ORIGINAL RESEARCH ARTICLE

Oxidized Phospholipids Promote NETosis and Arterial Thrombosis in LNK(SH2B3) Deficiency

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BACKGROUND: LNK/SH2B3 inhibits Janus kinase/signal transducer and activator of transcription (JAK/STAT) signaling by hematopoietic cytokine receptors. Genome-wide association studies have shown association of a common single nucleotide polymorphism in LNK (R262W, T allele) with neutrophilia, thrombocytosis, and coronary artery disease. We have shown that LNK(TT) reduces LNK function and that LNK-deficient mice display prominent platelet–neutrophil aggregates, accelerated atherosclerosis, and thrombosis. Platelet–neutrophil interactions can promote neutrophil extracellular trap (NET) formation. The goals of this study were to assess the role of NETs in atherosclerosis and thrombosis in mice with hematopoietic Lnk deficiency.

METHODS: We bred mice with combined deficiency of Lnk and the NETosis-essential enzyme PAD4 (peptidyl arginine deiminase 4) and transplanted their bone marrow into Ldlr−/− mice. We evaluated the role of LNK in atherothrombosis in humans and mice bearing a gain of function variant in JAK2 (JAK2V617F).

RESULTS: Lnk-deficient mice displayed accelerated carotid artery thrombosis with prominent NETosis that was completely reversed by PAD4 deficiency. Thrombin-activated Lnk−/− platelets promoted increased NETosis when incubated with Lnk−/− neutrophils compared with wild-type platelets or wild-type neutrophils. This involved increased surface exposure and release of oxidized phospholipids (OxPL) from Lnk−/− platelets, as well as increased priming and response of Lnk−/− neutrophils to OxPL. To counteract the effects of OxPL, we introduced a transgene expressing the single-chain variable fragment of E06 (E06-scFv). E06-scFv reversed accelerated NETosis, atherosclerosis, and thrombosis in Lnk−/− mice. We also showed increased NETosis when human induced pluripotent stem cell–derived LNK(TT) neutrophils were incubated with LNK(TT) platelet/megakaryocytes, but not in isogenic LNK(CC) controls, confirming human relevance. Using data from the UK Biobank, we found that individuals with the JAK2 VF mutation only showed increased risk of coronary artery disease when also carrying the LNK R262W allele. Mice with hematopoietic Lnk−/− and Jak2VF clonal hematopoiesis showed accelerated arterial thrombosis but not atherosclerosis compared with Jak2VFLnk+/+ controls.

CONCLUSIONS: Hematopoietic Lnk deficiency promotes NETosis and arterial thrombosis in an OxPL-dependent fashion. LNK(R262W) reduces LNK function in human platelets and neutrophils, promoting NETosis, and increases coronary artery disease risk in humans carrying JAK2VF mutations. Therapies targeting OxPL may be beneficial for coronary artery disease in genetically defined human populations.

Key Words: clonal hematopoiesis ■ coronary artery disease ■ extracellular traps ■ phospholipids ■ thrombosis
Clinical Perspective

What Is New?
• A common genetic variant in LNK (R262W) reduces its function, increases JAK/STAT (Janus kinase/signal transducer and activator of transcription) signaling, and associates with coronary artery disease.
• Human pluriplent stem cell-derived platelets and neutrophils carrying LNK(R262W) produce increased neutrophil extracellular traps (NETs) and hyperlipidemic LNK-deficient mice display increased NETs and arterial thrombosis that are reversed by oxidized phospholipid (OxPL) antibodies.
• LNK(R262W) increases coronary artery disease risk in humans carrying a gain of function in JAK/STAT signaling (JAK2(V515M)), paralleling increased NETosis and arterial thrombosis in LNK-deficient mice and suggesting that targeting NETosis and OxPLs in patients carrying the common LNK loss of function variant could reduce atherothrombotic risk.

What Are the Clinical Implications?
• Hypercholesterolemic mice with hematopoietic Lnk deficiency display accelerated arterial thrombosis with increased NETs in thrombi; these changes are reversed by PAD4 (peptidyl arginine deiminase) deficiency or OxPL antibodies.
• LNK-deficient platelets from hyperlipidemic mice expose and release increased OxPL when activated, promoting NETosis when incubated with Lnk-deficient neutrophils.
• An anti-OxPL antibody reduces plasma OxPL levels, NETosis, and arterial thrombosis specifically in Lnk-deficient mice.
• Targeting atherothrombotic risk using OxPL antibodies may be particularly effective in genetically defined populations with reduced Lnk function or increased JAK/STAT signaling.

Nonstandard Abbreviations and Acronyms

- CAD: coronary artery disease
- CHIP: clonal hematopoiesis of indeterminate potential
- G-CSF: granulocyte colony-stimulating factor
- H3Cit: citrullinated histone
- HSC: hematopoietic stem cell
- IL-1β: interleukin-1β
- iPSC: induced pluripotent stem cell
- JAK/STAT: Janus kinase/signal transducer and activator of transcription
- MMK: megakaryocyte
- NETs: neutrophil extracellular traps
- OxPL: oxidized phospholipid
- PAF: platelet-activating factor
- PAFR: platelet-activating factor receptor
- WT: wild-type
- WTD: Western-type diet

A common genetic variant in LNK rs3184504 (T allele, R262W) is associated with increased platelet and neutrophil counts, coronary artery disease (CAD), thrombotic stroke, and autoimmune diseases. LNK (also called SH2B3), a member of the SH2B family of adaptor proteins, is primarily expressed in hematopoietic and endothelial cells. In hematopoietic cells, LNK functions as a negative regulator of cytokine signaling and cell proliferation. Targeted deletion of LNK in mice causes expansion of hematopoietic stem cells (HSCs), increased myelopoiesis, thrombosis, and leukocytosis, attributable to lack of negative feedback regulation of thrombopoietin receptor signaling through the Janus kinase/signal transducer and activator of transcription (JAK/STAT) pathway. Similarly, in human cord blood HSCs, LNK(R262W) causes reduced LNK function and increased thrombopoietin signaling. Both the common T risk single nucleotide polymorphism and much rarer complete loss of function variants of LNK are associated with myeloproliferation and similar phenotypes occur in LNK-deficient mouse models. Thus, LNK-deficient mice are a relevant model to assess mechanisms of atherosclerosis and thrombosis associated with LNK deficiency.

We showed that hematopoietic LNK deficiency synergizes with hyperlipidemia to promote platelet production and activation, neutrophilia, platelet–neutrophil aggregates, atherosclerosis, and arterial thrombosis. Platelet activation and platelet–neutrophil interactions have been shown to promote neutrophil extracellular trap (NET) formation. NETs are formed when neutrophils release their contents, leading to the formation of web-like structures made of DNA, myeloperoxidase, citrullinated histones (H3Cit), and proteases that entrap and kill bacteria. NETs may help suppress infections, but the formation of NETs (NETosis) in blood vessels can promote atherosclerosis and thrombosis. NETs are found on the surface of and within atherosclerotic lesions and within arterial thrombi in humans and mice. NETs may serve as biomarkers predicting the risk of thrombosis. Moreover, chemical inhibition or genetic deletion of PAD4 (peptidyl arginine deiminase 4), an enzyme critical for NETosis, decreased lesion size and delayed arterial thrombosis in ApoE−/− mice. These studies suggest that NETs may promote formation of unstable atherosclerotic plaques and atherothrombosis.

This study was undertaken to investigate the hypothesis that Lnk deficiency might promote NETosis, leading to formation of unstable atherosclerotic plaques and arterial thrombosis. We also determined whether
Results in LNK(SH2B3)-Deficient Mice

As reported,26 NETs shown by coincident staining for H3Cit and MPO or Ly6G were seen mainly on the surface of thrombi that were outlined using Carstairs stain. NETs were increased 7-fold in thrombi of Lnkr−/− mice with almost complete reversal by PAD4 deficiency (Figure 1C and Figure S2). Similar results were obtained when histone H3 staining was used to assess NETs (Figure 1C and Figure S2). DNase 1, which can effectively degrade DNA in NETs, has been used to dissolve NETs in vivo.18,27 Treatment with DNase 1 ameliorated thrombosis in WTD fed Lnkr−/− mice (Figure S2). Together these data indicate that NETosis promotes arterial thrombosis in a PAD4-dependent fashion in hypercholesterolemic mice with hematopoietic LNK deficiency.

To assess the role of NETs in atherogenesis, proximal aortic lesions were stained for neutrophils and NETs (Figure S3). This showed prominent NET formation in lesions of mice with hematopoietic LNK deficiency compared with controls, with reversal by PAD4 deficiency (Figure S3). We also found increased staining for interleukin–1β (IL-1β), an inflammatory marker, especially in regions containing NETs in LNK-deficient mice, which was reversed by PAD4 deficiency (Figure S4). However, although there was a trend for PAD4 deficiency to reduce lesion and necrotic core area in LNK-deficient mice, these changes were not significant (Figure S3).

Platelet-Released Platelet-Activating Factor–Like Oxidized Phospholipid Induces NETosis in Lnkr−/− Neutrophils

Previous studies have shown a major role of activated platelets in promoting NETosis.8,28 To explore the underlying mechanisms of accelerated NETosis, we incubated WT or Lnkr−/− neutrophils with thrombin-activated WT or Lnkr−/− platelets and stained for NETs using H3Cit. We found that NETs were only significantly increased when Lnkr−/− platelets were incubated with Lnkr−/− neutrophils, indicating a role for both Lnkr−/− platelets and Lnkr−/− neutrophils in increased NETosis (Figure 2A). The increase in NETosis was significantly greater in samples from hyperlipidemic mice, consistent with augmented Lnkr−/− platelet activation and thrombosis in the setting of hyperlipidemia.7 To investigate whether soluble factors released from activated platelets promote NETosis, we prepared supernatants from activated platelets and showed similar activity of intact Lnkr−/− platelets or their supernatant, whereas supernatants of WT platelets promoted less NETosis (Figure 2B). To assess a potential role of the lipids released by platelets, we prepared a sonicated lipid extract and incubated it with Lnkr−/− neutrophils. The lipids extracted from Lnkr−/− platelets induced a significantly higher level of NETosis than those from Lnkr−/− platelets (Figure 2B). Oxidized phospholipids (OxPLs) have been implicated in platelet activation in hyperlipidemic mice.29

Methods

On reasonable request to the corresponding author, the data, analytic methods, and study materials will be made available to other researchers. All supporting data are available within the article and the Expanded Methods in the Supplemental Material. Institutional review board approval was obtained for this study where applicable.

Results

Effect of NETs on Thrombosis and Atherosclerosis in LNK-Deficient Mice

PAD4 has an essential role in chromatin decondensation and NETosis. Deficiency of PAD4 impairs venous and arterial thrombosis in mice.11,23,24 To study the role of NETosis in thrombosis, Ldlr−/− mice were transplanted with WT, Lnkr−/−, Pad4−/−, and Lnkr−/−Pad4−/− bone marrow and fed Western-type diet (WTD) for 10 weeks. As reported,7 hypercholesterolemic Ldlr−/− mice receiving Lnkr−/− bone marrow displayed monocytosis, neutrophilia, and thrombocytosis, in a PAD4-independent fashion (Figure S1). Compared with controls, carotid artery thrombosis was substantially accelerated in the Lnkr−/− background (Figure 1A). Because PAD4 is expressed by other leukocytes than neutrophils,25 we assessed the specific role of neutrophils by depleting them with Ly6G antibody (Figure 1B). Similar to PAD4 deficiency, depletion of neutrophils had no effect in WT mice but significantly prolonged occlusion time in Lnkr−/− mice (Figure 1B), suggesting that neutrophil PAD4 promoted NETosis in Lnkr deficiency.

Animals

Wild-type (WT; C57BL/6J), Ldlr−/− (B6.129S7-Ldlrtm1Her), and Pad4−/− (B6.Cg-Padi4tm1.1Kmow/J) mice were purchased from Jackson Laboratory. Ldlr−/− E06-scFv mice (on C57BL/6J background) were shared by the Witztum Laboratory at the University of California, San Diego.21,22

Statistics

Statistical analysis included mean±SEM, Student t test, and 2-way analysis of variance. Results were considered significant at P<0.05. Statistical analyses were conducted using Prism (GraphPad).

Discussion

NETosis in LNK(SH2B3)-Deficient Mice

LNK(R262W) might modify the increased CAD risk associated with JAK2V617F (JAK2v617f), a gain of function variant associated with rare myeloproliferative neoplasms19 and more commonly with clonal hematopoiesis of indeterminate potential (CHIP),20 as LNK acts as a negative regulator of both JAK2 and JAK2v617f-mediated signaling and myeloproliferation.5,6

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OxPLs are released by activated platelets and a recent report showed that OxPLs promote NETosis in vitro. This suggests that soluble OxPLs released from Lnk−/− platelets might be inducing NETosis in Lnk−/− neutrophils. We found that oxPAPCs markedly induced NETosis especially when added to Lnk−/− versus WT neutrophils (Figure 2C). oxPAPCs demonstrate platelet-activating factor (PAF)–like agonistic activity through the PAF receptor (PAFR), which is inhibited by the specific PAFR antagonist WEB2086. OxPAPCs consistently induced...
NETosis in Lnk\(^{-/-}\) but not WT neutrophils and this Ox-PAPC induced NETosis was inhibited by WEB2086 (Figure 2C). PAF stimulation resulted in significantly increased NETosis in Lnk\(^{-/-}\) but not WT neutrophils compared with controls (Figure S5), similar to the effects of oxPAPC. When Lnk\(^{-/-}\) platelets were incubated with Lnk\(^{-/-}\) neutrophils, the increased NETosis was reversed by WEB2086 (Figure S5). Thus, PAF-like lipids, most likely OxPLs, released by Lnk\(^{-/-}\) platelets promote NETosis by activation of the PAFR in Lnk\(^{-/-}\) neutrophils.

**Figure 2.** Platelet-released platelet-activating factor–like oxidized phospholipid induces neutrophil extracellular traps in Lnk\(^{-/-}\) neutrophils through platelet-activating factor receptor.  

**A.** Platelets activated with 1 nmol/L thrombin were incubated with neutrophils at a ratio=20:1 for 4 hours. Representative images of NETosis. Red, Citrullinated histone (H3Cit); blue, DAPI. Neutrophil extracellular trap (NET) numbers were normalized to total neutrophil number; 2-way analysis of variance. Scale bar, 100 µm.  

**B.** Lnk\(^{-/-}\) neutrophils were stimulated with wild-type (WT) or Lnk\(^{-/-}\) platelets, platelet-derived supernatant, or platelet supernatant–derived lipids for 4 hours. Red, H3Cit; blue, DAPI; n=7.  

**C.** Neutrophils were preincubated with WEB2086 (4 µmol/L) for 30 minutes followed by 3 hours oxidized phospholipid (OxPL; 20 µg/mL) treatment and then stained with H3Cit and DAPI (n=7). Quantification of NETs; 2-way analysis of variance. Scale bar, 100 µm. Data are expressed as mean±SEM. SN indicates supernatant; and WTD, Western-type diet.

**E06-scFv Ameliorates Thrombosis and Atherosclerosis in LNK-Deficient Mice**

A recent study using E06-scFv transgenic mice that express OxPL-neutralizing antibodies demonstrated a major role of OxPLs in promoting atherogenesis.\(^{21}\) To assess the role of OxPLs in thrombosis and atherosclerosis, we bred E06-scFv transgenic Ldlr\(^{-/-}\) mice. Bone marrow cells from WT and Lnk\(^{-/-}\) mice were transplanted into male and female Ldlr\(^{-/-}\) and Ldlr\(^{-/-}\) E06-scFv mice.

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Figure 3. E06-scFv reverses increased neutrophil extracellular traps and accelerated thrombosis in Lnk−/− mice.

FeCl₃-induced carotid artery occlusion in Western-type diet–fed (10 weeks) female (A) or male (B) recipients of designated genotype as indicated; 2-way analysis of variance. C. Representative images of Carstairs, fibrinogen, or neutrophil extracellular trap (NET) staining in carotid artery thrombi of the female recipients as in A. NETs were validated using MPO (green) with citrullinated histone (H3Cit; red). (Continued)
followed by 10 weeks of WTD feeding. After 6 weeks of WTD, there were high levels of E06-scFv in Ldr<sup>−/−</sup> E06-scFv recipient mice, especially in males (Figure S6). Body weight, spleen weight, and blood cell counts were not affected by the E06-scFv transgene in either sex (Figures S6 and S7).

Accelerated thrombosis in LNK-deficient female mice was almost completely reversed by the E06-scFv transgene, whereas E06-scFv had no effect in the WT controls (Figure 3A), paralleling the selective reversal of accelerated thrombosis in LNK deficiency by PAD4 deficiency or neutrophil depletion. Similar results were obtained in male mice (Figure 3B). In parallel, increased NETosis in the thrombi of Lnk<sup>−/−</sup> mice was reversed by E06-scFv (Figure 3C and Figures S8 and S9). E06 does not bind to PAF or lysoPAF. Plasma OxPL levels were significantly reduced in E06-scFv mice (Figure 3F). Whereas E06-scFv did not significantly reduce lesion area in controls, previous studies showed reduced lesion area in Ldr<sup>−/−</sup> E06-scFv compared with Ldr<sup>−/−</sup> mice fed with 1% high-cholesterol diet for 4, 7, or 12 months, compared with only 10 weeks in the current study. The effect of E06-scFv was less pronounced in male than in female mice, which had more than 2-fold larger lesions (Figure 3B and 3C). Together, these findings suggest that E06-scFv is more effective at later stages of lesion development. Given that PAD4 deficiency did not affect lesion development (Figure S3), the more prominent effect of E06-scFv in ameliorating lesions of Lnk<sup>−/−</sup> versus control Ldr<sup>−/−</sup> mice is unlikely to be solely attributable to reversal of NETosis, but rather could reflect the fact that the LNK-deficient mice had more advanced lesions.

**Granulocyte Colony-Stimulating Factor Promotes NETosis in LNK-Deficient Mice**

Our in vitro incubations suggest a NETosis-promoting role of LNK deficiency in neutrophils as well as in platelets (Figure 2). Granulocyte colony-stimulating factor (G-CSF) signaling through STAT3/5 is known to be inhibited by LNK<sup>35</sup> and G-CSF treatment primes neutrophils for NET release. To assess G-CSF signaling in LNK deficiency, we treated WT and Lnk<sup>−/−</sup> mice with recombinant human G-CSF and isolated neutrophils. Compared with WT, p-STAT3 expression was significantly increased in Lnk<sup>−/−</sup> neutrophils even under basal conditions and was increased further after G-CSF administration. H3Cit levels were also elevated in Lnk<sup>−/−</sup> neutrophils after G-CSF treatment (Figure 5A). To assess G-CSF induced NETosis further, we treated WT and Lnk<sup>−/−</sup> neutrophils with G-CSF in vitro. G-CSF stimulation resulted in a dose-dependent increase in NETosis in both WT and Lnk<sup>−/−</sup> neutrophils, but the effect was more pronounced in Lnk<sup>−/−</sup> neutrophils (Figure 5B). We next assessed whether G-CSF treatment in vivo would prime neutrophils for NETosis induced by other stimulants ex vivo. This showed increased NETosis in response to PAF treatment in Lnk<sup>−/−</sup> neutrophils compared with WT neutrophils (Figure 5C–5E), suggesting that increased G-CSF-mediated priming may be responsible for the increased NETosis observed in Lnk<sup>−/−</sup> neutrophils.

**Figure 3 Continued.** NETs shown were from the same area as in fibrinogen staining on consecutive sections. Quantification of NETs in carotid artery thrombosis of the female recipient (n=4 per group); 2-way analysis of variance. Scale bar, 50 µm. Each point represents 1 mouse. D, Plasma oxidized phospholipid (OxPL) level was assessed in the female recipients as in A (n=6–7). E, E06-scFv binding to the basal or activated platelets (n=8). F, Plasma P-selectin level was assessed by enzyme-linked immunosorbent assay in the female recipients as in A (n=8). Surface P-selectin (G) and active integrin α<sub>IIb</sub>β<sub>3</sub> (JON/A; H) levels on platelets with and without AYPGKF (50 μmol/L) stimulation (n=6); 2-way analysis of variance. I, Quantification of NETs from wild-type (WT) or Selplg<sup>−/−</sup> neutrophils stimulated by activated WT or Lnk<sup>−/−</sup> platelets (n=5); 2-way analysis of variance. Data are expressed as mean (red bar) ±SEM.
Increased NETosis in Human Induced Pluripotent Stem Cell–Derived LNK(TT) Platelet/Megakaryocytes and Neutrophils

To test the human relevance of these findings, we generated isogenic LNK(TT) and isogenic LNK(CC) induced pluripotent stem cells (iPSCs) through CRISPR/Cas9-mediated gene editing (Figure S14). To select an LNK homozygous parental iPSC line, we first genotyped a panel of 17 previously generated genetically diverse normal lines. Thirty-three lines were LNK(TT), 8 were LNK(CC), and 3 were heterozygous LNK(CT). We selected a LNK(TT) iPSC line (N=2,123) and used CRISPR/Cas9-mediated homology-directed repair to edit the LNK locus to derive an isogenic LNK(CC) line. Using a differentiation
protocol that produces definitive-type hematopoietic progenitor cells followed by protocols to generate megakaryocyte/platelets or neutrophils, we generated mature megakaryocyte (MMK)/platelets and neutrophils from the isogenic iPSC pair. 39,40 By day 15 of differentiation, >70% of the cell population were CD41a+CD42b+ MMKs generated from megakaryocyte progenitor cells and large megakaryocytes and
proplatelet morphology was observed (Figure S15). LNK(TT) MMK/platelet mixtures showed increased surface P-selectin level under basal and stimulated conditions compared with LNK(CC) control cells (Figure 6A). p-STAT5 was also significantly increased in LNK(TT) neutrophils (CD45+CD14− gated) cultured in G-CSF–containing medium (Figure 6B), corroborating our finding of increased G-CSF signaling through STAT3/5 in Lnk-deficient neutrophils. To assess NETosis induced by MMK/platelets in human iPSC-derived neutrophils, we incubated LNK(TT) and LNK(CC) neutrophils with thrombin-activated MMK/platelets. There was significantly increased NETosis when LNK(TT) neutrophils were incubated with LNK(TT) MMK/platelets, but not when LNK(CC) neutrophils were incubated with LNK(CC) MMK/platelets (Figure 6C). Paralleling our findings in murine neutrophils, PAF stimulation resulted in significantly increased NETosis in LNK(TT) but not LNK(CC) neutrophils (Figure 6D).

**Figure 6.** Increased platelet activation, STAT signaling, and NETosis in human induced pluripotent stem cell–derived LNK(TT) cells. A, Surface P-selectin level on megakaryocytes (MMKs)/platelets with and without ADP (20 µmol/L) and thrombin (1 U/mL) or AYPGKF (50 µmol/L) stimulation from day 15 LNK(TT) and LNK(CC) cells (n=3). Unpaired t test. B, p-STAT5 level was analyzed in day 13 CD45+CD14− LNK(TT) and LNK(CC) cells (n=3). Unpaired t test. C, MMKs/platelets activated with 1 nmol/L thrombin were incubated with LNK(TT) and LNK(CC) neutrophils for 4 hours. Representative images of NETosis. Green, Neutrophil elastase; red, citrullinated histone (H3Cit); blue, DAPI. Neutrophil extracellular trap (NET) numbers were normalized to neutrophil elastase positive cell numbers; 2-way analysis of variance. Scale bar, 50 µm. D, LNK(TT) and LNK(CC) neutrophils were stimulated with 5 µmol/L platelet-activating factor (PAF) for 3 hours and quantified for NETs (n=6); 2-way analysis of variance. Scale bar, 50 µm. Data are expressed as mean±SEM.
Increased CAD in JAK2VF Subjects Homozygous for LNK(R262W)

We evaluated the effect of the LNK(R262W) reduced function variant on CAD in JAK2VF subjects in the UK Biobank first release (Table). There were 232 JAK2VF carriers (rs77375493) in UK Biobank. We examined the association of this variant with CAD by LNK R262W (rs3184504) status. This suggested a dosage-related effect of the LNK R262W allele to increase CAD. A formal test for trend by genotype is significant \((P=0.03)\), with the JAK2VF variant predisposing to CAD among LNK R262W T/T carriers but not among LNK R262W C/C carriers.

To model acquired JAK2VF mutation as would occur in individuals with JAK2VF clonal hematopoiesis on the germline LNK(R262W) reduced function background, we transplanted a mixture of 20% Jak2VF Lnk+/− bone marrow with 80% Lnk−/− bone marrow into Ldlr−/− mice, followed by treatment with polyIC to activate the Jak2VF transgene,7,11,42 and then fed mice WTD for 10 weeks. Control mice were transplanted with 20% Jak2VF Lnk+/− bone marrow mixed with 80% Lnk+/− bone marrow and then received similar treatment. Ldlr−/− mice receiving Jak2VF Lnk+/− bone marrow displayed slightly increased monocytes and hematocrit but neutrophil and platelet counts, body weight, and spleen weight were not different compared with the latter group (Figure 7A and 7B). Arterial thrombosis was markedly accelerated in Jak2VF Lnk+/− mice compared with Jak2VF Lnk−/− mice (Figure 7C) with parallel effects on NETs in thrombi (Figure 7D and Figure S17). Although NETs were also more prominent in lesions of Jak2VF Lnk+/− mice (Figure 7F and Figure S18), atherosclerotic lesion size and necrotic core area were not different comparing the Jak2VF Lnk+/− and Jak2VF Lnk−/− groups (Figure 7E).

DISCUSSION

We found that NETs are increased in atherosclerotic lesions and arterial thrombi of mice with hematopoietic LNK deficiency. Despite having little effect on atherogenesis, PAD4 deficiency almost completely reversed NETosis and accelerated carotid arterial thrombosis in hyperlipidemic LNK-deficient mice, suggesting a major role of NETosis in the accelerated formation of thrombi on early carotid atherosclerotic lesions. On a mechanistic level, our studies suggest that OxPLs, expressed on the surface or released from LNK-deficient platelets, bind to the neutrophil PAF receptor, acting in parallel to increased P-selectin signaling to enhance NETosis (Figure S19). Neutralization of OxPLs through the E06 transgene reversed platelet activation and accelerated thrombosis.

On a translational level, we confirmed that LNK(R262W) causes reduced inhibition of cytokine signaling in isogenic human iPSC-derived megakaryocyte/platelets and neutrophils, leading to increased NETosis. Together with our earlier studies,7 these observations further support that LNK(R262W) is a reduced function variant that leads to increased JAK/STAT signaling. We discovered that the CAD risk associated with a gain of function in JAK/STAT signaling (JAK2VF) was significantly increased by the reduced function variant LNK(R262W) variant, correlating with an increase in NETosis and thrombosis in a mouse model of partially reduced LNK function and Jak2VF clonal hematopoiesis. Together these studies point to a potential therapeutic role of antioxidant phospholipid antibodies in suppressing NETosis and thrombosis and suggest they might be particularly effective in genetically defined subpopulations.

Our studies suggest a major role of OxPLs in plasma and platelets in the increased NETosis and accelerated thrombosis observed in Lnk−/− mice. OxPLs are formed during tissue injury and have been implicated in the pathogenesis of several diseases. The recent availability of E06-scFv transgenic mice has allowed direct assessment of the role of OxPL in disease pathogenesis. In addition to reducing atherosclerosis, the E06-scFv transgene also ameliorated nonalcoholic steatohepatitis and reduced myocardial ischemia/reperfusion injury.21,22,43 Our findings provide the first evidence that inactivating OxPL inhibits arterial thrombosis. We found that in parallel with reversing accelerated thrombosis, E06-scFv reduced increased plasma OxPL levels and bound in increased amounts to the surface of activated Lnk−/− platelets. Conceivably, E06-scFv bound to plasma OxPL and prevented assessment of free OxPL in the assay, which uses exogenous E06 in a competitive immunoassay. In either case, this represents the first demonstration that E06-scFv can reduce plasma OxPL levels, or effective free OxPL levels, in parallel with an improved biological outcome. Our data suggest that Lnk−/− platelets expose or release OxPLs that activate neutrophils through the PAFR-promoting NET release. PAF is a phospholipid with phosphatidylycholine as headgroup and sn1 linked ether and sn2 acetyl group. 

| Subgroup | CAD in JAK2VF carriers | CAD in noncarriers | HR for CAD | \( P \) value |
|----------|------------------------|-------------------|-----------|-------------|
| All of UK Biobank | 17/232 | 5427/132032 | 1.50 | 0.12 |
| SH2B3 R262W C/C | 2/53 | 1469/39879 | 0.77 | 0.73 |
| SH2B3 R262W C/T | 8/117 | 2688/66431 | 1.33 | 0.46 |
| SH2B3 R262W T/T | 7/62 | 1269/30931 | 2.48 | 0.03 |

A formal test for trend by genotype is significant \((P=0.03)\), with the JAK2VF variant predisposing to coronary artery disease (CAD) among SH2B3 R262W T/T carriers (hazard ratio [HR], 2.48; \( P=0.03)\), but not among SH2B3 R262W C/C carriers (odds ratio, 0.77; \( P=0.73)\).
Figure 7. Increased neutrophil extracellular traps and accelerated thrombosis in Jak2<sup>wt</sup>Lnk<sup>+/−</sup> mice.

A. Surface P-selectin level on platelets in whole blood with and without AYPGKF (50 µmol/L) stimulation from mice after 2, 6, and 10 weeks Western-type diet (WTD; n=6–7); 2-way analysis of variance. *P<0.05, **P<0.01, ***P<0.001. 

B. Surface P-selectin level on platelets in whole blood with and without ADP (20 µmol/L) and AYPGKF (50 µmol/L) after 10 weeks WTD feeding (n=7–8).

C. FeCl<sub>3</sub> induced carotid occlusion in bone marrow Ldlr<sup>−/−</sup> recipient mice (n=15–17).

D. Representative images of Carstairs, fibrinogen, or neutrophil extracellular trap (NET) staining in thrombi of the recipients. NETs were validated using MPO (green) with citrullinated histone (H3Cit; red). Quantification of NETs in thrombi of the recipients (n=4 per group). Unpaired t test. Scale bar, 50 µm. Each point represents 1 mouse.

E. Representative images showing staining for MPO (green) and citrullinated histone (H3Cit; red) with DAPI (blue) in aortic root lesions of Ldlr<sup>−/−</sup> recipients fed with WTD for 10 weeks and quantification of NETs (MPO<sup>+</sup>H3Cit<sup>+</sup>) in lesions. Unpaired t test. Scale bar, 50 µm.
does not bind to either PAF or lysoPAF. Thus, the in vivo experiments suggest that inhibiting PAF is not likely the reason for effectiveness of E06, and that effectiveness of the PAFR antagonist in vitro more likely reflects antagonism of OxPLs released by or exposed on the surface of Lnk–/– platelets.

Consistent with evidence that platelet activation stimulates NETosis, we found NET formation was increased in neutrophils incubated with Lnk–/– or Lnk(TT) platelets that showed higher levels of activation compared with WT or Lnk(CC) megakaryocyte/platelets. The reversal by E06 of elevated platelet activation markers such as P-selectin suggests that in addition to providing a signal to activate the PAFR on neutrophils, OxPL may be acting indirectly to increase platelet activation and NETosis in Lnk-deficient mice. Previous studies have shown that increased OxPLs in hyperlipidemic mouse plasma activate platelets through CD36 signaling and that TLR4 activation in platelets increases their ability to promote NETosis. Decreased plasma OxPL and inhibition of platelet activation by OxPL–platelet CD36 interactions or OxPL–TLR interactions could contribute to the in vivo effects of E06. Because platelet P-selectin/neutrophil P-selectin receptor interaction is a key pathway in the stimulation of NETosis, this could be an additional mechanism leading to increased NETosis. Indeed, Selplg–/– neutrophils were resistant to increased NETosis induced by Lnk–/– platelets. Because both PAF receptor inhibition and Selpg knockout inhibited the stimulation of NETosis by Lnk–/– platelets, these receptors may be acting in parallel pathways to increase NETosis in Lnk deficiency (Figure S19). There is precedent for cooperative signaling by P-selectin and PAF receptors in neutrophils. We also found that Lnk–/– and Lnk(TT) neutrophils were more susceptible to NETosis, an effect that we attribute to increased priming by G-CSF/JAK/STAT signaling, rendering neutrophils more susceptible to NETosis in response to OxPL or PAF (Figure S19).

Our findings that PAD4 deficiency reduced thrombosis while having little effect on atherosogenesis are consistent with a recent study showing that although PAD4 deficiency did not reduce plaque size in WT bone marrow transplanted Ldir–/– mice, it decreased the extent of endothelial injury and thrombosis in a model of superficial plaque erosion. NETs may contribute to features of plaque instability under conditions of increased neutrophil production and entry into plaques. Thus, NETs seem to contribute to plaque development and features of instability under specific experimental conditions but not in the standard Ldir–/– model. Whereas platelets do not readily enter plaques, NETosis stimulated by platelet-associated OxPL may trap platelets and stimulate fibrin deposition during thrombus formation in Lnk deficiency.

Consistent with its role in inhibiting JAK/STAT signaling, we showed a genetic interaction between the reduced function LNK variant (R262W) and the gain of function JAK2VF to increase CAD risk. LNK promotes the association of the E-3 ligase CBL with JAK2, promoting its ubiquination and degradation. LNK also inhibits the increased signaling of JAK2 and the common LNK(R262W) variant increases the risk of developing myeloproliferative neoplasm in individuals with the JAK2VF mutation. We show that LNK(R262W) increases CAD risk in individuals with JAK2VF, identified in a general European population and therefore much more likely to have JAK2VF CHIP than myeloproliferative neoplasms. Modeling CHIP in mice with Lnk–/– in all hematopoietic cells and JAK2VF induced in a subset of hematopoietic cells, we showed accelerated thrombosis but not atherosclerosis. This suggests that the increased CAD in JAK2VF subjects carrying LNK(R262W) might reflect increased thrombotic risk. Among different clonal hematopoiesis variants, JAK2VF most dramatically increases thrombotic risk, including both arterial and venous thrombosis and venous thrombo-embolism. Similar to our findings, these studies showed increased NET formation in venous thrombi of Jak2VF mice. However, the LNK(R262W) variant has not been associated with venous thrombo-embolism in genome-wide association studies and its role in modifying venous thrombotic risk in patients with JAK2VF remains uncertain. To reduce CAD risk, patients carrying JAK2VF might benefit from more effective antithrombotic therapy, such as multiple dosing with aspirin to compensate for increased platelet production or perhaps anti-inflammatory or anti-OxPL antibodies.

In summary, using genetic approaches to interfere with NETosis, we show a major contribution of NETs to arterial thrombosis in hyperlipidemic LNK-deficient mice (Figure S19). Novel strategies to decrease NETosis such as PAD4 inhibitors or treatment with OxPL neutralizing antibodies could reduce the risk of atherothrombosis in individuals with the common LNK(R262W) risk variant, including those with JAK2VF CHIP. Information on genetic risk conferred by CHIP mutations and LNK R262W could help to define populations that would benefit most from these or similar therapeutic strategies.

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