Background and Purpose—For stroke prevention, patients with atrial fibrillation typically receive oral anticoagulation. The commonly used anticoagulant warfarin increases the risk of hemorrhagic transformation (HT) when a stroke occurs; tissue-type plasminogen activator treatment is therefore restricted in these patients. This study was designed to test the hypothesis that 12/15-lipoxygenase (12/15-LOX) inhibition would reduce HT in warfarin-treated mice subjected to experimental stroke.

Methods—Warfarin was dosed orally in drinking water, and international normalized ratio values were determined using a Coaguchek device. C57BL/6J mice or 12/15-LOX knockout mice were subjected to transient middle cerebral artery occlusion with 3 hours severe ischemia (model A) or 2 hours ischemia and tissue-type plasminogen activator infusion (model B), with or without the 12/15-LOX inhibitor ML351. Hemoglobin was determined in brain homogenates, and hemorrhage areas on the brain surface and in brain sections were measured. 12/15-LOX expression was detected by immunohistochemistry.

Results—Warfarin treatment resulted in reproducible increased international normalized ratio values and significant HT in both models. 12/15-LOX knockout mice suffered less HT after severe ischemia, and ML351 reduced HT in wild-type mice. When normalized to infarct size, ML351 still independently reduced hemorrhage. HT after tissue-type plasminogen activator was similarly reduced by ML351.

Conclusions—In addition to its benefits in infarct size reduction, 12/15-LOX inhibition also may independently reduce HT in warfarin-treated mice. ML351 should be further evaluated as stroke treatment in anticoagulated patients suffering a stroke, either alone or in conjunction with tissue-type plasminogen activator. (Stroke. 2017;48:445-451. DOI: 10.1161/STROKEAHA.116.014790.)

Key Words: atrial fibrillation ▪ brain ▪ hemorrhage ▪ stroke ▪ warfarin

Atrial fibrillation increases the risk of stroke, which is a leading cause of death and disability worldwide. The prevalence of atrial fibrillation was between 2.7 and 6.1 million cases in 2010 and is estimated to rise ≥12.1 million by 2030. Administration of long-term oral anticoagulation reduces the risk of ischemic stroke in patients with atrial fibrillation. However, the use of the vitamin K antagonist warfarin can be challenging because it significantly increases the risk of hemorrhagic transformation (HT) in patients with stroke. Warfarin-treated patients with effective anticoagulation, that is, an international normalized ratio (INR) value >1.7 are contraindicated for tissue-type plasminogen activator (tPA) treatment.

Accumulating data suggest that increased permeability of the blood–brain barrier (BBB) after ischemia is one of the major causes of HT. We have previously shown that after experimental stroke, 12/15-lipoxygenase (12/15-LOX) is increased in both neurons and vascular endothelial cells in the peri-infarct region. 12/15-LOX inhibition or knockout reduced the loss of endothelial tight junction protein claudin-5 and the leakage of IgG after transient ischemia, both evidence for a role of 12/15-LOX in the disruption of the BBB seen after an ischemic event. We also demonstrated that the 12/15-LOX inhibitor LOXBlock-1 significantly reduced BBB leakage after tPA thrombolysis in a distal clot mouse model.

We have recently introduced ML351 as a novel inhibitor of 12/15-LOX, which has shown effective neuroprotection both in vitro and in vivo. On the basis of an established animal model of warfarin-associated HT, this study was designed to investigate whether inhibition of 12/15-LOX or its gene...
knockout reduces warfarin-induced HT after experimental stroke. Furthermore, we evaluated whether 12/15-LOX inhibition reduces the risk of excess HT caused by thrombolytic treatment with tPA in warfarin-anticoagulated mice.

Materials and Methods

Mouse Model of Oral Anticoagulation
Male C57BL/6J mice (12 weeks old) from Jackson Laboratories and Alox15 knockout mice bred in our animal facility (in which the gene encoding 12/15-LOX is deleted) were used in accordance with the National Institutes of Health Guide for the Care and Use of Laboratory Animals. After an established protocol,15,16 we administered warfarin by oral uptake through bottled drinking water. Briefly, different doses of Coumadin tablet (0, 1.25, 2.5, and 5 mg; warfarin sodium, crystalline; Bristol Myers Squibb, Munich, Germany) were dissolved in 375 mL tap water, and the mice were fed for 3 hours with free access to the treated water. The INR measurements were performed on venous blood sample drawn from the tail vein by a commercially available point of care coagumeter (Coaguchek XS, Roche, Mannheim, Germany) immediately after warfarin withdrawal.

Immunohistochemistry
Mice were treated with warfarin and subjected to MCAO as indicated. After 24 hours, anesthetized mice were perfused transcardially with ice-cold PBS (pH 7.4), followed with ice-cold 4% paraformaldehyde in PBS (pH 7.4). The brains were removed, immersed in the same fixative overnight at 4°C, and cryoprotected in 15% and 30% sucrose solutions in PBS at 4°C. Frozen coronal sections (20 μm thick) were prepared using a cryostat. After blocking with PBS containing 0.2% Triton X-100 and 3% bovine serum albumin, sections were incubated at 4°C overnight in a PBS/0.2% Triton X-100/2% bovine serum albumin solution with an affinity-purified rabbit polyclonal antibody directed toward the C terminus of 12/15-LOX (characterized in Peckce et al,14 diluted 1:100), followed by incubation with an Alexa-488-labeled secondary anti-rabbit antibody (Invitrogen, 1:200). Brain sections from 3 mice/group were imaged using a Nikon Eclipse Ti fluorescent microscope with NIS Elements software, keeping the settings constant. For each mouse, 6 images were taken from cortex and striatum, respectively, of both hemispheres, giving a total of 24 images per mouse. Signal intensity was measured for the ipsilateral brain sections using NIH ImageJ software and normalized as the ratio of Max/Mean intensity.

Statistical Analysis
The Wilcoxon–Mann–Whitney test was used to compare data in 2 groups. For tests with ≥3 groups, ANOVA with Dunnett test was used. Data taken over time were analyzed with 2-way ANOVA. All data are expressed as mean±SEM. Probability values <0.05 were considered significant.

Results

In a Model of Severe Ischemia in Conjunction With Warfarin Pretreatment, 12/15-LOX Knockout Mice Show Less Hemorrhage Than the Corresponding Wild-Type Mice

In a preliminary study, we determined a warfarin dose to provide effective and safe anticoagulation in C57Bl6 mice, according to Foerch et al.18 The INR values after 24 hours of warfarin treatment increased in a dose-dependent manner (Figure 1A). Whereas 5 mg warfarin added to the drinking water resulted in a very high INR value at the detection limit of the Coaguchek measuring device, the 2.5 mg dose of warfarin led to an INR value of 3.21±0.32 (mean±SEM.), and we used this dosage for all subsequent experiments. We next established 2 models of warfarin-associated HT, characterized by severe (3 hours) focal ischemia (experiment A in Figure 1B) or by 2 hours of MCAO followed by tPA thrombolysis (experiment B). To determine whether or not the absence of 12/15-LOX reduces hemorrhage in an established model of anticoagulation followed by severe ischemia,15 after pretreatment with warfarin we subjected 12/15-LOX knockout mice (in which the ALOX15 gene has been deleted) and the corresponding wild-type mice to 3 hours of MCAO and 21 hours of reperfusion. Using a spectrophotometric hemoglobin assay, we observed high levels of hemorrhage in the ipsilateral hemispheres of wild-type mice (Figure 1C). In contrast, the 12/15-LOX knockout mice showed...
significantly less hemorrhage, suggesting that the absence of 12/15-LOX is protective. This was not because of any effect on the coagulation system because the INR values were equal in both cohorts of mice (12/15-LOX knockout 3.14±0.06; wild-type 3.20±0.06). Fewer knockouts died than wild-type mice (wild type 5/10, 12/15-LOX knockout 3/10).

Reduced HT in Anticoagulated Mice After Administration of a 12/15-LOX Inhibitor

We next attempted to replicate these findings with pharmacological treatment. We have recently introduced a novel, highly selective inhibitor of 12/15-LOX. ML351 reduced infarct size in a permanent focal ischemia model but had not been tested in a transient focal ischemia model at the time. Using the classical filament model of transient focal ischemia with 2 hours of occlusion, we now found that ML351 reduced infarct size in a dose-dependent manner, with significant reductions at 25 and 50 mg/kg (Figure in the online-only Data Supplement). Because the 50 mg/kg dose was more effective, with an infarct size reduction of 43%, we used this dosage for subsequent experiments with warfarin anticoagulation. We administered ML351 to warfarin-pretreated mice at the time of reperfusion 3 hours after onset of ischemia. These mice developed significantly less hemorrhage on the ischemic side of the brain, compared with vehicle-treated mice (Figure 2A). In a separate study but using the same experimental parameters, we changed the workup procedure to include imaging of the hemorrhage both on the surface of the brain and in coronal brain sections. Mice with effective oral anticoagulation from the vehicle-treated group showed massive hemorrhages in brain sections after MCAO (Figure 2B). This was again significantly reduced in the mice treated with ML351. Subsequent triphenyltetrazolium hydrochloride staining showed that this was accompanied by a slight but statistically significant

Figure 1. A, Increasing amounts of warfarin added to the drinking water of mice led to increased international normalized ratio (INR) values 24 h later (**P<0.01, ***P<0.001 compared with no added warfarin; n=6 per group). B, Experimental setup with 24 h warfarin pre-administration, followed by either 3 h of MCAO (experiment A) or 2 h of MCAO, followed by tissue-type plasminogen activator (tPA) treatment 1 h after reperfusion (experiment B). Timeline not to scale. C, Photometrically determined hemoglobin levels in the ipsilateral brain hemisphere after warfarin pretreatment and 3 h of middle cerebral artery occlusion (MCAO) were significantly lower in 12/15-lipoxygenase (12/15-LOX) knockout mice compared with matching wild-type mice (**P<0.01; n=5 per group). Expt indicates experiment.

Figure 2. A, Significantly reduced hemoglobin levels in C57Bl6 mice treated with the 12/15-lipoxygenase (12/15-LOX) inhibitor ML351 after warfarin and 3 h of middle cerebral artery occlusion (MCAO), compared with vehicle treatment (**P<0.01). B, Less hemorrhage was detected in brain sections of mice treated with vehicle and ML351 (**P<0.01). C, Infarct size comparison after triphenyltetrazolium hydrochloride staining in mice subjected to warfarin plus 3 h of MCAO showed large infarcts in the vehicle-treated mice, which were slightly but significantly reduced after ML351 treatment (**P<0.05). D, After normalization for infarct size, hemorrhage remained significantly reduced by ML351 treatment (*P<0.05). tPA indicates tissue-type plasminogen activator.
reduction in infarct size (Figure 2C). Mortality was also lower in the inhibitor-treated mice, but the effect was not statistically significant (Vehicle 7/17, ML351 4/17; \( P = 0.464 \)).

**ML351 May Be Associated With Reduced HT Independently of Its Ability to Reduce Infarct Size**

One important issue arising here is whether or not the reduced hemorrhage is because of the reduction in infarct size, rather than a specific protective effect of ML351 on the brain vasculature. To address this question, we normalized the data by dividing the hemorrhage values by the infarct size determined for the same mouse. This normalized hemorrhage still showed a significantly reduced value in the ML351-treated mice (Figure 2D), suggesting that 12/15-LOX inhibition protects the vasculature against leakage in this anticoagulant model of severe ischemia, in addition to its neuroprotective effects.

**Lipoxygenase Inhibition Reduces HT Associated With tPA Treatment in Warfarin-Treated Mice**

Compounding the increased risk of HT, another major factor leading to increased hemorrhage under warfarin anticoagulation is the use of thrombolytics, particularly tPA. To test if 12/15-LOX inhibition can provide a benefit under anticoagulation with thrombolysis, we subjected mice to 24 hours of anticoagulant treatment, followed by 2 hours of MCAO and intravenous infusion of tPA 1 hour later. Treatment with either ML351 or vehicle was initiated by IP injection at the time of tPA infusion, and mice were euthanized 24 hours after the onset of ischemia (Figure 1B). In 2 separate cohorts of mice, we determined either the hemorrhage volume photometrically (Figure 3A) or the hemorrhages in brain sections (Figure 3B). In both measurements, treatment with ML351 led to a significant reduction in hemorrhage. In addition, fewer mice died in the ML351 group, but again the difference in mortality was not statistically significant (Vehicle 7/17, ML351 5/18; \( P = 0.488 \)). Taken together, these results indicated that ML351 reduced the HT that occurred subsequent to tPA infusion in warfarin-treated mice.

**Moderate Anticoagulation With Warfarin Leads to Increased Expression of 12/15-LOX After MCAO**

To determine if warfarin pretreatment impacts the level of 12/15-LOX after experimental stroke, we compared 12/15-LOX signal in brain sections from mice pretreated with warfarin to mice without warfarin pretreatment, euthanized after 2 hours of MCAO and 22 hours of recovery. A third cohort of mice was pretreated with warfarin, followed by 2 hours of MCAO and subsequent infusion of tPA. As expected from our previous studies, immunohistochemistry showed increased 12/15-LOX protein on the ipsilateral side of the brains of mice subjected to MCAO under all 3 conditions (data not shown), but the 12/15-LOX signal was significantly higher in

![Figure 3.](http://stroke.ahajournals.org/)

**Figure 3.** A, After warfarin pretreatment, then 2 h of middle cerebral artery occlusion (MCAO), and tissue-type plasminogen activator (tPA) infusion 1 h later, IP ML351 delivered concomitantly reduced hemoglobin levels in the ipsilateral side of the brain at 24 h (*\( P < 0.05 \)). B, Hemorrhage in brain sections was significantly reduced by treatment with ML351 (*\( P < 0.05 \); \( n = 5 \) for vehicle, \( n = 7 \) for ML351). C, 12/15-lipoxygenase (12/15-LOX) protein, detected by immunohistochemistry, was increased in the brains of warfarin-pretreated mice (**\( P < 0.001 \), *\( P < 0.05 \) compared with MCAO only; \( n = 3 \) per group, 24 images per brain) after MCAO. D, Representative images show that most of the increased 12/15-LOX signal after warfarin treatment is vessel derived (scale bar=100 \( \mu m \)). W, warfarin.
the warfarin model mice (Figure 3C). Representative images taken from the ischemic hemisphere indicated that the signal is mostly increased in the vasculature, consistent with the hypothesis that increased vascular 12/15-LOX may contribute to HT (Figure 3D).

Long-Term Effects of 12/15-LOX Inhibition Under Warfarin Anticoagulation

In a first attempt to gauge the effects of 12/15-LOX inhibition on extended outcome beyond the 24-hour time window, we subjected mice to warfarin pretreatment, followed by 3 hours of MCAO. Only 2 of 22 mice lived past day 2 and none longer than 4 days (data not shown), suggesting that this model in our hands is too severe for long-term outcome studies. We next reduced the MCAO period to 2 hours, allowing for improved survival. Unfortunately, using a new preparation of warfarin, some of the mice developed unusually high INR values, and we excluded mice with INR >4 from the study, leaving 7 mice per group. With 2 hours of MCAO after warfarin pretreatment, survival was better in the ML351 treatment group, although the difference was not statistically significant (Figure 4A). The behavioral score using a modified Garcia Score according to Imai et al.20 showed a significant benefit for the ML351 treatment group (Figure 4B).

![Figure 4. A. Seven days after warfarin plus 2 h of middle cerebral artery occlusion (MCAO) treatment, survival was better in ML351-treated animals, but the difference was not statistically significant (P=0.24; n=7 per group). B. Behavioral outcome was significantly improved in the ML351 treatment group (P<0.05 by 2-way ANOVA).](http://stroke.ahajournals.org/)

Discussion

Although warfarin greatly reduces the risk of cardioembolic ischemic stroke, many patients with atrial fibrillation nonetheless experience a stroke while under anticoagulation.21-22 In addition, warfarin increases the risk of HT in case of stroke.23 However, no effective treatment for this warfarin-associated HT has been identified. Both our laboratory and others have previously shown that 12/15-LOX enzyme inhibition or gene knockout reduces ischemic injury in animal models of transient MCAO.11,12,24 In this study, we demonstrate for the first time that the 12/15-LOX inhibitor ML351 effectively protects against HT in 2 mouse models of warfarin-associated hemorrhage after ischemic stroke.

We have previously found that ML351 significantly reduced infarct size after permanent focal ischemia in a mouse model of ischemic stroke.19 In this study, we found that ML351 also has a protective effect on infarction in transient MCAO (Figure in the online-only Data Supplement). With 43% infarct size reduction when used at 50 mg/kg, the degree of protection afforded by ML351 is comparable to our previous findings with other 12/15-LOX inhibitors, as well as with infarct size reductions in the 12/15-LOX knockout mice (30%-42% in MCAO studies with 90 minutes or 2 hours of occlusion).11,13 reinforcing the notion that 12/15-LOX is a valid target for neuroprotection. Given the severity of 3 hours of occlusion under conditions of warfarin anticoagulation, we were somewhat surprised to see that ML351 could still reduce the infarct size, even with these very large infarcts (Figure 2C). In human stroke patients, an infarct size threshold around 70 cm³ is proposed where current stroke treatments are no longer capable of providing a benefit.25 Our results suggest that a 12/15-LOX inhibitor may provide neuroprotection across a fairly broad range of stroke severity. Under warfarin anticoagulation, HT in the ML351 group was significantly lower than in the vehicle group (Figure 2A and 2B), and this protection against HT remained significant even after adjusting for the different infarct sizes (Figure 2D). Our findings suggest that ML351 can reduce both the infarct lesion and HT under effective anticoagulation, and these 2 effects may be independent of each other. Mortality in each experiment favored the group with reduced 12/15-LOX, although the differences did not reach statistical significance.

The safety of tPA in acute ischemic stroke patients treated with warfarin is a topic of controversy.23,26,27 The current American Heart Association/American Stroke Association guidelines accept tPA treatment for patients with an INR ≤1.7 as a measure of limited anticoagulant effect.28 However, the reality of the situation is that tPA is the only potentially available therapy for patients under effective anticoagulation (with INR value 2–4) in case of a stroke. We have previously shown that another 12/15-LOX inhibitor, LOXBlock-1, reduced tPA-associated hemorrhage in a thrombotic mouse model.13 Therefore, 12/15-LOX might be a therapeutic target for HT associated with tPA treatment. In this study, we investigated the effect of ML351 when combined with tPA in a warfarin-associated HT model. We found that ML351 administered at the same time as tPA at a clinically relevant time point can significantly reduce HT (Figure 3). Several compounds, including
a blocker of the prostaglandin receptor EP1,29 an activator of Nrf2,20 as well as recombinant annexin 2,30 have been shown to reduce BBB leakage in rodent models of ischemic stroke. Aside from blocking the activity of warfarin itself,6 this is to our knowledge the first report of protection against tPA-induced HT in the context of warfarin anticoagulation.

It has long been considered that the main mechanism of HT subsequent to ischemic stroke is the leakage of blood by the disruption of BBB.31,32 Although the precise mechanisms that mediate BBB disruption and HT after ischemic insults are complex, there is likely to be a strong correlation with oxidative stress, and 12/15-LOX might be a major contributor.33,34 We previously reported that the expression of 12/15-LOX was increased in endothelial cells after transient focal ischemia, and its inhibitor baicalin reduced the loss of the endothelial tight junction protein claudin-5 after ischemia. We also demonstrated that 12/15-LOX knockout mice had less leakage of IgG into the brain parenchyma, as did animals treated with inhibitor.13 In this study, we measured hemispheric swelling as 1 manifestation of edema formation in a subset of mice with warfarin pretreatment and 3 hours of MCAO. We found no statistically significant difference for the ratio of ipsilateral to contralateral area in the mice treated with ML351 (1.13±0.03 versus 1.18±0.04 for vehicle-treated mice, P=0.31; n=5 per group). However, we agree with the reviewers that effects on edema can potentially be important as well, and future studies with larger N may be warranted. Warfarin pretreatment led to increased 12/15-LOX protein levels after stroke, especially in the brain vasculature (Figure 3C and 3D). This does not automatically mean that warfarin itself upregulates 12/15-LOX, but it supports the idea that increased vascular 12/15-LOX contributes to BBB leakage leading to the increased HT and provides a rationale for how 12/15-LOX inhibition may reduce HT.

There are several limitations to our study. First, we purposefully chose severe models of stroke and stroke with thrombolysis, to obtain high levels of hemorrhage. Whether or not a similar benefit would be apparent with less severe strokes remains to be seen. A first indication that this benefit is present can be taken from our experiment with 7-day survival where we combined warfarin with 2 hours of MCAO (Figure 4), but the relative contributions of infarct size reduction versus protection against HT in that setting need to be further studied. Second, we here used the classical filament model of MCAO, so directly thrombus-related effects on HT are not evaluated in this study. A thrombin injection model with subsequent tPA thrombolysis has been established,35 but its use in conjunction with warfarin pretreatment may be challenging. We saw very similar protection against HT through 12/15-LOX inhibition in a distal MCAO thrombosis model, however, so the benefits of ML351 may be broadly applicable.13 Third, the results shown here are specific to the use of warfarin as anticoagulant. Clinically, besides warfarin, new oral anticoagulants, including dabigatran, rivaroxaban, or apixaban, are now in use. Further studies will be needed to evaluate the effects of 12/15-LOX inhibition in the context of these novel anticoagulants. Finally, although it seems likely that a reduction in HT provides a sustained benefit for warfarin-treated animals, this will have to be probed further in additional long-term outcome experiments with more extensive behavioral testing.

In summary, our results have shown that the novel inhibitor of 12/15-LOX, ML351, significantly reduced warfarin-associated HT with and without tPA administration. These results suggest that ML351 could be a candidate for the treatment of anticoagulated patients suffering a stroke, either alone or in conjunction with tPA thrombolysis.

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Disclosures
Dr Foerch received speaker honoraria from Boehringer Ingelheim (modest). A patent for the use of ML351 to treat stroke has been applied for (PCT/US2014/052269). The other authors report no conflicts.

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12/15-Lipoxygenase Inhibition or Knockout Reduces Warfarin-Associated Hemorrhagic Transformation After Experimental Stroke
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Infarct size in the classical filament model of MCAO. C57Bl6] mice were subjected to 90 minutes transient MCAO according to van Leyen et al\textsuperscript{1}. ML351 at the indicated dosage [mg/kg] or an equal volume of vehicle was injected intraperitoneally two hours after onset of ischemia. Following sacrifice at 24 hours, infarct sizes were determined by staining with 2,3,5-triphenyltetrazolium chloride (TTC) (n=8/group; *p<0.05, ***p<0.001).