and ER stress in macrophages and islets (37, 38). Notably, levels of 12-HETE were shown to be elevated in the circulation of youths and adults with new-onset T1D (compared with healthy controls and those with established T1D) (39).

Considering the biological contributions of the 12-LOX enzymes (mouse 12/15-LOX and human 12-LOX) to T1D and other inflammatory disorders, the development of enzyme inhibitors offers an attractive approach to disease modification. Inhibition of 12/15-LOX using ML351 demonstrated promising outcomes in NOD mice, with reductions in insulitis and improvements in glucose homeostasis (13). An inhibitor that showed mouse and human cross-species reactivity, ML127, unfortunately also displayed evidence of off-target cytotoxicity (13). By contrast, VLX-1005 (also known as ML355) is a potent inhibitor  $(IC_{so} \sim 300 \text{ nM})$  of human 12-LOX (14) that has shown efficacy in reversing the insulin secretory defects of cytokine-treated human islets in vitro without evidence of cytotoxicity (17); however, the benefit of this inhibitor in disease states in vivo has remained speculative. To address the translational challenge of testing 12-LOX inhibitors in preclinical disease models in vivo, we developed a human gene replacement mouse model on both the C57BL/6J and NOD autoimmune diabetes backgrounds. The utility of this model as a platform for 12-LOX inhibitor testing was confirmed in our multiple low-dose STZ studies, which showed that VLX-1005 administration to B6.hALOX12 mice precluded hyperglycemia, whereas WT controls developed hyperglycemia over time. STZ is a toxin whose full effects involve communication between  $\beta$  cells and macrophages (19), and our findings with systemic administration of VLX-1005 are consistent with similar STZ studies in mice harboring the global deletion of Alox15 (18) — collectively suggesting that human ALOX12 gene replacement mice respond appropriately to a 12-LOX inhibitor and that the human ALOX12 gene can sufficiently replace the functionality of the mouse Alox15 gene.

In recent studies, our group showed that loss of the mouse *Alox15* gene in either myeloid cells or islet  $\beta$  cells could protect animals from the development of autoimmune diabetes on the NOD background (2, 11). The pathological phenotypes of these animals were similar, with reductions in islet invasion by T cells, B cells, and myeloid cells and a characteristic increase in PD-L1 in either macrophages or  $\beta$  cells. PD-L1 is an immune checkpoint protein whose interaction with its receptor PD-1 on adaptive immune cells leads to a more immunosuppressive response (21). In our studies using *NOD.ALOX12* mice, we found similar responses to VLX-1005 treatment, with striking reductions in both innate and adaptive immune cell infiltration into the islets and a notable increase in PD-L1 in the insulitic (immune cell) component. We interpret these latter findings to suggest that the effect of systemically administered VLX-1005 may be greater in macrophages than in  $\beta$  cells. The reductions in markers of  $\beta$  cell proliferation, death, and oxidative stress that we observed with VLX-1005 treatment may also be reflective of this preferential effect on macrophages since these responses are otherwise characteristic of the effects of cytokines on  $\beta$  cells (40). Our RNA sequencing studies of isolated NOD mouse BMDMs polarized

to the proinflammatory M1 state support this contention, as VLX-1005 treatment resulted in reductions in the IFN response. Our studies open the possibility that VLX-1005 may be useful in autoimmune diabetes resulting from PD-L1 or PD-1 blockade (checkpoint inhibitor therapy) often used to treat cancers (41).

Some limitations to our study should be acknowledged. First, because our mouse model replaces the mouse Alox15 with the human ALOX12 globally, we cannot be certain that the effects we observed are exclusively related to the inhibition of the human enzyme in only macrophages and  $\beta$  cells; a recent study suggested that mouse *Alox15* may contribute to pro-resolving functions of Tregs (42). It remains unclear, therefore, whether the loss of Alox15 globally in our mouse model might have affected Treg function or the function of other cell types that have low levels of *Alox15* expression. In this respect, our mouse model may have limited utility in other disease states where ALOX12 might not fully replace the function of Alox15. Moreover, we cannot rule out the possibility that *Alox15* and *ALOX12* mediate distinct pathways that nevertheless still allow for shared phenotypes. For example, our RNA sequencing studies of macrophages treated with VLX-1005 reveal distinct reductions in IFN-mediated responses, but such effects have not been directly explored as a consequence of Alox15 in mice. Second, our studies do not fully address the timing and duration of VLX-1005 treatment. We only treated mice for a 4-week period (6-10 weeks of age); it is possible that a longer duration of treatment might have yielded even more robust T1D prevention outcomes. Therefore, the timing and duration of human treatment may require further investigation in our preclinical model. Limitations regarding timing and duration may also apply to diabetes reversal studies, in which we observed a reduction in glycemia but no reversal in disease. We cannot rule out the possibility that the effect of VLX-1005 to suppress  $\beta$  cell proliferation may have prevented a more robust outcome in this case. Finally, our study does not assess the potential for 12-LOX inhibition in combination with other immunomodulatory agents. Whereas the relatively modest impact of VLX-1005 treatment in the diabetes reversal studies in NOD.ALOX12 mice might suggest diminishing returns on the treatment of humans at the time of disease diagnosis, this effect may be amplified in the presence of T or B cell blockade.

Despite these limitations, our studies identify a platform on which to study a class of LOX inhibitors for their utility in ameliorating human autoimmune diabetes. Our human gene replacement mouse model demonstrates a functional equivalence between mouse *Alox15* and human *ALOX12* in the context of T1D since the whole-body replacement of the mouse gene with the human (under the mouse upstream control elements) preserves islet pathology and the frequency of diabetes incidence in NOD mice. Therefore, beyond its utility to test inhibitors of human 12-LOX, our mouse model also provides a platform to interrogate the cause-and-effect relationship of human 12-LOX in T1D and possibly other inflammatory diseases in vivo.

#### Methods

Sex as a biological variable. For C57BL/6J mice, our study examined only male mice, because comparative data for these mice in the literature are primarily from males. For NOD mice, our study mostly examined females, because T1D in the NOD strain is more frequently observed in females. However, some data from NOD male mice are included that parallel those seen in females, suggesting that the effects observed in females may be relevant to males.

Animals. Male and female C57BL/6J mice and NOD/ShiLTJ mice were procured from The Jackson Laboratory. All mice were kept under pathogen-free housing conditions with standard 12-hour light/dark cycles and fed ad lib normal chow. To generate a humanized *ALOX12* mouse model, the coding region of the mouse *Alox15* gene was replaced with the coding region of the human *ALOX12* gene while retaining all the mouse regulatory elements (Figure 1A) (mice were generated by a contract to Ingenious Targeting Laboratory). Targeted iTL BF1 (C57BL/6 FLP) embryonic stem cells were microinjected into BALB/c blastocysts. Chimeras with high percentage of black coat color resulting from this procedure were then mated to WT C57BL/6J mice to generate germline neo-deleted mice. The following primers were used to genotype the mice: 5'-TCTGATCTGTGTATGCCTGTGTGTGTGG-3' (forward) and 5'-TTCCAAGGAAAAAGG-CATGGTTTCTGAGG-3' (reverse). These primers generate a 478 bp band for WT and 581 bp band for knockin mice (Supplemental Figure 1A).

Human *ALOX12* alleles were introgressed onto both the C57BL/6J and *NOD.ShiLT/J* mouse backgrounds using a speed congenics approach based on microsatellite genotyping at The Jackson Laboratory. Genome scanning was also performed at The Jackson Laboratory to confirm successful backcrossing onto the C57BL/6J and *NOD.ShiLT/J* mouse background (*B6.hALOX12* and *NOD.hALOX12*; Supplemental Table 1). Body mass was measured by EchoMRI. Intraperitoneal GTTs (IPGTTs) were performed in mice after overnight fasting (16 hours). Mice were i.p. injected with glucose at a dose of 1 or 2 g/kg body weight and blood glucose levels were measured at specific time points: 0, 10, 20, 30, 60, 90, and 120 minutes after glucose injection using an AlphaTrak glucometer.

*Formulation of VLX-1005.* VLX-1005 was obtained from Veralox Therapeutics Inc. A VLX-1005 spraydried dispersion (for p.o. administration) was prepared by dissolving VLX-1005 and HPMC-E3 in a 90:10 w/w mixture of tetrahydrofuran and water to attain a total solids concentration of 5% w/w; 1575 g of solution was then spray dried using a Buchi B-290 laboratory spray dryer. The yield after spray drying was 67.8 g. The collected material was further dried in an oven at 40°C under vacuum to remove residual tetrahydrofuran.

*Pharmacokinetic analysis and lipidomics*. Following i.p. injection or oral gavage (p.o.), VLX-1005 was quantified in plasma using high-performance LC-MS/MS (Triple Quad 6500+; Sciex) after separation by HPLC (column: Agilent Poroshell 120 EC-C18; HPLC: Shimadzu DGU-405). Pharmacokinetic parameters for VLX-1005 were estimated by noncompartmental model using WinNonlin 8.3 (https://www.certara. com/software/phoenix-winnonlin/). The bioavailability (*F*%) was calculated as the following: AUC<sub>last</sub>-p.o./ AUC<sub>inf</sub>-p.o. > 80%:  $F = (AUC_{inf}$ -p.o. × Dose-i.v.)/(mean AUC<sub>inf</sub>-i.v. × Dose-p.o.), where AUC<sub>last</sub> is the AUC from the time of dosing to the last measurable concentration and AUC<sub>inf</sub> is the AUC from the time of dosing extrapolated to infinity. Lipidomics on serum samples was performed by the New York Medical College Lipidomics Core using a Shimadzu LC-MS/MS 8050 system equipped with a UHPLC and autosampler.

*STZ induction*. Male C57BL/6J and *B6.hALOX12* mice (8–10 weeks of age) were injected with either vehicle (0.5% methylcellulose) or 30 mg/kg/day VLX-1005 by i.p. injection for 15 days: 5 days prior to the start of multiple low-dose STZ (55 mg/kg/day; 5 consecutive days), 5 days during STZ treatment, and 5 days after STZ injections. Male *B6.hALOX12* mice (8–10 weeks of age) were injected with either vehicle (0.5% methylcellulose) or 30 mg/kg/day VLX-1005 p.o. for 11 days: 3 days prior to the start of multiple low dose STZ (55 mg/kg/day; 5 consecutive days), 5 days during STZ treatment, and 3 days after STZ injections. Random-fed glucose levels were measured by tail snip using a glucometer (AlphaTrak), and mice were followed for 20 days after STZ injections. IPGTT was performed on day 4 after STZ treatments after overnight fasting. At the end of each study, mice were euthanized, and pancreas and blood samples were collected.

*Diabetes incidence and treatment*. Both male and female *NOD.hALOX12* mice were given either vehicle or 30 mg/kg/day VLX-1005 (p.o.) for 4 weeks in the prediabetic stage (6–10 weeks of age) and then followed for diabetes incidence until 25 weeks of age or until diabetes diagnosis. Diabetes incidence was determined by observing 2 consecutive blood glucose values greater than 250 mg/dL. At the end of each study, mice were euthanized, and pancreas and blood samples were collected.

For diabetes treatment studies, female *NOD.hALOX12* mice were followed for random-fed blood glucose from 12 to 20 weeks of age. At diabetes incidence (2 consecutive blood glucose values greater than 250 mg/dL), mice were administered 30 mg/kg/day VLX-1005 SDD or vehicle for up to 6 weeks via daily oral gavage or until the mice exhibited signs of physical deterioration from hyperglycemia (loss in body weight, dishevelment). At the end of each study, mice were euthanized, and pancreas and blood samples were collected.

Islet and macrophage isolation. Islets were isolated from NOD.hALOX12 mice with either vehicle or 30 mg/kg/day of VLX-1005 treatment using collagenase digestion. Briefly, collagenase was injected into the pancreatic bile duct to digest the connective tissue and release pancreatic cells (43). A Histopaque-HBSS gradient was applied to the dissociated pancreas and centrifuged at 900g for 18 minutes. The isolated islets were cultured in RPMI 1640 medium. The collected islets were handpicked and allowed to recover overnight before processing. RNA was isolated for use in RNA sequencing or quantitative PCR.

BMDMs were isolated from 8-week-old *NOD.hALOX12* mice as described previously (2). The isolated BMDMs were cultured for 7 days in complete medium (RPMI 1640 containing 10% FBS, 10 mM HEPES, and 100 U/mL penicillin/streptomycin) supplemented with 10 ng/mL M-CSF. On day 7 of culture, the BMDMs were pretreated with either vehicle (0.1% DMSO) or 10  $\mu$ M VLX-1005. After 1-hour pretreatment, the BMDMs were further stimulated with 10 ng/mL LPS and 25 ng/mL IFN- $\gamma$  for 18 hours for M1-like polarization. RNA was isolated and used for sequencing.

*RNA isolation and quantitative PCR*. RNA was isolated from mouse tissues and macrophages using an RNeasy Mini Kit from Qiagen. The isolated RNA was used to synthesize cDNA using a High-Capacity cDNA Reverse Transcription kit (Applied Biosystems) according to manufacturer's instructions. Quantitative PCR was performed using a Bio-Rad CFX Opus with a predesigned TaqMan assay probe for human and mouse genes: human *ALOX12*: Hs00167524\_m1; mouse *Alox15*: Mm00507789\_m1; mouse *Actb*:

Mm01205647\_m1 (Invitrogen). The relative gene expression levels were calculated using the comparative threshold cycle value (Ct) and normalized to *Actb*.

*Immunostaining,*  $\beta$  *cell mass, and insulitis scoring.* Pancreatic tissues were fixed using 4% paraformaldehyde. After fixation, the tissues were embedded in paraffin and sectioned with a thickness of 5 µm. Three sections per mouse were used for analysis, with each section being spaced 100 µm apart. Tissue sections were immunostained with anti-insulin (ProteinTech, 15848-1-AP; 1:200), anti-glucagon (Abcam, ab92517; 1:200), anti-12-LOX (Thermo Fisher Scientific, PA5-26020; 1:200), anti-12/15-LOX (Abcam, ab80221; 1:200), anti-CD3 (Abcam, ab16669; 1:200), anti-F4/80 (Sigma-Aldrich, D2S9R; 1:150), anti-MAC2 (Thermo Fisher Scientific, EbioM3/38; 1:200), and anti-H2A.X (Cell Signaling Technology, 9718s; 1:200) primary antibodies followed by conjugated anti-rabbit Ig (Vector Laboratories) secondary antibody. A DAB (3,3'-diaminobenzedine) Peroxidase Substrate Kit from Vector Laboratories was used for detection. After immunostaining, the tissue sections were counterstained with hematoxylin (Sigma-Aldrich). Images were acquired using a BZ-X810 fluorescence microscope (Keyence) and  $\beta/\alpha$  cell mass was quantified by insulin<sup>+</sup> or glucagon<sup>+</sup> area and whole pancreas area. Insulitis score reflects the degree of immune cell infiltration within pancreatic islets. The score system used as follows: 1 = no insulitis, 2 = infiltrate <50% circumference, 3 = infiltrate >50% circumference, 4 = infiltration within islet. Data are shown as the average insulitis score per mouse.

For immunofluorescent staining, pancreatic sections were stained with the following antibodies: anti-insulin (Dako, IR002; 1:4), anti-glucagon (Santa Cruz Biotechnology, sc514592; 1:50), anti-B220 (BioLegend, 03201; 1:100), anti-CD3 (Abcam, ab16669; 1:200), anti-PCNA (Santa Cruz Biotechnology, sc-7907; 1:100), anti-4-HNE (Abcam, ab46545; 1:200), and anti-GPx1 (Santa Cruz Biotechnology, sc-22145; 1:100). Highly cross-adsorbed Alexa Fluor secondary antibodies (Thermo Fisher Scientific) were used at a dilution of 1:500. Tissue sections were stained with DAPI (Thermo Fisher Scientific) to label cell nuclei. The Nikon A1 confocal microscopy was used to capture images. CellProfiler v4.1 software (https:// cellprofiler.org/) was used for image analysis.

*TUNEL staining.* A TUNEL assay was used to determine  $\beta$  cell death in pancreatic islets. The assay was performed according to the protocol provided by the manufacturer (Abcam) and HRP-DAB chemistry was used for detection. Two sections, spaced 100  $\mu$ m apart, were used for each mouse. Images were captured using a BZ-X810 fluorescence microscope system (Keyence). The number of TUNEL<sup>+</sup> cells was assessed manually per islet.

*NanoString spatial proteomics.* Paraffin embedded pancreata were used for NanoString spatial proteomics analysis. Tissues were stained with morphology markers: AF-647–conjugated insulin (Cell Signaling Technology, 9008s; 1:400) and nuclei marker (SYTO13). Tissues were hybridized using a pre-validated mouse GeoMx Immune cell panel (NanoString, GMX-PROCONCT-MICP) comprising the following markers: PD-1, CD11c, CD8a, PanCk, MHC II, CD19, CTLA4, SMA, CD11b, CD3e, fibronectin, Ki-67, CD4, GZMB, F4/80, CD45, PD-L1; housekeeping genes: histone H3, S6, GAPDH; and IgG antibodies: Rb IgG, Rat IgG2a, and Rat IgG2b for background subtraction. All markers were conjugated to unique UV-photocleavable oligos for indexing. At least 5 or 6 islets with insulitis were chosen as regions of interest (ROIs) per mouse based on the morphology markers (insulin and nuclei). The ROIs were segmented into insulitic region and insulin<sup>+</sup> region for each islet. Oligonucleotides from the segmented ROIs were photocleaved, collected in a 96-well plate, and reads were counted using nCounter (NanoString). Scaling was performed to normalize for any differences in tissue surface area and depth. After scaling, reads were normalized to housekeeping markers and background was subtracted using IgG markers.

*Flow cytometry*. Spleen and pLNs were harvested, homogenized, and passed through a 70  $\mu$ m strainer to obtain a single-cell suspension. Cell pellets were resuspended in red blood cell (RBC) lysing solution to remove red blood cells. A total of 2.5 × 10<sup>5</sup> to 1 × 10<sup>6</sup> cells per condition were incubated with blocking solution (eBioscience, 14-0161-86) containing anti-mouse CD16/CD32 to block the Fc receptors for 20 minutes on ice. The following surface markers were used — CD4-FITC (BioLegend, 100510; 1:100), CD8–PerCP-Cy5.5 (BioLegend, 100734; 1:100), and CD19-AF700 (BioLegend, 152414; 1:100). Following incubation of surface antibodies, cells were washed with stain buffer and then permeabilized using fix/perm buffer (BD Biosciences, 554722) before intracellular staining. The following intracellular antibodies from BD Biosciences were used: FoxP3-AF647 (560401; 1:100), IFN- $\gamma$ –PE (554412; 1:50), and IL-17a–APC-CY7 (560821; 1:50). Cells were analyzed on the Attune NxT Flow Cytometer (Thermo Fisher Scientific). Data were analyzed by FlowJo software (BD Biosciences).

*RNA sequencing*. RNA extraction was performed using RLT Buffer, according to the manufacturer's instructions (Qiagen). Samples were submitted for library generation and sequencing by the University of Chicago sequencing core using a NovaSeq 6000 (Illumina). Data were analyzed using Galaxy (https://use-galaxy.org/). Reads were aligned to the *Mus musculus* genome build mm10 using HISAT2 (https://daehwank-imlab.github.io/hisat2/). Individual sample reads were quantified using HTseq-count and normalized using DESeq2 (https://bioconductor.org/packages/release/bioc/html/DESeq2.html). DEseq2 was also used to calculate FCs and *P* values and to perform optional covariate correction. GO was used for pathway analysis.

*Statistics*. All data are represented as mean  $\pm$  SEM. When comparing more than 2 conditions, 1-way ANOVA was performed. Tukey's or Dunnett's post hoc test was used to determine specific differences between individual group means. When comparing only 2 conditions, 2-tailed Student's *t* test was performed. Mantel-Cox log-rank test was specifically used for analyzing the NOD diabetes incidence experiments. Data analyses were performed using GraphPad Prism 10 software. The differences were considered statistically significant at a *P* value of less than 0.05.

*Study approval.* All experiments involving mice were performed at the University of Chicago and the procedures were conducted according to protocols approved by the University of Chicago Institutional Animal Care and Use Committee (Chicago, Illinois, USA).

*Data availability*. The islet RNA sequencing data have been uploaded to the NCBI Gene Expression Omnibus (https://www.ncbi.nlm.nih.gov/geo/) with accession number GSE272668. The BMDM sequencing data have been uploaded to the Gene Expression Omnibus with accession number GSE272687. Values for all data points in graphs are reported in the Supporting Data Values file.

#### Author contributions

JLN, DJM, MBB, SAT, and RGM conceptualized the research. TN, CM, JRE, JEW, KBK, AC, KF, SP, JBN, and SAT performed the investigation. SAT and RGM provided project supervision. TN, SCM, SAT, and RGM wrote the original draft of the manuscript. All authors contributed to discussion, edited the manuscript, and approved the final version of the manuscript.

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