Lysophosphatidic Acid Receptors 1 and 2 Play Roles in Regulation of Vascular Injury Responses but Not Blood Pressure

Manikandan Panchatcharam, Sumitra Miriyala, Fanmuyi Yang, Mauricio Rojas, Christopher End, Christopher Vallant, Anping Dong, Kevin Lynch, Jerold Chun, Andrew J. Morris, Susan S. Smyth

Abstract—Phenotypic modulation of vascular smooth muscle cells (SMCs) is essential for the development of intimal hyperplasia. Lysophosphatidic acid (LPA) is a serum component that can promote phenotypic modulation of cultured SMCs, but an endogenous role for this bioactive lipid as a regulator of SMC function in vivo has not been established. Ligation injury of the carotid artery in mice increased levels in the vessel of both autotaxin, the lysophospholipase D enzyme responsible for generation of extracellular LPA, and 2 LPA responsive G protein–coupled receptors 1 (LPA1) and 2 (LPA2). LPA1−/−/2−/− mice were partially protected from the development of injury-induced neointimal hyperplasia, whereas LPA1−/− mice developed larger neointimal lesions after injury. Growth in serum, LPA-induced extracellular signal-regulated protein kinase activation, and migration to LPA and serum were all attenuated in SMCs isolated from LPA1−/−/2−/− mice. In contrast, LPA1−/− SMCs exhibited enhanced migration resulting from an upregulation of LPA3. However, despite their involvement in intimal hyperplasia, neither LPA1 nor LPA2 was required for dedifferentiation of SMCs following vascular injury or dedifferentiation of isolated SMCs in response to LPA or serum in vitro. Similarly, neither LPA1 nor LPA2 was required for LPA to elicit a transient increase in blood pressure following intravenous administration of LPA to mice. These results identify a role for LPA1 and LPA2 in regulating SMC migratory responses in the context of vascular injury but suggest that additional LPA receptor subtypes are required for other LPA-mediated effects in the vasculature. (Circ Res. 2008;103:662-670.)

Key Words: arterial injury ■ lipids ■ lysophosphatidic acid ■ vascular remodeling ■ vascular smooth muscle cells

Phenotypic modulation of vascular smooth muscle cells (SMCs) occurs in response to vascular injury and is a critical component in the development of atherosclerotic and restenotic lesions.1,2 Changes in the extracellular environment promote this response, which is characterized by alterations in the differentiation state of SMCs and in their acquisition of the capacity to proliferate and migrate. Isolated vascular SMCs from human and rodent species can be stimulated to dedifferentiate, proliferate, and migrate by serum. The lipid mediator lysophosphatidic acid (LPA) has been proposed as 1 of the factors present in serum that may promote phenotypic modulation of vascular SMCs.3–5

LPA, the simplest glycerophospholipid, promotes dedifferentiation, proliferation,6–8 and migration9 of isolated vascular SMCs. Although the specific signaling systems involved in SMC responses are not known, in other cell systems, LPA acts through at least 5 G protein–coupled receptors, LPA1-5, to stimulate a wide variety of intracellular signaling pathways important in health and disease.10 Prominent among the pathways that are activated by LPA include Rho GTPases and extracellular signal-regulated protein kinase (ERK), which are known to play important roles in vascular SMC function. LPA has also been proposed to serve as an endogenous activator of peroxisome proliferator-activated receptor (PPAR)γ,11 and there may be additional, as yet unidentified receptor targets for LPA.

LPA is produced in locations that position it to be a pathophysiologic mediator of vascular cell function. LPA is present in relatively low levels in plasma but is more abundant in serum, where it is thought to be derived at least in part through processes that require platelet activation.12–18 Therefore, local concentrations of LPA may be increased along vessels at sites of platelet adhesion and thrombus formation. LPA is also found in abundance in the lipid-rich core of atherosclerotic plaque, where it may be derived from mildly oxidized LDL.19 Thus, LPA is present, or can be

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formed, in the settings associated with alterations in SMC function. The lysophospholipase D autotaxin (ATX), which catalyzes the hydrolysis of lysophospholipid substrates, is responsible for generation of biologically active LPA in circulation. Mice that are heterozygous for the wild-type (WT) and null ATX allele have 50% normal circulating LPA levels, and mice that transgenically overexpress ATX in the liver under control of the α-antitrypsin promoter have elevated circulating levels of ATX and LPA.

Exogenous administration of LPA to animals elicits responses that are consistent with it serving as an endogenous mediator of vascular cell function. For example, intravenous injection of LPA elevates arterial blood pressure in rats, and local application causes cerebral vasodilation and constriction in pigs. Moreover, local infusion of LPA in the rat common carotid artery induces vascular remodeling by stimulating neointimal formation. A similar response is observed in mice and may be mediated by PPARγ.

Until recently, a lack of appropriate and specific tools has limited our ability to define pathophysiologic roles of endogenous LPA in the vasculature. In particular, the most extensive in vitro studies reported to date have used rat or human SMCs, whereas responses of murine SMCs to LPA have not been examined in detail. A concerted analysis of murine SMC responses to LPA in vitro, coupled with a phenotypic analysis of vascular injury responses of LPA receptor–deficient mice, is required to provide definitive insights into the role of LPA in this important aspect of vascular function. In this report, we identify an upregulation of ATX and LPA receptors 1 and 2 (LPA1 and LPA2) following vascular injury. We use mice deficient in LPA1 and -2 alone and in combination to define roles for these receptors and LPA in the pathophysiologically relevant vascular responses and in the phenotypic modulation of SMCs that occurs with vascular injury.

Materials and Methods

Mice
All procedures conformed to the recommendations of Guide for the Care and Use of Laboratory Animals (Department of Health, Education, and Welfare publication number NIH 78-23, 1986) and were approved by the Institutional Animal Care and Use Committee. The production and initial characterization of mice deficient in LPA receptors 1 and 2 (LPA1 and LPA2) following vascular injury. We use mice deficient in LPA1 and -2 alone and in combination to define roles for these receptors and LPA in the pathophysiologically relevant vascular responses and in the phenotypic modulation of SMCs that occurs with vascular injury.

Vascular Injury
At various intervals after carotid surgery, 5 mm of aorta proximal to the suture was removed and processed for RNA analysis by qualitative PCR (qPCR) or analysis of protein markers by immunoblotting. Neointimal formation along the length of the vessel was assessed at 4 weeks after surgery using computer assisted morphometry as has been previously described. Digital images were taken with high performance digital camera (resolution 3840×3072 pixels) attached to a Nikon 80i microscope with a ×10 (NA=0.3) or ×20 (NA=0.5) objective and analyzed with Metamorph software.

Femoral artery denudation injury was performed and analyzed at 4 weeks as previously described.

Isolation of SMCs
Mouse aortic SMCs were obtained from thoracic aortas by removing the adventitia and endothelium by digestion with collagenase type II ( Worthington; 175 U/mL). The media were further digested in solution containing collagenase type II (175 U/mL) and elastase (Sigma; 0.5 mg/mL), which yielded 100 000 cells per aorta. Cells were grown in DMEM containing 0.5 ng/mL EGF, 5 μg/mL insulin, 2 ng/mL basic fibroblast growth factor, 10% FBS, 100 U/mL penicillin, and 100 μg/mL streptomycin and incubated at 37°C with 5% CO2/95% air. SMC lineage was confirmed by the presence of immunoreactivity for α-actin (Sigma) in >99% of the cells. Experiments involving SMCs were performed by using cells with a passage number of ≤5.

An expanded Materials and Methods section is available in the online data supplement at http://circres.ahajournals.org.

Statistical Analysis
All results were expressed as mean±SD. In vitro studies were repeated a minimum of 3 times, and results were analyzed by Student t test or ANOVA. Statistical significance within strains was determined using ANOVA with multiple pair-wise comparisons. Statistical analysis was performed using Sigma-STAT software version 3.5 (Systat Software Inc). A probability value of less than 0.05 was considered significant.

Results
Exposure of isolated smooth muscle cells or intact vessels to exogenous LPA elicits SMC phenotypic modulation; however, the role of endogenous LPA in mediating SMC responses in the context of vascular injury is not known. The aim of this study was to determine whether LPA contributes to vascular injury responses in intact organisms. To accomplish this aim, we used mice deficient in candidate LPA receptors and pharmacological antagonists of these receptors to attenuate normal LPA signaling in vascular SMCs and determine the consequences on vascular and isolated cells responses.

Upregulation of ATX and LPA Receptors Following Vascular Injury
Following ligation injury, we observed a time-dependent increase in vessel-associated levels of ATX protein that was 2.5-fold after 1 to 3 days (P<0.05 at 1 day and 3 days versus uninjured; Figure 1A). ATX is a secreted lysophospholipase D responsible for generation of biologically active extracellular LPA. Increasing plasma levels of ATX elevates circulating LPA levels; therefore, extracellular LPA production along the vessel wall likely increases following injury. In many systems, LPA elicits cellular effects via G protein–coupled receptor signaling. We therefore assessed the level of gene expression of the 5 known LPA receptors (LPA1–5) following injury and observed that expression of LPA1 and LPA2 increased following carotid injury by 2.1- and 6.2-fold, respectively (Figure 1B).

LPA1+/−, LPA2+/− Mice Are Partially Protected From Development of Intimal Hyperplasia in Vascular Injury Models, Whereas LPA1+/− Mice Exhibit Positive Remodeling Characterized by an Enhanced Neointimal Response
Mice that are singly or doubly deficient in LPA1 and LPA2 (LPA1−/−, LPA2−/−, and LPA1+/−, LPA2−/− mice) are viable and...
fertile, allowing us to use them to investigate a role for these LPA signaling receptors in the regulation of vascular SMC responses. Carotid ligation was performed in WT, LPA1/−/−, LPA2/−/−, and LPA1/−/−2/−/− double knockout mice. Four weeks after carotid ligation, neointima developed along the length of the carotid in WT mice (Figure 2A and 2B). The extent of intimal area and the intima/media ratio was similar in mice lacking LPA2 (Figure 2B and 2D). In contrast, LPA1/−/− mice displayed an ∼3-fold enhanced intimal area (Figure 2B), an increase in intima/media ratio ($P<0.017$ versus WT; Figure 2D), a larger lumen area (Figure 2E), and a resulting increase in the area within the external elastic lamina, consistent with positive remodeling (ie, maintenance of luminal area) in the presence of heightened neointimal formation. Unlike either the LPA2/−/− or LPA1/−/− mice, which displayed normal and enhanced neointima, respectively, LPA1/−/−2/−/− double knockout were partially protected from the development of intimal hyperplasia with an ∼4-fold lower intimal areas ($P<0.05$; Figure 2B) and intima/media ratios after carotid ligation (Figure 2D). Medial areas were similar in all mice (Figure 2C).

To determine whether these results were model-specific, we denuded the endothelium of the mouse femoral artery to elicit the development of intimal hyperplasia and observed an enhanced injury response that was broadly similar to that observed in the carotid artery ligation model in LPA1/−/− mice (see Figure I in the online data supplement). Additional experiments were not performed with the other strains of mice.

Figure 2. Alterations in the development of intimal hyperplasia in mice lacking LPA receptors. A, Representative combined Masson elastin–stained sections of carotid arteries at 4 weeks after surgery taken ∼1.2 mm from the site of ligation in WT, LPA1/−/−, LPA1/−/−2/−/− mice. B through E, Intimal area (B), medial area (C), intima/media ratio (D), and luminal areas (E) in micrometers squared along the length of vessels in WT (n=8), LPA1/−/− (n=6), LPA2/−/− (n=6), and LPA1/−/−2/−/− (n=6) mice. *$P<0.05$ in comparison to WT by ANOVA with multiple pairwise comparisons.

Figure 1. Increases in ATX/lysophospholipase D and LPA receptor expression following arterial injury. Carotid arteries were isolated from control, uninjured mice, or animals that underwent carotid ligation. A, ATX expression was determined by immunoblotting and was significantly higher ($P<0.05$ by ANOVA) at 1 and 3 days after injury in comparison to uninjured control levels. B, LPA receptor expression was determined by quantitative PCR. Results are expressed relative to uninjured vessel and presented as means±SD from 3 separate experiments.
LPA receptor–deficient mice using this femoral artery model because of a high incidence of acute arterial thrombosis in WT Balb/C.

Vascular injury elicits a characteristic sequence of events that includes inflammatory cell recruitment and SMC dedifferentiation, proliferation, and migration. To investigate the mechanism(s) by which LPA receptors might regulate vascular injury responses, we measured markers for these events after ligation. Following injury, there was an increase in ERK activation, as measured by the ratio of phospho-ERK to total ERK, and the increase in ERK activity returned to baseline by 5 days in LPA1−/− mice and by 7 days in LPA1−/− mice (Figure 3A). However, expression of the proliferation marker PCNA was upregulated to similar extents in WT, LPA1−/−, and LPA1−/− mice (Figure 3B), suggesting that cell proliferation following injury was similar in all genotypes. Likewise, lack of LPA1 and LPA2 did not affect SMC dedifferentiation following injury because downregulation of SMC markers occurred in all genotypes of mice examined (Figure 3B). Expression of interleukin-6 and the neutrophil- and monocyte-derived inflammatory marker myeloperoxidase increased following injury and was also unaffected by the absence of LPA1 and LPA2 (Figure 3C). As detected by immunoblotting, vessel-associated CD68, myeloperoxidase, and P-selectin, a marker of platelet activation, increased following injury in all genotypes (see supplemental Figure II). Taken together, these results indicate that vascular inflammation, SMC dedifferentiation, and cell proliferation are similar in WT, LPA1−/−, and LPA1−/− arteries following injury. Because these markers of proliferation and differentiation were unaltered by combined LPA1 and LPA2 deficiency, we considered the possibility that the attenuation of injury-induced intimal hyperplasia in LPA1−/− mice involves a requirement for these receptors as regulators of SMC migration that, as discussed below, we evaluated using primary cultures of mouse SMCs.

**LPA1−/−2−/− Vascular SMCs Exhibit Attenuated Proliferative Responsiveness In Vivo and In Vitro**

Although LPA responses of cultured SMCs from humans and rats have been investigated in detail, these types of studies have not been reported with mouse SMCs. To identify the mechanistic basis for vascular injury response phenotypes observed in the LPA receptor–deficient mice, we examined proliferation, dedifferentiation, and migration responses to LPA of isolated mouse vascular SMCs. No significant differences in growth of WT and LPA1−/− SMCs in serum were observed, but LPA1−/− SMCs grew more slowly (Figure 4A). This difference appeared to be dependent on the LPA component of serum because there was no difference between platelet-derived growth factor (PDGF)-induced growth of WT and LPA1−/− SMCs (Figure 4B). LPA regulates cell proliferation through ERK activation in many cell systems. Therefore, we examined LPA-induced ERK responses in isolated SMCs. LPA treatment of isolated SMCs increased ERK activation maximally at 5 to 10 minutes (Figure 4C). This response was concentration-dependent, with maximal activation being observed with 1 μmol/L LPA (data not shown). A similar increase in LPA-induced ERK activity in SMCs cultured from LPA2−/− mice was observed (data not shown).

**Figure 3.** LPA1 and LPA2 are not required for injury-induced alterations in proliferation, differentiation, or inflammation markers. Carotid injury increases ERK activity in vessels from WT, LPA1−/−, and LPA1−/− mice (A), although responses in the LPA1−/− vessels return to baseline within 3 days. pERK indicates phospho-ERK. B, Injury upregulates PCNA gene expression and downregulates SMC differentiation markers in WT, LPA1−/−, and LPA1−/− vessels. HC indicates heavy chain. C, Upregulation of myeloperoxidase (MPO) and interleukin-6 (IL-6) occurs following injury and is similar in WT, LPA1−/−, and LPA1−/− mice. Results are presented as mean ± SD from 3 separate experiments in vessels harvested at 5 days after injury.
shown). In contrast, there was an \( \approx \)2-fold reduction in the initial burst of LPA-induced ERK activation in LPA1\(^{-/-}\) SMCs and a 5-fold reduction in LPA1\(^{-/-}\)2\(^{-/-}\) SMCs (Figure 4C).

**LPA1\(^{-/-}\)2\(^{-/-}\) SMCs Exhibit Decreased Migration in Response to LPA, Whereas LPA1\(^{-/-}\) SMCs Exhibit Enhanced Migration in Response to Upregulation of the LPA3 Receptor**

Although the reduced proliferative responses observed in LPA1\(^{-/-}\)2\(^{-/-}\) SMCs in vitro are consistent with the protection of LPA1\(^{-/-}\)2\(^{-/-}\) mice from the formation of neointima in response to injury, the lack of a change in proliferation markers following injury in LPA1\(^{-/-}\)2\(^{-/-}\) vessels does not support the idea that protection from the development of intimal hyperplasia is a direct result of attenuated cell proliferation. In many systems, LPA is a potent stimulus for migration; therefore, we compared migration responses to LPA in SMCs derived from WT and LPA receptor–deficient mice (Figure 5). LPA, serum, and PDGF stimulated 4.7±0.94-, 6.44±0.046-, and 11.5±0.79-fold increases, respectively, in migration of SMCs (Figure 5). The migration of LPA1\(^{-/-}\)2\(^{-/-}\) cells was lower at baseline (39±1% that of WT cells; \( P < 0.01 \)), and the cells did not display any increase in migration to LPA and serum (Figure 5), although migration to PDGF was preserved (10.5±1.7-fold increase). In contrast, LPA1\(^{-/-}\) cells were hypermigratory at baseline (7.9±1.8-fold higher than WT cells; \( P = 0.003 \)) and displayed a further exaggerated migration response to LPA and serum (Figure 5).

To determine whether expression of an alternate LPA receptor might be responsible for the hypermigratory LPA response observed in the LPA1\(^{-/-}\) cells, the assays were conducted in the presence of VPC32301, a selective antagonist of the LPA3 receptor.\(^{33} \) LPA3 antagonism had a modest 30% inhibition of WT migration (22,978±3507 versus 33,680±6329 \( \mu \)m\(^2\) in antagonist-treated and vehicle-treated cells, respectively) and inhibited LPA-induced ERK activation in WT cells by 10% to 15%. LPA3 receptor antagonism reduced both baseline and LPA-induced migration in LPA1\(^{-/-}\)2\(^{-/-}\) cells to levels observed in WT cells (Figure 5) and inhibited LPA-induced ERK activation in LPA1\(^{-/-}\) cells by \( \approx \)35%. We found that LPA3 receptor expression was elevated 3.7±0.2-fold in LPA1\(^{-/-}\) but not LPA2\(^{-/-}\) or LPA1\(^{-/-}\)2\(^{-/-}\) cells as compared to WT cells (see supplemental Table I). To determine whether upregulation of LPA3 expression alone could account for the enhanced migration observed in LPA1\(^{-/-}\) cells, LPA3 was overexpressed in WT SMCs using recombinant lentivirus vectors, which resulted in enhanced migration toward LPA and serum phenocopying the behavior of LPA1\(^{-/-}\) cells (Figure 5).
LPA1 and LPA2 Are Not Required for Rho Activation or SMC Dedifferentiation

Migration of SMCs to LPA was attenuated by ERK and Rho kinase inhibitors (see supplemental Figure III). Therefore, we examined LPA-induced Rho responses in WT, LPA1+/−/−, and LPA1+/−/− cells toward 1 μmol/L LPA, 10% serum, or 20 ng/mL PDGF. Bottom right, Comparison of migratory responses of WT and LPA1−/− SMCs. Enhanced migration in LPA1−/− SMCs is attenuated by the LPA3 antagonist VPC32301 (10 μmol/L) and overexpression of LPA3 increases WT migration. Results from at least 3 separate experiments are reported as means±SD of surface coverage by migrated cells in micrometers squared. nd indicates not determined.

LPA1 and LPA2 Do Not Mediate Acute Blood Pressure Responses to LPA or Regulate Systemic Blood Pressure

The above results suggest that LPA regulates the vascular injury response by modulating migratory properties of SMCs. In other species, intravascular administration of LPA alters blood pressure, which likely results from direct effects on vascular smooth muscle cell contractility. We sought to determine whether the same receptors that regulate vascular SMC migration responses to LPA also contribute to the effects of LPA on blood pressure. Intravenous infusion of BSA-conjugated LPA transiently increased mean arterial blood pressure in anesthetized mice, with 10 pmol of LPA being the lowest dose that reproducibly increased mean arterial blood pressure above vehicle by 17±6 mm Hg in WT mice. A similar response was observed in mice lacking either LPA1 or LPA2 or both, suggesting that neither receptor is responsible for acute increase in blood pressure elicited by LPA (Table 1). Likewise, there was no difference in blood pressure or heart rate in conscious WT, LPA1−/−, LPA2−/−, or LPA1−/−/− mice (Table 2).

Discussion

In this study, we provide the first demonstration for a role for the LPA signaling nexus in endogenous regulation of pathophysiologic vascular responses. Specifically, we report that mice deficient in 2 defined LPA receptors, LPA1 and LPA2, are protected from intimal hyperplasia in response to vascular

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**Figure 5.** LPA receptor deficiency alters SMC migratory responses. SMCs were stained with Diff-Quik on the undersurface of a membrane with a 5-μm pore following migration to media containing vehicle in 0.5% FBS or 1 μmol/L LPA (top). Bottom left, Cell migration in WT, LPA2−/−, LPA1+/−/−, and LPA1+/−/− cells toward 1 μmol/L LPA, 10% serum, or 20 ng/mL PDGF. Bottom right, Comparison of migratory responses of WT and LPA1−/− SMCs. Enhanced migration in LPA1−/− SMCs is attenuated by the LPA3 antagonist VPC32301 (10 μmol/L) and overexpression of LPA3 increases WT migration. Results from at least 3 separate experiments are reported as means±SD of surface coverage by migrated cells in micrometers squared. nd indicates not determined.
injury through a mechanism that can primarily be accounted for by a decrease in the ability of LPA to promote migration of vascular SMCs. Interestingly, we also found that mice lacking LPA1 alone exhibited enhanced vascular injury responses, which correlated with an upregulation of a third LPA receptor, LPA3, and appeared to result from an enhanced LPA3-dependent vascular smooth muscle cell migratory response. These findings suggest that LPA is a relevant regulator of the murine vascular injury response and focus attention on possible sources of LPA at sites of vascular injury. We found that injury elevated vessel-associated levels of ATX, the lysophospholipase D responsible for generation of extracellular LPA. This may reflect localized expression of ATX or, perhaps more likely, recruitment of circulating ATX. Because ATX is responsible for generating circulating LPA and the ATX substrate lysocephatidylcholine is abundant in the circulation, the increase in vessel-associated ATX likely drives localized production of biologically active, extracellular LPA at the site of injury, which in turn promotes SMC migration via LPA1 and LPA2.

Comparison of the in vivo injury and in vitro SMC responses from mice that are singly or doubly deficient in LPA1 and LPA2 identifies functional redundancy and crosstalk between LPA receptors that has been reported in other cell types including cardiomyocytes and embryonic fibroblasts. For example, LPA-induced ERK activity is normal in LPA2−/− SMCs, slightly reduced in LPA1−/− cells, and nearly completely attenuated in LPA1−/−2−/− SMCs, indicating that both LPA1 and LPA2 are required to produce maximal LPA activation of ERK, which likely accounts for the reduced proliferative responses in LPA1−/−2−/− cells in vitro. LPA1−/−2−/− SMCs display an attenuated migration response to LPA, which may also related to the reduced ERK activity in response to LPA. These results also underscore the importance of the LPA component to the promigratory effects of serum on these cells, because LPA1−/−2−/− SMCs display reduced migration to both serum and LPA but not to PDGF.

Our results also point to LPA-dependent vascular signaling pathways that are not regulated by LPA1 or LPA2. Specifically, SMCs lacking both receptors appear to undergo essentially normal dedifferentiation in response to serum, LPA, or vascular injury. Because of the possibility that localized production of LPA could promote vasoconstriction at sites of vascular injury, we were particularly interested in evaluating the possible roles of LPA1 and LPA2 in blood pressure regulation. However, although

Figure 6. Neither LPA1 nor LPA2 is required for isolated SMC dedifferentiation. A, Expression of SMC-specific myosin heavy chain in cultured SMCs after serum deprivation or following reexposure to either 10% FCS or 1 μmol/L LPA for 24 hours. B, Rho activity was examined 10 minutes after exposure of cells to LPA (1 μmol/L) by Rhotekin pull-down assay. Values represented results obtained from 3 separate experiments and are reported as a fold change from baseline in WT cells (means±SD). S1P indicates sphingosine-1-phosphate.

Table 1. Blood Pressure and Heart Rate in WT Mice and Mice Lacking LPA1 and/or LPA2

<table>
<thead>
<tr>
<th>Genotype</th>
<th>N</th>
<th>SBP (mm Hg)</th>
<th>Heart Rate (bpm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>WT</td>
<td>6</td>
<td>124±26</td>
<td>621±38</td>
</tr>
<tr>
<td>LPA1−/−</td>
<td>9</td>
<td>126±12</td>
<td>621±46</td>
</tr>
<tr>
<td>LPA2−/−</td>
<td>12</td>
<td>133±11</td>
<td>623±46</td>
</tr>
<tr>
<td>LPA1−/−2−/−</td>
<td>4</td>
<td>120±24</td>
<td>588±66</td>
</tr>
</tbody>
</table>

Values for systolic blood pressure (SBP) and heart rate are presented as means±SD. There was not a statistically significant difference in blood pressure (P=0.500) or heart rate (P=0.621).

Table 2. Increase in Mean Arterial Pressure in Response to Intravenous Administration of LPA

<table>
<thead>
<tr>
<th>Genotype</th>
<th>N</th>
<th>Change in MAP (mm Hg)</th>
<th>Weight (g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>WT</td>
<td>5</td>
<td>17±6</td>
<td>29±4</td>
</tr>
<tr>
<td>LPA1−/−</td>
<td>3</td>
<td>25±8</td>
<td>22±4</td>
</tr>
<tr>
<td>LPA2−/−</td>
<td>3</td>
<td>19±8</td>
<td>23±1</td>
</tr>
<tr>
<td>LPA1−/−2−/−</td>
<td>2</td>
<td>20±2</td>
<td>29±4</td>
</tr>
</tbody>
</table>

Values for change in mean arterial pressure (MAP) and for weight are presented as means±SD. No statistically significant difference between groups was observed (P=0.466).
these receptors are clearly expressed and functional in vascular SMCs, we found that both systemic blood pressure and the acute, transient increase in arterial blood pressure provoked by direct administration of exogenous of LPA were unaltered in LPA1−/−, LPA2−/−, and LPA1−/−LPA2−/− mice. LPA exerts a range of effects in the cardiovascular system that include modulation of platelet activation, recruitment and activation of inflammatory cells, and regulation of blood pressure that likely results from the ability of LPA to promote SMC contractility. Clearly, much more work will be required to associate these signaling functions of LPA with the identified LPA receptors. One particularly interesting possibility is that the nuclear receptor PPARγ may mediate some G protein–coupled receptor–independent signaling actions of LPA, and we note that exogenously administered LPA may elicit neointimal formation in rodent vessels in a PPARγ-dependent manner.

Emerging evidence supports a role for LPA in regulation of vascular development and function. Mice lacking the LPA synthetic enzyme ATX/lysophospholipase D (Enpp2−/−) die embryonically with defects in vascular development, and homozygous inactivation of the LPP3 gene, which encodes a phosphatase enzyme responsible for inactivation of LPA, also results in early embryonic lethality resulting from defects in vascular development and patterning. Our data add to the weight of the evidence that the LPA signaling axis may be important in the regulation of pathophysiologically important vascular cells responses in adult animals. Taken together, our results suggest that interference with the enzymes and receptors responsible for LPA production and signaling would be a viable strategy for pharmacological intervention in the process of intimal hyperplasia.

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Disclosures

None.

References


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Supplemental Methods

Expression of LPA3
A cDNA clone corresponding to human LPA3 was obtained from the IMAGE consortium clone collection and the ORF amplified by PCR using a forward primer that incorporated an N-terminal HA-tag sequence CACCATGTACCCTACAGATGTCCAGATTACGCTaatgagtgtcactatgacaagcacatg). The PCR product was subcloned into pENTR D/TOPO (Invitrogen) and then used to generate a lentivirus expression vector by in vitro recombination with pLenti6.2 nV5 DEST (Invitrogen). Recombinant lentiviruses were obtained by transfection of HEK293FT cells with this vector and appropriate helper plasmids.

RNA Isolation and Quantitative PCR
Total RNA was extracted from carotid arteries and primary cultured mouse aortic smooth muscle cells using the RNeasy mini kit (Qiagen, Chatsworth, CA) following manufacturer’s instructions. cDNA was prepared with Multiscribe reverse-trasncriptase enzyme (High-Capacity cDNA Archive Kit; Applied Biosystems, Foster City, CA), and mRNA expression was measured in a RT-PCR reaction using TaqMan® gene expression assays and TaqMan® Universal PCR Master Mix No Amp Erase® (Applied Biosystems) in an ABI 7500 Fast Real-Time PCR System (Applied Biosystems). Threshold cycles (C\textsubscript{T}) were determined by an in-program algorithm assigning a fluorescence baseline based on readings prior to exponential amplification. Fold change in expression was calculated using the 2\textsuperscript{ΔΔC\textsubscript{T}} method using 18s RNA as an endogenous control\textsuperscript{1}.

SMC Proliferative Responses
SMC were serum deprived for 72 hours by growth in media containing 0.5% fetal bovine serum and then exposed to indicated concentrations of oleoyl (18:1) LPA dispersed in fatty acid free BSA or vehicle or serum (10%) or PDGF (20 ng/ml) for the indicated times. For growth curves, the cells were plated in 6-wells dishes (2 x 10\textsuperscript{4} cells/well), harvested with EDTA at the indicated time point, and viable cells counted. For ERK activity, cells were washed with ice cold PBS and lysed (10 mM Tris-HCl pH 7.2, 1% Nonidet P-40, 158 mM NaCl, 1 mM EDTA, 50 mM NaF, 1 mM PMSF, 10 μg/ml aprotinin, 10 μg/ml leupeptin, 1 mM sodium orthovanadate, 10 mM sodium pyrophosphate). Total protein in clarified cell lysates (16,000g x 10 min) was determined (BCA protein assay; Pierce) and equal amounts loaded onto 10% SDS-PAGE. Immunoblotting was performed with antibodies to actin (Santa Cruz Biotechnology) and ERK/phospho-ERK Thr202/Tyr204 and quantified using the Odyssey infrared imaging system (LI-COR). Equal sample loading was confirmed by Coomassie blue staining. Results are presented as the ratio of phosphoERK (p-ERK) to total ERK.

SMC Migration
After serum starvation in DMEM with 0.1% FBS, SMCs (1.8 x 10\textsuperscript{4} cells/well) were placed into the upper well of a chamber of a multiwell chambers (Neuroprobe Inc., Gaithersburg, MD) with apolyvinylpyrrolidone-free polycarbonate filter (5 μm pore). The bottom chamber was filled with DMEM with 0.1% FBS and the indicated chemotactic
agent (1 μM LPA, 10% FBS, 20 ng/ml PDGF) or vehicle. Where indicated, the ERK pathway inhibitor U0126 (Promega) or the Rho kinase inhibitor Y-27632 (Calbiochem) was included in the incubation. The chamber was incubated for 12-h at 37°C in a CO₂ incubator, at which time the filter was removed from the chamber and non-migrated cells were scraped from the upper surface. The membranes were fixed, and the migrated cells stained with Diff-Quik® (VWR Scientific Products). Digital images of the membranes were obtained with a Nikon 80i microscope using a 20x objective (NA = 0.5). The total area (µm²) occupied by migrated cells was determined with Metamorph imaging software.

Rho Activity
Rho activity was measured by incubating SMC lysates with GST-rhotekin Rho binding domain fusion protein immobilized to glutathione-beads. Proteins were separated on a 15% SDS-PAGE and immunoblotted with an anti-Rho antibody (Upstate). Results are expressed as a ratio of active to total Rho in the lysate.
Online Figure I. Femoral injury was performed in mice as previously described\textsuperscript{2, 3}. Four weeks after injury, vessels were isolated, serially sectioned, and stained with CME. The intimal area was 2.29 ± 0.48 fold greater in LPA1\textsuperscript{-/-} mice (n = 6) as compared to wild-type (WT) animals (n = 7).
Online Figure II. Changes in inflammatory markers post-injury. CD68 and P-selectin expression were measured by immunoblotting at the indicated times post-injury (A). At five days after injury, expression of CD68 and P-selectin was measured in WT, LPA1-/-, and LPA1-/-2-/- carotids (B). Myeloperoxidase (MPO) activity in the vessel was measured at the indicated times as previously described (C). nd = not determined.
Online Figure III. SMC migration to LPA is attenuated by ERK and Rho kinase inhibitors. SMC migration was performed as described in Online Supplemental Methods in the presence of the ERK inhibitor PD98059 (10 μM) or the Rho kinase inhibitor Y-27632 (10 μM).
### Online Table I. LPA receptor expression in LPA1<sup>−/−</sup>, LPA2<sup>−/−</sup>, and LPA1<sup>−/−</sup>-2<sup>−/−</sup> SMC relative to WT SMC

<table>
<thead>
<tr>
<th>Receptor</th>
<th>LPA1&lt;sup&gt;−/−&lt;/sup&gt;</th>
<th>LPA2&lt;sup&gt;−/−&lt;/sup&gt;</th>
<th>LPA1&lt;sup&gt;−/−&lt;/sup&gt;-2&lt;sup&gt;−/−&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>LPA1</td>
<td>ND</td>
<td>0.7 ± 0.1</td>
<td>ND</td>
</tr>
<tr>
<td>LPA2</td>
<td>1.5 ± 0.1</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>LPA3</td>
<td>3.7 ± 0.2</td>
<td>1.2 ± 0.1</td>
<td>0.99 ± 0.1</td>
</tr>
<tr>
<td>LPA4</td>
<td>0.8 ± 0.05</td>
<td>1.3 ± 0.1</td>
<td>1.3 ± 0.05</td>
</tr>
</tbody>
</table>

Results are presented as fold change relative to WT. ND = not detected. LPA5 expression was too low to be quantitated.

