Autophagy Is Induced Upon Platelet Activation and Is Essential for Hemostasis and Thrombosis

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PLATELETS AND THROMBOPOIESIS

Autophagy is induced upon platelet activation and is essential for hemostasis and thrombosis

Madhu M. Ouseph, Yunjie Huang, Meenakshi Banerjee, Smita Joshi, Laura MacDonald, Yu Zhong, Huijuan Liu, Xianting Li, Binggang Xiang, Guoying Zhang, Masaaki Komatsu, Zhenyu Yue, Zhenyu Li, Brian Storrie, Sidney W. Whiteheart, and Qing Jun Wang

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Key Points

- Autophagy, an essential degradation pathway, is constitutively active in resting platelets and is induced upon platelet activation.
- Platelet autophagy is indispensable for hemostasis and thrombus formation.

Autophagy is important for maintaining cellular homeostasis, and thus its deficiency is implicated in a broad spectrum of human diseases. Its role in platelet function has only recently been examined. Our biochemical and imaging studies demonstrate that the core autophagy machinery exists in platelets, and that autophagy is constitutively active in resting platelets. Moreover, autophagy is induced upon platelet activation, as indicated by agonist-induced loss of the autophagy marker LC3II. Additional experiments, using inhibitors of platelet activation, proteases, and lysosomal acidification, as well as platelets from knockout mouse strains, show that agonist-induced LC3II loss is a consequence of platelet signaling cascades and requires proteases, acidic compartments, and membrane fusion. To assess the physiological role of platelet autophagy, we generated a mouse strain with a megakaryocyte- and platelet-specific deletion of Atg7, an enzyme required for LC3II production. Ex vivo analysis of platelets from these mice shows modest defects in aggregation and granule cargo packaging. Although these mice have normal platelet numbers and size distributions, they exhibit a robust bleeding diathesis in the tail-bleeding assay and a prolonged occlusion time in the FeCl3-induced carotid injury model. Our results demonstrate that autophagy occurs in platelets and is important for hemostasis and thrombosis. (Blood. 2015;126(10):1224-1233)

Introduction

Autophagy, one of the major degradation pathways in eukaryotes, is important for cellular homeostasis and is implicated in a broad spectrum of human diseases including cancer, neurodegeneration, and cardiovascular diseases.1 There are 3 main types: chaperone-mediated autophagy,2 microautophagy,3 and the primary form, macroautophagy (referred to hereafter as autophagy).3 Autophagy involves de novo synthesis of double-membrane organelles called autophagosomes that contain cytosolic constituents including damaged organelles (eg, mitochondria) and protein aggregates. Autophagosomes fuse with multivesicular bodies, late endosomes, and lysosomes to form autolysosomes,6 in which waste elimination, energy production, and recycling of cellular components take place.

Autophagosome biogenesis and maturation use several protein complexes.7-10 In mammals, cellular signals (eg, starvation) activate the Ulk1 complex (Ulk1, FIP200, Atg13, and Atg101),11-14 which together with syntaxin 17,15 localizes Atg14/Atg14L16-19 to the endoplasmic reticulum.15,20 This recruits the Beclin 1-Vps34 core complex (Beclin 1, Vps34, and Vps15) to autophagosome initiation sites and promotes phosphatidylinositol 3-phosphate production for recruitment of additional autophagy pathway components.20,21 Two ubiquitin-like conjugation systems are also important for autophagosome biogenesis. One uses Atg7 and Atg10 (E1 and E2 enzymes, respectively) to form an Atg12–Atg5 conjugation.22-24 A second system uses Atg7, Atg3, and the Atg12–Atg5–Atg16L1 complex (E1, E2, and E3 enzymes, respectively) to catalyze the addition of phosphatidylethanolamine to the microtubule-associated protein 1A/B light chain 3 (LC3), producing LC3II.25,26 LC3II is attached to both inner and outer autophagosome membranes,27 and thus is a marker for isolation membranes (ie, phagophores), autophagosomes, and autolysosomes. During maturation, autophagosomes fuse with multivesicular bodies, late endosomes, and lysosomes, forming acidic compartments in which LC3II on the inner autophagosome membrane is degraded, along with the luminal contents. This fusion process is mediated by syntaxin 17, SNAP29, VAMP3, VAMP8, and Atg14/Atg14L.28-30

Although autophagy is important in nucleated cells, its role in anucleate cells is less recognized. In the eye, deletion of Atg5 or Vps34, although causing cataracts, does not impair organelle clearance in the lens organelle-free zone.31 The role of autophagy in differentiated anucleate red blood cells is unclear, despite its importance for mitochondrial...
clearance during erythroblast maturation.22–38 Autophagy in platelets is understudied. Only recently, a role for autophagy in megakaryopoiesis and thrombopoiesis was indicated in a study on hematopoietic lineage-specific Atg7 deletion mice carrying Vav promoter-driven Cre.39 Feng et al reported starvation/rapamycin-induced autophagy in platelets.40 They also found that platelets isolated from whole-body heterozygous Beclin1 knockout mice are defective in ex vivo collagen-induced aggregation, as well as flow-based platelet adhesion and thrombus formation.40 However, they did not detect autophagy induction in response to platelet agonists, nor were they able to distinguish the unique contribution of platelet autophagy deficiency to hemostasis impairment, given Beclin1-deficiency in other cell types (eg, endothelial cells). Our studies extend these initial observations and show that autophagy not only occurred constitutively in resting platelets but also was induced during platelet activation. We further show that megakaryocyte- and platelet-specific deletion of Atg7 (using platelet factor 4 [PF4] promoter-driven Cre) caused modest defects in platelet aggregation and granule cargo packaging, but had a profound effect on hemostasis and thrombosis, despite normal platelet counts and mean platelet volumes. Our results demonstrate a critical role for autophagy in platelet activation during hemostasis and thrombosis.

**Method**

**Mouse strains**

Animal procedures were approved by the Institutional Animal Care and Use Committees at the relevant institutions. GFP-LC3+/+ mice, in which GFP-LC3 expression is controlled by an actin promoter, were obtained from Noboru Mizushima (University of Tokyo, Japan).14–16 Beclin1-EGFP/+43 and EGFP-Atg5/+ mice were generated using bacterial artifical chromosome (BAC) transgenics.44 To make EGFP-Atg5/+ mice, BAC clone (#RP24-117F24) containing the entire mouse Atg5 gene was purchased from Children’s Hospital Oakland Research Institute (Oakland, CA). The sequence for EGFP was inserted after the Atg5 start codon, and the modified BAC was injected into B6C3F1 hybrid eggs at the Mount Sinai School of Medicine core facility (New York, NY). GFP-LC3+/+, Beclin1-EGFP/+, and EGFP-Atg5/+ mice were genotyped using 5'-CCT ACG GCC TGC AGT GCT TCA GC-3' (forward) and 5'-CGG CGA CGT GCC GCA CGT GCC TGC CTC-3' (reverse).

VAMP8+/+ mice were described previously.45 Platelet-specific VAMP2/3+/− mice were generated by PF4-Cre-mediated expression of tandem neurotoxin endopeptidase, which cleaves both VAMP2 and VAMP3.55 VAMP2/3−/− mice were generated from VAMP2/3+/− and VAMP8+/− mice and will be described elsewhere. Arg2/2, PF4-Cre+/+ mice were generated by crossing Arg2/2 mice with PF4-Cre mice.56 Floxed Arg7 alleles were genotyped using 5'-TGG CTA CTT CTC GCA AAG TGA TG-3' (forward) and 5'-CGG AGC GAC AGC CAC GTC CTC-3' (reverse). PF4-Cre was genotyped using 5'-CCC ATA CAG CAC ACC TTT TG-3' (forward) and 5'-TGC ACA GTC AGC AGG TT-3' (reverse).

**Ex vivo assays**

Human platelets were prepared as described46 from platelet-rich plasma (obtained from the Kentucky Blood Center). Mouse platelets were prepared as described.49 Platelet-rich plasma was separated at 675 g for 10 minutes. Platelet pellets were washed from the retroorbital plexus into heparinized capillary tubes containing EDTA (10 mM). Platelet counts and mean platelet volumes were analyzed with a Hematometer (Log, Havertown, PA).

For western blot analysis, washed platelets (1.2 × 10⁹/mL) were preincubated with the indicated inhibitors and stimulated with the indicated agonists. Proteins were resolved by sodium dodecyl sulfate-polyacrylamide gel electrophoresis on 4% to 12% (or 12% for LC3) Bis-Tris gels (Invitrogen, Grand Island, NY) before western blotting, using appropriate antibodies (see the supplemental Methods, available on the Blood Web site). Films were exposed for different periods to optimize signals. Images of films were scanned in positive mode and analyzed using ImageJ (National Institutes of Health).

Platelet aggregation and adenosine triphosphate release were measured with washed platelets (2.5 × 10⁹/mL), as described.47–49 In a Lumi-Dual Aggregometer (Model 460VS) with Aggro/Link interface (Model 810) and software (Chrono-Log, Havertown, PA).

Dense granule, α-granule, and lysosome secretion were assayed as described.45–47 Specifically, PF4 was assessed by anti-PF4/CXCL4 ELISA, [H]-serotonin levels determined by scintillation counting, and β-hexosaminidase activity evaluated using a colorimetric assay.50 Total PF4 levels, total uptake of [H]-serotonin, and total β-hexosaminidase activity were recorded, and the percentage secretion was calculated.51

**In vivo tail-bleeding assay**

This assay was performed as described.57 To measure bleeding, tails of 5- to 6-week-old sedated (ketamine, 75 mg/kg, intraperitoneally) mice were transected 3 mm from the tip and immersed in saline at 37°C. The time required for bleeding cessation was recorded, and the mice were observed for an additional minute. Experiments were terminated at 10 minutes. The transected tail tips were used for genotyping.

**In vivo FeCl3-induced carotid injury model**

This assay was performed as described.58 Mice at 21 to 32 weeks of age were anesthetized with avertin (tribromoethanol; 200 mg/kg; intraperitoneally). The left carotid artery was exposed and visualized through a dissecting microscope. A Doppler, microvascular probe was placed on the exposed artery to measure vascular blood flow, and the artery was injured by placing a small piece of filter paper saturated with 6% FeCl3 directly on the artery for 3 minutes. Times to form stable, occlusive thrombi were recorded. Experiments were terminated at 3 minutes postocclusion or 30 minutes postinjury, whichever occurred first.

**Platelet count and volume measurements**

Mice were anesthetized with isoflurane, and blood (50–200 μL) was collected from the retroorbital plexus into heparinized capillary tubes containing EDTA (10 mM). Platelet counts and mean platelet volumes were analyzed with a HEMAVET HV950FS analyzer (Drew Scientific, Waterbury, CT).

**Results**

**Autophagy machinery is present in murine and human platelets**

Autophagy gene transcripts57,58 and proteins59–62 have been detected in platelets. To confirm, we probed murine and human platelet extracts with antibodies to autophagy-related proteins. We detected components of the Ulk1 complex (eg, Ulk1 and FIP200), the Atg12-Atg5 and LC3II conjugation systems (eg, Atg7, Atg3, Atg12, Atg5, and LC3II), and the Beclin 1-Vps34 complex (eg, Beclin 1, Vps34, Vps15, Atg14L, Nrfl2, and UVRRAG) (Figure 1A; supplemental Figure 1, summarized in supplemental Table 1). We next examined platelets from transgenic mice expressing GFP-fused autophagy markers (LC3,43 Atg5, and Beclin 1)43 to visualize isolation membranes, autophagosomes, and autolysosomes in resting platelets. GFP-LC3, expressed at higher levels than endogenous LC3II
Basal autophagy occurs in resting platelets

To determine whether basal autophagy occurred in resting platelets, we monitored LC3II levels in unstimulated platelets in the presence or absence of lysosomal inhibitors (e.g., NH₄Cl and chloroquine). Treatment with NH₄Cl and, less effectively, with chloroquine, increased LC3II levels over time in unstimulated human platelets (Figure 2A). Consistently, unstimulated GFP-LC3/⁺ platelets containing GFP-LC3-positive puncta increased from 21% in the absence of NH₄Cl to 64% ± 10% in the presence of NH₄Cl (Figure 2B; P < 10⁻⁷). Together, these findings indicate the presence of a basal autophagic flux (ie, turnover of autophagosomes) in resting platelets, which can be inhibited by lysosomal inhibitors.

Because anucleate platelets have few means for transcription, we asked whether translation contributed to basal autophagy, and thus accounted for the changes in LC3II seen in lysosomal inhibitor-treated resting platelets. Neither cycloheximide nor puromycin affected LC3II levels, regardless of whether lysosomal proteolysis was inhibited or not (Figure 2C). These results imply that in resting platelets, LC3II and autophagosomes are continuously produced, independent of LC3 translation.

Platelet activation leads to LC3II reduction and altered autophagic flux

Given that autophagic flux appeared to occur in resting platelets, we sought to determine whether it changed upon platelet activation. We found that LC3II levels decreased in response to thrombin in both murine and human platelets; this observation was confirmed using 4 different anti-LC3 antibodies (Figure 3A). We tested other platelet agonists and found that LC3II levels in human platelets were reduced by ~2.5-fold in response to strong activators (thrombin, convulxin [CVX], and PAR1 peptide), whereas weaker activators (ADP, collagen, and U46619) led to modest LC3II reduction (~1.5-fold) (Figure 3B). Similar agonist-induced LC3II reductions were observed in murine platelets (supplemental Figure 3). These results indicate that platelet activation results in a loss of LC3II, indicating an alteration in autophagic flux.

To delineate which platelet activation step or steps are required for agonist-induced LC3II reduction and to determine whether autophagy is downstream of the known platelet signaling cascades, we tested the effects of platelet activation inhibitors on agonist-induced LC3II reduction. The phospholipase C inhibitor, U-73122, but not its inactive analog, U-73343, blocked thrombin-induced LC3II reduction (Figure 3C). The permeant calcium chelator BAPTA-AM and the protein kinase C inhibitor Ro-31-8220 also blocked thrombin-induced LC3II reduction (Figure 3C). In addition, PP2, the inhibitor of Src-family tyrosine kinases, blocked CVX-induced LC3II reduction (Figure 3C). Together, our results demonstrate that platelet agonists induce LC3II reduction, which requires known elements of platelet activation signaling cascades; for example, phospholipase C, Ca²⁺, protein kinase C, and Src-family kinases.
Agonist-induced LC3II reduction involves autolysosomal proteolysis

To determine whether agonist-induced LC3II reduction involved proteolysis, we preincubated human platelets with a protease inhibitor cocktail (PIC) before stimulation. The PIC pretreatment blocked thrombin-, PAR1-, and CVX-induced LC3II reduction in platelets (Figure 4A). Although inhibition of thrombin-induced reduction in LC3II could be attributed to a direct effect of serine protease inhibitors in the PIC on thrombin, inhibition of PAR1- and CVX-induced LC3II reduction was likely a result of inhibition of proteolysis unrelated to thrombin. The ubiquitin–proteasome system and autophagy are 2 major degradation pathways in eukaryotes. The ubiquitin–proteasome system has been reported to be important for thrombopoiesis and platelet function.63,64 We asked whether agonist-induced LC3II reduction involved proteasomes. Inhibition of proteasomes with MG132 had no effect on CVX-induced LC3II reduction in mouse platelets (supplemental Figure 4A),
VAMP2/3 lysosome fusion. Notably, the resting platelets puriﬁed from mice lacking 3 of the 4 major VAMPs (ie, VAMP2/3/8) showed greatly impaired thrombin-induced LC3II reduction, whereas those deﬁcient in either VAMP2/VAMP3 or VAMP8 showed no overt impairment relative to control platelets, which expressed normal levels of VAMP2, VAMP3, and VAMP8, from mice with defective membrane fusion machinery. Platelets from mice with defective membrane fusion machinery. Platelets lacking 3 of the 4 major VAMPs (ie, VAMP2/3/8−/−) showed greatly impaired thrombin-induced LC3II reduction, whereas those deﬁcient in either VAMP2/VAMP3 or VAMP8 showed no overt impairment (Figure 4C). These results suggest that VAMP2, VAMP3, and VAMP8 have redundant function in agonist-induced autophagosome–lysosome fusion. Notably, the resting platelets puriﬁed from VAMP2−/−, VAMP3−/−, and VAMP8/3/8−/− mice had increased LC3II levels relative to control platelets, which expressed normal levels of VAMP2, VAMP3, and VAMP8, with VAMP2/3/8−/− platelets having the highest LC3II levels (Figure 4C). This is consistent with VAMP-mediated autophagosome–lysosome fusion being required for basal autophagy in resting platelets. Together, our results suggest that agonist-induced LC3II reduction likely results from platelet autolysosomal proteolysis; that is, canonical degradative autophagy is induced upon platelet activation.

Of note, a new role of autophagy in unconventional secretion is emerging recently. We asked whether exocytosis of LC3II-decorated exosomes occurred upon agonist stimulation. We activated mouse platelets with CVX, PAR4 peptide, or thrombin before separating platelet-containing pellets from releasate-containing supernatants. PF4, a known α-granule cargo, was the releasate control. Despite the appearance of PF4, LC3II was not detected in the supernatants of activated platelets (supplemental Figure 4B), suggesting the absence of agonist-induced exocytosis of LC3II-decorated exosomes.

Platelet spreading (data not shown), or clot retraction (either in the absence or presence of glucose; supplemental Figure 5D). In summary, we detected modest aggregation and granule cargo packaging defects, but normal cargo release and integrin signaling in the Atg7−/−;PF4-Cre/+ platelets.

Atg7-deﬁcient platelets are deﬁcient in ex vivo platelet aggregation and granule cargo packaging

Given the observed effects activation exerted on autophagy, we asked whether platelet autophagy had a functional role. We generated a mouse strain with a megakaryocyte- and platelet-speciﬁc deletion of Atg7 (ie, Atg7+/-;PF4-Cre/+). Although Atg7 levels were only partially reduced in platelets from Atg7−/−;PF4-Cre/+ mice, LC3II was largely depleted (Figure 5A), suggesting Atg7 is likely the rate-limiting enzyme for LC3II production. We then systematically characterized platelets from these Atg7-deﬁcient mice, using standard ex vivo assays. First, we monitored platelet aggregation by lumi-aggregometry and found that Atg7 deﬁciency caused defects in aggregation in response to modest-to-low concentrations of thrombin and collagen (Figure 5B). Second, we assayed thrombin-dependent platelet exocytosis by monitoring secretion of PF4 from α-granules, [3H]-serotonin from dense granules, and β-hexosaminidase from lysosomes. Our data show that the total levels of α-granule cargo PF4, the total uptake of [3H]-serotonin into dense granules, and the total lysosomal β-hexosaminidase activity were slightly, but signiﬁcantly, reduced in platelets from Atg7−/−;PF4-Cre/+ mice (by 15%, 26%, and 18%, respectively; P < .05), suggesting modestly impaired granule cargo packing (Figure 5C). However, Atg7 deﬁciency did not impair percentage release of PF4, [3H]-serotonin, or β-hexosaminidase (supplemental Figure 5A). Consistently, we also found no overt alterations in the levels of the fusion machinery (eg, STX11, SNAP23, VAMP3, and VAMP8) and the α-granule cargo von Willebrand factor (supplemental Figure 5B). Third, to determine the cause of the observed granule cargo packaging defects, we assessed overall morphology of Atg7−/− and Atg7−/−;PF4-Cre/+ platelets by electron microscopy. Our micrographs show that Atg7-proficient platelets (Figure 5Di-iii) and Atg7-deﬁcient platelets (Figure 5Div-vi) had no overt difference in morphology. Of note, we observed isolation membranes in platelets (Figure 5Div-iii, arrowheads and inset). Fourth, we examined integrin inside-out and outside-in signaling and found that Atg7 deﬁciency had no effect on activation-dependent Jon/A (supplemental Figure 5C) or ﬁbrinogen (data not shown) binding, platelet spreading (data not shown), or clot retraction (either in the absence or presence of glucose; supplemental Figure 5D). In summary, we detected modest aggregation and granule cargo packaging defects, but normal cargo release and integrin signaling in the Atg7−/−;PF4-Cre/+ platelets.

Deletion of Atg7 in platelets leads to impaired hemostasis and thrombus formation

As we observed defects in the Atg7-deﬁcient platelets ex vivo, we next determined the consequence of deleting Atg7 in platelets in vivo. Using
the tail-bleeding assay for hemostasis, we showed that 11 (−65%) of 17 of Atg7fl/fl;PF4-Cre+/− mice were unable to stop bleeding within 10 minutes after tail transection, whereas Atg7fl/fl littermates demonstrated normal bleeding time (median, 2.5 minutes) (Figure 6A). Moreover, using the FeCl3-induced carotid injury model, we further showed that 7 (−47%) of 15 of Atg7fl/fl;PF4-Cre+/− mice were unable to form occlusive thrombi within 30 minutes after injury, whereas all Atg7fl/fl littermates could do so (median occlusion time, 5.6 minutes) (Figure 6B). Of note, the occlusion times of Atg7fl/fl;PF4-Cre+/− mice appeared to show a bimodal distribution for an unknown reason (Figure 6B). Because PF4-Cre expression deletes the floxed Atg7 gene at a later stage of megakaryocyte differentiation, it is possible that bleeding in Atg7fl/fl;PF4-Cre+/− mice is a result of defective platelet production, especially given the thrombocytopenia phenotype of the Atg7fl/fl;Vav-Cre+/− mice, where Atg7 is deleted at an earlier stage of the hematopoietic cell lineage. To investigate this possibility, we measured platelet counts and mean platelet volumes as metrics of thrombocytopenia and faulty platelet production. Compared with Atg7fl/fl mice, Atg7fl/fl;PF4-Cre+/− mice had indistinguishable platelet counts (Figure 6C) and mean platelet volumes (Figure 6D), suggesting that defects in Atg7-mediated platelet function or functions, but not thrombocytopenia, accounted for the severe bleeding in Atg7fl/fl;PF4-Cre+/− mice. Together, our results show that even partial reduction of Atg7 protein levels in Atg7fl/fl;PF4-Cre+/− platelets profoundly impairs hemostasis and thrombus formation. As both LC3II depletion and impaired hemostasis and thrombus formation were observed in Atg7fl/fl;PF4-Cre+/− platelets, our results suggest that Atg7 may be a rate-limiting factor in the role that platelet autophagy plays during hemostasis and thrombosis.

**Discussion**

Although autophagy-related mRNAs and proteins have been reported in transcriptomic and proteomic studies, the functional ramifications of autophagy in platelets have only recently begun to be addressed. Here we confirm the presence of many autophagy-related proteins in platelets. Using transgenic mice expressing GFP-fused LC3, Beclin 1, and Atg5, we further show that these autophagy markers

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**Figure 5.** Atg7-deficient platelets showed modest defects in ex vivo platelet function assays. (A) Representative western blot analysis showing reduced Atg7 and depleted LC3II levels in platelets from Atg7fl/fl;PF4-Cre+/− mice compared with platelets from Atg7fl/fl mice. For each genotype, results from 2 different mice are shown. (B) Lumi-aggregometry measurements of platelets from Atg7fl/fl (red traces) and Atg7fl/fl;PF4-Cre+/− (blue traces) mice when stimulated with thrombin (upper) or collagen (lower). Different agonist concentrations were titrated. The images shown are representative of at least 3 independent experiments. (C) Total levels of PF4 (left), total [3H]-serotonin uptake (middle), and total β-hexosaminidase activity (right) in platelets purified from Atg7fl/fl and Atg7fl/fl;PF4-Cre+/− mice. Data represent an average of 21 measurements. *P < .05, **P < .001, Student t test. (D) EM images of platelets from Atg7fl/fl (i-iii) and Atg7fl/fl;PF4-Cre+/− (iv-vi) mice. Samples were prepared by the freeze substitution method. Scale bars, 500 nm. The images shown are representative micrographs from 2 independent experiments.
autophagosomes and is itself degraded in autolysosomes, have also
actional. The levels of p62, an adaptor protein that delivers substrates to
may exceed the time during which cultured platelets are fully fun-
chase labeling. This assay is not applicable to platelets, as chase time
assay measures the long-lived proteind e g r a d a t i o nr a t e st h r o u g hp u l s e-
platelet activation-induced autophagy. A classical autophagic
biogenesis, increased degradation, or both, a standard assay for evalu-
reduction in human platelets. We postulate that the L7543 antibody
dected thrombin-induced LC3II reduction in both human and murine
 lethal agonist-induced LC3II reduction as a readout for agonist-
activating phenotype reported by Feng et al, in a global heterozygous
were shown to cause abnormal megakaryopoiesis or thrombopoiesis, suggesting key degradation processes may be needed for pro-
expression caused by lack of autophagy in other cell types. The bleed-
formation without thrombocytopenia. (A) Tail-bleeding
time measurements taken from $\text{Atg7}^{+/+}$ (C) and $\text{Atg7}^{+/+}$
(+) mice. For A and B, each data point represents an individual mouse, with the horizontal line marking the median. Statistical signifi-
endothelial cells, although it caus es no abnormality in platelets,
consistent, other studies have shown that loss of autophagy in en-
selecting cargoes to autophagosomes, we cannot rule out the possibility
that agonist-induced platelet autophagy uses adaptors other than p62. Together, our work establishes agonist-induced LC3II reduc-
tion as a useful measure for platelet activation-induced autophagy.
To our knowledge, this is the first report of autophagy induction by platelet activation.

Deleting $\text{Atg7}$ early in platelet production by Vav-Cre$^{39}$
prolonged platelet proteasome inhibition by genetic means$^{63}$ have
been shown to cause abnormal megakaryopoiesis or thrombopoi-
ysis. The use of the $\text{PF4-Cre}$ system allows us to examine the effects of autophagy
deficiency on platelets and hemostasis without the complication of
deficient platelet production.

The use of $\text{PF4-Cre}$ also allows us to examine the effects of autophagy deficiency on platelets and hemostasis without the compi-
lation caused by lack of autophagy in other cell types. The bleeding
phenotype reported by Feng et al, in a global heterozygous $\text{Becn1}$ mouse,$^{40}$ is likely the culmination of Beclin 1 depletion in several cell types; for example, endothelial cells and platelets. Consis-
tently, other studies have shown that loss of autophagy in end-
these results, decreases secretion of von Willebrand factor, prolongs thrombus
formation times, and yields smaller thrombi.$^{71,72}$ Our work is the
first direct report of platelet autophagy being essential for hemo-
stasis and thrombosis. Our work clearly demonstrates that platelet
$\text{Atg7}$ is indispensable and likely a rate-limiting factor for platelet
autophagy’s role in hemostasis and thrombosis. This is confirmed

are localized to punctate structures in platelets. We demonstrate that autophagic flux occurs in resting platelets and is enhanced in response to various platelet agonists. Using mice with $\text{PF4-Cre}$-driven, megakaryocyte/
platelet-specific $\text{Atg7}$ deletion, we further show that platelet autophagy contributes to hemostasis and thrombus formation. Collectively, our data demonstrate physiological relevance of autophagy in platelets and add autophagy to a growing list of cellular processes (eg, translation, secretion, cytoskeletal rearrangements) that are unexpectedly important for platelet functions.

Feng et al reported that platelet autophagy was independent of activation.$^{40}$ We tested several anti-LC3 antibodies, including the one used by Feng et al (ie, Sigma L7543). All but the L7543 antibody detected thrombin-induced LC3II reduction in both human and murine platelets; even the L7543 antibody detected thrombin-induced LC3II reduction in human platelets. We postulate that the L7543 antibody may recognize a subpopulation of mouse LC3II that is less affected by platelet activation. As LC3II reduction could result from decreased biogenesis, increased degradation, or both, a standard assay for evaluating autophagic flux is to monitor changes in LC3II levels in the absence vs presence of lysosomal inhibitors.$^{66,67}$ Using NH$_4$Cl, we confirmed agonist-induced LC3II reduction as a readout for agonist-induced enhancement of autophagic flux. Other potential metrics of autophagic flux, however, appear to be inappropriate for monitoring platelet activation-induced autophagy. A classical autophagic flux assay measures the long-lived protein degradation rates through pulse-chase labeling. This assay is not applicable to platelets, as chase time may exceed the time during which cultured platelets are fully functional. The levels of p62, an adaptor protein that delivers substrates to autophagosomes and is itself degraded in autolysosomes,$^{68}$ have also been used as indicators of autophagic flux in certain cell types.$^{18,69,70}$ However, p62 did not accumulate in $\text{Atg7}$-deficient platelets (supplemental Figure 5B). As p62 is one of many adaptors that bring

Figure 6. Megakaryocyte- and platelet-specific $\text{Atg7}$
deletion causes impaired hemostasis and thrombus formation
without thrombocytopenia. (A) Tail-bleeding
time measurements taken from $\text{Atg7}^{+/+}$ (C) and $\text{Atg7}^{+/+}$
$\text{PF4-Cre}^/+$ (filled gray triangles) mice. (B) Occlusion
time measurements taken from $\text{Atg7}^{+/+}$ (C) and $\text{Atg7}^{+/+}$
$\text{PF4-Cre}^/+$ (filled gray triangles) mice in the FeCl$_3$
carotid injury model of thrombosis. For A and B, each
point represents an individual mouse, with the horizontal line marking the median. Statistical signifi-
cance was evaluated with the log-rank test using SigmaPlot12 software. (C) Platelet counts and (D) mean
platelet volumes from $\text{Atg7}^{+/+}$ (C) and $\text{Atg7}^{+/+}$/$\text{PF4-Cre}^/+$
(filled gray triangles) mice. For C and D, statistical significance was evaluated with the Student $t$ test. NS, not significant.
by our unpublished data showing similar bleeding diatheses in mice with megakaryocyte/platelet-specific deletion of other key autophagy genes; for example, Beclin1\textsuperscript{BAC}PF4-Cre/+ and Pik3c3\textsuperscript{BAC}PF4-Cre/+ (data not shown).

Despite robust in vivo phenotypes, megakaryocyte/platelet-specific Atg7 deficiency only modestly affected aggregation and granule cargo packaging and had limited effects on cargo release or integrin signaling. We performed additional ex vivo platelet assays to examine effects of platelet Atg7 loss on phosphatidylserine (PS) exposure and energetics. Both Atg5 and Beclin 1 are essential for apoptotic cell clearance during development by mediating expression of extracellular PS.\textsuperscript{73} As PS exposure is also important for hemostasis,\textsuperscript{74,75} we thought that autophagy-mediated PS exposure in platelets might be required for hemostasis. Unfortunately, this hypothesis is unlikely, as our studies with annexin V (data not shown) and lactadherin (supplemental Figure 5E) binding showed no impairment in activation-dependent PS exposure on Atg7\textsuperscript{-}/PF4-Cre/+ platelets. As the conventional role of autophagy is for adenosine triphosphate production and nutrient recycling through catabolism, and autophagy is essential for the clearance of damaged organelles such as mitochondria, we speculated that platelet autophagy may be critical for energetics. However, there was no clot retraction defect observed in the Atg7-deficient platelets, even when glucose was withheld (supplemental Figure 5D), suggesting Atg7-deficiency does not impair glucose use in platelets.

At this stage, we have insufficient data to define the mechanism by which Atg7-deficient mice demonstrate severe hemostasis and thrombosis defects. There are ample published examples in which deletion of a signaling cascade protein causes only modest defects in platelet activation in ex vivo assays, yet the protein contributes significantly to in vivo hemostasis and thrombosis.\textsuperscript{56,76-79} In a recent example, STXBPS deletion yielded modest defects on dense granule release but was absolutely required for hemostasis in the tail transection model and for thrombosis in the FeCl\textsubscript{3}-induced carotid injury model.\textsuperscript{80} Therefore, it is possible that the sensitivity of the ex vivo assays we employed is insufficient to fully delineate the effects of deleting the autophagy machinery. There may be an additive or even synergistic effect resulting from several singularly mild impairments.

Alternatively, it is possible that autophagy deficiency may cause some unidentified platelet dysfunction, leading to the observed severe hemostasis and thrombosis defects. Our current hypothesis is that platelet autophagy-mediated degradation of a specific substrate or substrates is required for effective hemostasis and thrombosis. Consistent with this concept, there are functional similarities between the 2 major degradation systems: the ubiquitin–proteasome system and autophagy in platelets. Both occur in resting platelets and are activated platelet autophagy-mediated degradation of a specific substrate is required for effective hemostasis and thrombosis defects. Our current hypothesis is that platelet autophagy-mediated degradation of a specific substrate or substrates is required for effective hemostasis and thrombosis. Consistent with this concept, there are functional similarities between the 2 major degradation systems: the ubiquitin–proteasome system and autophagy in platelets. Both occur in resting platelets and are activated.

Note in added proof. While this manuscript was under review, a case report was published illustrating autophagosome-like structures in platelets from a dog with severe nonregenerative anemia.\textsuperscript{85} Moreover, another work using hematopoietic system conditional Atg7 knockout (ie, Atg7\textsuperscript{Pp}Vav-Cre) mice was published that demonstrates that in addition to severe defects in megakaryogenesis, megakaryocyte differentiation and thrombopoiesis observed in these mice, platelets isolated from these mice are defective in activation and aggregation.\textsuperscript{90} Our work using Atg7\textsuperscript{Pp}PF4-Cre mice, which deletes the Atg7 genes at a later stage in platelet production, distinguishes the roles of autophagy in platelets and in hemostasis from roles in megakaryogenesis and in megakaryocyte differentiation.

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Authorship

Contributions: Q.J.W. conceived the project and coordinated the study. M.M.O., Y.H., Z.L., B.S., S.W.W., and Q.J.W. designed the experiments. M.M.O., Y.H., M.B., S.J., L.M., Y.Z., H.L., X.L., B.X., G.Z., M.K., Z.Y., Z.L., S.W.W., and Q.J.W. carried out the experiments. M.M.O., Y.H., M.B., S.W.W., and Q.J.W. analyzed the data. Q.J.W. wrote the manuscript, and S.W.W. helped with the editing. For experiments, both M.M.O. and Y.H. contributed to all figures; M.B. contributed to the aggregometry and cargo release assays; S.J. and L.M. contributed to the electron microscopic analysis; S.J. also contributed to the generation of the VAMP-deficient platelets; H.L. and S.W.W. contributed to the in vivo tail bleeding assay; G.Z. and H.L. contributed to the FeCl\textsubscript{3} carotid injury model. Y.Z. contributed to Figures 1-3; H.L., Z.L., B.X., and G.Z. contributed to Figures 5-6; X.L., Q.J.W., and Z.Y. generated the EGFP-Atg5\textsuperscript{+/} BAC transgenic mice; Z.Y. also provided the Beclin1-EGFP+/ BAC transgenic mice; and M.K. and M.M.O. generated the Atg7\textsuperscript{Pp}PF4-Cre/+ mice. The authors declare no competing financial interests.

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Autophagy is induced upon platelet activation and is essential for hemostasis and thrombosis

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