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Deficiency or inhibition of lysophosphatidic acid receptor 1 protects against hyperoxia-induced lung injury in neonatal rats

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Abstract

Aim—Blocking of lysophosphatidic acid (LPA) receptor (LPAR) 1 may be a novel therapeutic option for bronchopulmonary dysplasia (BPD) by preventing the LPAR1-mediated adverse effects of its ligand (LPA), consisting of lung inflammation, pulmonary arterial hypertension (PAH) and fibrosis.

Methods—In Wistar rats with experimental BPD, induced by continuous exposure to 100% oxygen for 10 days, we determined the beneficial effects of LPAR1 deficiency in neonatal rats with a missense mutation in cytoplasmic helix 8 of LPAR1 and of LPAR1 and -3 blocking with Ki16425. Parameters investigated included survival, lung and heart histopathology, fibrin and collagen deposition, vascular leakage, and differential mRNA expression in the lungs of key genes involved in LPA signalling and BPD pathogenesis.

Results—LPAR1 mutant rats were protected against experimental BPD and mortality with reduced alveolar septal thickness, lung inflammation (reduced influx of macrophages and neutrophils, and CINC1 expression), and collagen III deposition. However, LPAR1 mutant rats were not protected against alveolar enlargement, increased medial wall thickness of small arterioles, fibrin deposition, and vascular alveolar leakage. Treatment of experimental BPD with Ki16425 confirmed the data observed in LPAR1 mutant rats, but did not reduce the pulmonary

Conflict of interest

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Authorship Contributions

F.W., E.C., G.F. and G.W. participated in research design. X.C., E.L., R.S., R.B. and G.W. conducted experiments. X.C., E.L., R.S., R.B., M.D. and G.W. performed data analysis. X.C., F.W., R.B., E.C., G.F. M.D. and G.W. wrote or contributed to the writing of the manuscript.

The authors report no conflicts of interest to disclose.

influx of neutrophils, CINC1 expression, and mortality in rats with experimental BPD. In addition, Ki16425 treatment protected against PAH and right ventricular hypertrophy.

Conclusion—LPAR1 deficiency attenuates pulmonary injury by reducing pulmonary inflammation and fibrosis, thereby reducing mortality, but does not affect alveolar and vascular development and, unlike Ki16425 treatment, does not prevent PAH in neonatal rats with experimental BPD.

Keywords

bronchopulmonary dysplasia; fibrosis; lung inflammation; lysophosphatidic acid receptor; right ventricular hypertrophy

Introduction

Premature infants at risk for neonatal chronic lung disease or bronchopulmonary dysplasia (BPD) suffer from lung damage due to lung immaturity and subsequent treatment for respiratory distress with mechanical ventilation and/or reactive oxygen species generated by (prolonged) exposure to supplemental oxygen. BPD leads to permanently enlarged alveoli, caused by an arrest in alveolar and vascular development, and a subsequent reduction of the alveolar surface and lung function (Baraldi & Filippone 2007, Jobe 1999). Serious complicating factors in the perinatal period are inflammation and oxidative stress, and at later stages pulmonary arterial hypertension (PAH)-induced right ventricular hypertrophy (RVH) and fibrosis (Abman 2009, Baraldi & Filippone 2007, Steinhorn 2010, Tuder *et al.* 2009). Similar to premature infants suffering from BPD, neonatal rats that are exposed to hyperoxia show chronic lung inflammation, persistent alveolar simplification, fibrosis, PAH and RVH (de Visser *et al.* 2010, 2012).

Lysophosphatidic acid (LPA) is a small glycerophospholipid, generated by the enzymatic removal of the choline group by extracellular autotaxin (lysophospholipase D) or phospholipase A1 or A2. LPA has a widespread tissue distribution and is secreted in the circulation (Rancoule et al. 2011). LPA exerts its multiple biological effects on cell proliferation, migration, survival, differentiation, motility, cytoskeletal change, inflammation and cell-cell interaction after binding to G protein coupled receptors (LPAR1-6) via multiple signaling pathways (Choi et al. 2010). LPA is involved in many pathological processes, including neurological disorders, cardiovascular disease, inflammation, lung and renal fibrosis, and cancer (Choi et al. 2010, Lin et al. 2010, Pyne et al. 2013). Experimental and clinical evidence strongly suggests that LPA is involved in lung pathology and disease, including airway repair and remodeling, inflammation, and fibrosis via LPA receptor signaling (Oikonomou et al. 2012, Shea & Tager 2012, Tager et al. 2008, Toews et al. 2002, Zhao & Natarajan 2013). LPAR1 or endothelial differentiation gene (EDG) family member 2 (EDG2), is the first identified LPA receptor with high affinity to LPA. In humans and rodents, LPAR1 is expressed in most organs and tissues, including the gastro-intestinal tract, heart, lung, brain, kidney, male and female reproductive organs, spleen, and thymus (Choi et al. 2010). LPAR1 couples with three types of G proteins: Ga12/13, Gaq/11, Gai/o, resulting in the initiation of several downstream signaling cascades: Rho-ROCK pathway, phospholipase C pathway, Ras- MAPK pathway, Akt

pathway, and adenylyl cyclase inhibition (Lin *et al.* 2010). Beneficial effects of reduced LPAR1 signaling in pulmonary disease have been demonstrated in fibroblasts of patients with idiopathic pulmonary fibrosis (Tager *et al.* 2008) and in rodents, as LPAR1 deficient mice are protected against fibrosis and mortality after bleomycin-induced pulmonary fibrosis (Tager *et al.* 2008) and LPS-induced lung inflammation (Zhao *et al.* 2011), and pharmacological blocking of LPAR1 protects against lung fibrosis (Swaney *et al.* 2010) and LPS-induced lung injury (Zhao *et al.* 2011). LPA is also a potent bronchoconstrictor, suggesting a role in airway hyperresponsiveness and asthma (Toews *et al.* 2002).

The role of LPAR1 deficiency in BPD is unknown, but may result in the identification of a novel therapy with specific LPAR1 antagonists for chronic lung disease, including BPD. To advance our knowledge on LPAR1 signaling in neonatal cardiopulmonary disease *in vivo*, we studied the beneficial effects of LPAR1 blocking in neonatal rats with experimental BPD induced by prolonged exposure to hyperoxia using two different models: (I) LPAR1 mutant rats (van Boxtel *et al.* 2011) and (II) pharmacological treatment with the LPAR1 and -3 inhibitor Ki16425 (Rancoule *et al.* 2011). We then investigated inflammation, alveolarization, fibrosis, PAH and RVH in the lung as described previously (de Visser *et al.* 2010).

Materials and Methods

Animals

The research protocol was approved by the Institutional Animal Care and Use Committee of the Leiden University Medical Center and the Royal Dutch Academy of Sciences. LPAR1 mutant rats carry a missense mutation with a methionine instead of arginine at position 318 in the cytoplasmatic 8th helix, resulting in a loss of function phenotype. This mutation was induced by an N-ethyl-N-nitrosourea (ENU)-driven target-selected mutagenesis approach in Wistar rats (van Boxtel et al. 2010, 2011). LPAR1 mutant rats were backcrossed for 7 generations on a Wistar background. For each experiment newborn wild type Wistar rat pups from 2 litters and homozygous LPAR1 mutant (LPAR1M318R/M318R) newborn rat pups from 2 litters were pooled. Hereafter, wild type and LPAR1^{M318R/M318R} mutant neonatal rat pups were equally distributed over the experimental groups: an oxygen group (N=12) and two room air (RA)-exposed control groups (N=6). All pups were fed by wild type Wistar foster dams. For the intervention experiments neonatal rat pups were pooled and distributed over four experimental groups (N=6): an oxygen-DMSO, oxygen-Ki16425, RA-DMSO and RA-Ki16425 group injected either with DMSO or Ki16425. The oxygen concentration, body weight, evidence of disease and mortality were monitored daily. Adult Wistar rats (6 months old; N=5) were exsanguinated after induction of anesthesia with an intraperitoneal injection of ketamine (50 mg/kg) and xylazine (50 mg/kg). Lungs were stored at -80°C until isolation of RNA for real time RT-PCR and hearts were fixed in formalin. On neonatal days 7 and 15 the thorax was opened under anaesthesia to visualize the ductus arteriosus in wild type and LPAR1^{M318R/M318R} mutant neonatal rat pups in situ (N=2). Hereafter, hearts were removed, fixed in formalin and processed for histology.

Hyperoxia-induced neonatal lung injury—Pups (a mix of both sexes) were continuously exposed to 100% oxygen for 10 days. In the intervention experiments pups received from day 2 either 5 mg kg⁻¹ day⁻¹ Ki16425 (Selleckchem S1315, Bioconnect, Huisen, The Netherlands) or DMSO (10%) in 0.9% NaCl (100 μ l, subcutaneously). Lung and heart tissue were collected on day 10 under anesthesia. Separate experiments were performed for [1] histology, [2] frozen lung tissue and [3] collection of bronchoalveolar lavage fluid. Lungs were snap-frozen in liquid nitrogen for *real-time* RT-PCR and fibrin deposition assay or fixed in formalin for histology studies as previously described (de Visser *et al.* 2009).

Primary rat embryonic fibroblast isolation

Embryonic fibroblasts, known to express LPAR1 (Stortelers *et al.* 2008), were used to functionally characterize the molecular consequences of the LPAR1 mutation. Heterozygous LPAR1 mutant rats were mated and at day 13.5 embryos were isolated. After washing the embryo thoroughly, the head and visceral organs were removed and used for DNA isolation and genotyping by PCR and sequencing using the following primer set: forward primer: 5'-AGATCCTCTCAGGCTGGTC-3' and reverse primer (5'-

ACAGGAGTGTCTTCGTCTCC-3') as described previously (van Boxtel et al. 2010). The embryos were minced and treated with trypsin to get a single cell suspension. Adherent rat embryonic fibroblasts (REFs) were grown in DMEM supplemented with 10% FCS.

In vitro fusion protein expression studies

Wild type and mutant receptors were N-terminally hemagglutinin (HA)-tagged by cloning into the expression vector pcDNA3.1 (Invitrogen, Life Technologies, Carlsbad, CA, USA). The receptor fusion protein constructs were transfected with a similar efficiency for both HA-tagged wild type and mutant LPAR1 plasmids and expressed in COS-7 cells (Figure S1). COS-7 cells were seeded on a coverslip and 24 hours after transfection the cells were placed on ice and incubated with DMEM-buffered HEPES containing 0.2% fatty acid-free bovine serum albumin (DHB) for 15 minutes. Subsequently, the cells were incubated for 1 hour with a polyclonal rabbit anti-HA (ab137838, Abcam Inc, Cambrigde, UK) on ice in DHB at a 1:500 dilution. For cell surface expression analysis the cells were immediately methanol-fixed and washed thoroughly with PBS and incubated for one hour in blocking buffer (1% BSA in 0.1% PBS-Tween) at room temperature. The cells were washed three times with PBS and incubated for 1 hour with a secondary anti-rabbit antibody conjugated with FITC (ab97050, Abcam Inc, Cambridge, UK), diluted 1:500 at room temperature in the dark. After three times washing with PBS the coverslips were mounted using Vectashield with DAPI (Brunschwig chemie, Amsterdam, The Netherlands) and analyzed using confocal microscopy. For western blotting COS-7 cells were lysed 24 hours after transfection, the proteins were separated on a SDS gel (10% acrylamide gradient, Bio-Rad, Veenendaal, The Netherlands) and transferred to a nitrocellulose membrane. The membrane was incubated for 1 hour at room temperature with a 1:500 or 1:1,000 dilution of respectively a polyclonal rabbit anti-HA antibody (ab13834, Abcam Inc, Cambridge, UK) or a polyclonal rabbit antiactin antibody (A2066, Sigma Aldrich, St. Louis, MO, USA) in blocking buffer followed by an incubation for 1 hours with peroxidase-conjugated, anti-rabbit IgG diluted 1:5,000 in blocking buffer at room temperature. Protein bands were detected by using the enhanced

chemiluminescence detection method (ECL, Amersham Biosciences/GE Healthcare, Eindhoven, The Netherlands).

Cell surface ELISA

Surface expression of N-terminally HA-tagged receptors was quantified as described (Gehret et al. 2010). Briefly, the N-terminally tagged receptors were transiently expressed in COS-7 cells and 24 hours after transfection the cells were harvested, seeded in 12 well plates and serum-starved overnight. To specifically measure the pool of cell surface expressed receptors, the cells were placed on ice and incubated with DMEM-buffered HEPES containing 0.2% fatty acid-free bovine serum albumin (DHB) for 15 minutes. Subsequently, the cells were incubated for 1 hour with a polyclonal rabbit anti-HA (ab137838, Abcam Inc, Cambridge, UK) on ice in DHB at a 1:500 dilution, rinsed with PBS on ice, fixed with ice-cold methanol: aceton (1:1) and air dried. Cells were then washed 5 minutes with PBS, blocked with RIPA/milk buffer (150 mM NaCl, 50 mM Tris, 1 mM EDTA, 10 mM NaF, 100nM sodium orthovanadate, 1% Triton X-100, 0.1% SDS, 0.5% sodium deoxycholate, pH 8.0, and 5% nonfat dried milk). To measure the total pool of Nterminally tagged receptors, cells were first fixed with ice-cold methanol: aceton (1:1), air dried and then washed 5 minutes with PBS. Subsequently, the cells were blocked with RIPA/milk buffer and incubated with 1:500 anti-HA antibody (ab13834, Abcam Inc, Cambridge, UK). All cells were then washed 3 times 5 minutes with PBS and incubated with peroxidase-conjugated, anti-rabbit IgG diluted 1:5,000 in RIPA/milk buffer. The cells were washed 3 times 5 minutes with PBS and incubated with ELISA TMB reagent (Sigma-Aldrich, St. Louis, MO, USA). The reaction was terminated with stop reagent for TMB substrate (Sigma-Aldrich, St. Louis, MO, USA) and the absorbance was measured at 450 nm. Data was presented as the A450 of the pool of cell surface expressed receptors as a percentage of the A₄₅₀ of the total pool of HA-tagged receptors.

ERK1/2 phosphorylation analysis

Primary REFs were seeded in 12-wells plates and allowed to attach overnight. Subsequently, the cells were serum-starved overnight in DMEM supplemented with 0.2% fatty acid-free BSA, and with or without the presence of 100 ng ml⁻¹ pertussis toxin (PTX). Cells were treated with 1 mM LPA and lysed with RIPA buffer supplemented with complete protease inhibitor cocktail (Roche Diagnostics, Woerden, The Netherlands) and PhosSTOP phosphatase Inhibitor Cocktail (Roche Diagnostics, Woerden, The Netherlands) at the indicated time points. Western blot analysis was performed as described above with a mouse antibody against pT183 and pY185 in human ERK1 and ERK2 (ab50011, Abcam Inc, Cambrigde, UK; diluted 1:5,000) and a rabbit antibody against total rat ERK1 and ERK2 (91021, Cell Signaling Technology, Danvers, MA, USA; diluted 1: 1,000).

Histology

Formalin-fixed, paraffin-embedded, 4 µm-thick heart and lung sections were stained with hematoxylin and eosin or Hart's stained to visualize elastin (Simon *et al.* 2010). Lungs were immunostained additionally with anti-LPAR1 (AP01253PU-N or SP4371P, Acris antibodies, San Diego, CA, USA, diluted 1:1,000), anti-ED-1 (monocytes and macrophages;

1:5), anti-myeloperoxidase (MPO, RB-373-A1, Thermo Fisher Scientific, Fremont, CA, USA; diluted 1:1,500), anti-α smooth muscle actin (ASMA, A2547, Sigma-Aldrich, St. Louis, MO, USA; diluted 1:20,000), anti-collagen III (COL3A1, ab7778, Abcam, Cambridge, United Kingdom; diluted 1:3,000) or anti-von Willebrand factor (vWF, A0082, Dako Cytomation, Glostrup, Denmark; diluted 1:4,000) using standard methods (de Visser *et al.* 2009). Quantitative morphometry was performed by two independent researchers blinded to the treatment strategy as previously described (de Visser *et al.* 2009, Yi *et al.* 2004, Wagenaar *et al.* 2013).

Fibrin detection assay

Quantitative fibrin deposition in lung tissue homogenates was determined by Western blotting as described previously (de Visser et al. 2009, Wagenaar et al. 2013). Tissue samples were dissolved in reducing sample buffer (10 mM Tris pH 7.5, 2% SDS, 5% glycerol, 5% -mercaptoethanol, and 0.4 mg/ml of bromophenol blue), subjected to SDS-PAGE (7.5% gel; 5% stacking gel) and blotted onto PVDF membrane (Immobilon-FL, Millipore, Bedford, MA). The 56-kDa fibrin β -chains were detected with monoclonal 59D8 (Oklahoma Medical Research Foundation, Oklahoma City, OK USA; diluted 1:1000), infrared labeled goat-anti-mouse secondary antibody (IRDye 800CW; Licor Biosciences, Lincoln, NE, USA, diluted 1:5000), and quantified using an infrared detection system (Odyssey infrared imaging system, Licor Biosciences, Lincoln, NE, USA). Fibrin deposition was quantified using rat fibrin as a reference.

Bronchoalveolar lavages, protein assay and CINC1 ELISA

Lung lavages, protein and CINC1 ELISA (ADI-900-074, Enzo Life Sciences, Raamsdonksveer, The Netherlands) were performed as previously described (de Visser *et al.* 2009, Wagenaar *et al.* 2013).

Real-time RT-PCR

Total RNA isolation from lung tissue homogenates (RNA-Bee, Tel-Test Inc, Bio-Connect BV, Huissen, the Netherlands), first-strand cDNA synthesis (SuperScript Choice System (Life Technologies, Breda, the Netherlands)) and *real-time* quantitative PCR, using β -actin as a housekeeping gene reference, were performed on a LightCycler 480 (Roche, Almere, the Netherlands) of the Leiden Genome Technology Center (Leiden, The Netherlands) as described previously (Wagenaar *et al.* 2004, 2013). Primers are listed in Table 1.

Statistical analysis

Values are expressed as mean \pm SEM. Differences between groups were analyzed with ANOVA, followed by Tukey's multiple comparison test. Differences at *p* values <0.05 were considered statistically significant.

Results

Dose finding for Ki16425 in experimental BPD

To determine the optimal dosing of Ki16425, we performed a pilot experiment in which hyperoxia-exposed rat pups were treated with 1-5 mg kg⁻¹ day⁻¹ Ki16425 or DMSO (2-10% in 0.9% NaCl. Ki16425 has anti-inflammatory properties (Zhao *et al.* 2011) and preliminary data with the LPAR1 deficient animals showed a reduced influx of macrophages in mutant rat pups with experimental BPD compared to wild type controls. Therefore, we used the influx of macrophages to the lung as a read-out and demonstrated that 5 mg kg⁻¹ day⁻¹ of Ki16425 was the most optimal dose to reduce the influx of macrophages into the lung in hyperoxia-induced BPD (Figure 1).

Effects of LPAR1^{M318R} mutation on LPAR1 expression in the lung

In the neonatal rat lung on day 10 highest LPAR1 protein expression was observed in the bronchial epithelial layer (Figure 2A). In addition protein expression was observed in alveolar type 2 cells and in arterioles in the endothelium and vascular smooth muscle cells, and in inflammatory cells. A similar pattern of expression was observed after exposure to hyperoxia (Figure 2B) and in LPAR1 deficient rat pups under normoxia (Figure 2C) and hyperoxia (Figure 2D).

LPAR1^{M318R} results in a hypomorphic phenotype in rats

Using N-ethyl-N-nitrosourea (ENU)-driven target-selected mutagenesis, a mutation was identified in LPAR1 that resulted in the substitution of methionine 318 into an arginine in the putative helix 8 of the receptor (Figure 3A; van Boxtel *et al.* 2011). The hydrophobicity of the affected residue is highly conserved in the GPCR class A family. Furthermore, this position is analogous to the phenylalanine of the NPxxY(x)5,6F motif found in many GPCRs, including the prototypic Rhodopsin and β 2 adrenergic receptor (ADRB2; Figure 3A). The residue sticks into a hydrophobic pocket and contacts the tyrosine of the same motif, which is important for the folding of the 8th helix. Computational analyses suggest that the amino acid change in the LPAR1 mutant will disrupt this hydrophobic interaction and additionally, an arginine is too big to fit in this pocket and will most probably result in incorrect packing of helix 8 (van Boxtel *et al.* 2011). In line with this, LPAR1 mutant rats showed a craniofacial malformation with an increase of the distance between both eyes and a shortening of the eye to nose tip distance (Figure 3B; van Boxtel *et al.* 2011), similarly as observed in LPAR1 knockout mice (Contos *et al.* 2000).

To test the molecular effect of the mutation on LPAR1 function, we first tested if LPAR1^{M318R} could still be transported and expressed in the plasma membrane, the site of action of the receptor. Although membrane expression was still observed in cells overexpressing mutant LPAR1 (LPAR1^{M318R}), it was considerably less compared to wild-type receptor (Figure 3C). Interestingly, cell surface-expressed mutant LPAR1 displayed a punctate appearance (Figure 3C), which can be indicative for receptor clustering in clathrin-coated pits and aberrant internalization of mutant LPAR1. Western blot analysis of lysates of cells overexpressing LPAR1 revealed a slightly decreased expression level of mutant compared to wild-type receptor (Figure 3D), which could reflect receptor instability or

degradation. Therefore, we quantified cell membrane expression using a cell surface ELISA and indeed observed that the pool of cell surface expressed HA-LPAR1 $^{\rm M318R}$ as a percentage of the total expressed mutant receptor was significantly decreased compared to HA-LPAR1^{WT} (Figure 3E). One of the pathways activated by LPA is the mitogen-activated protein (MAP) kinase cascade via pertussis toxin (PTX)-sensitive G_i in a tyrosine kinasedependent manner (Moolenaar et al. 2004). This pathway is thought to be the main mediator of the stimulatory effect of LPA on cell proliferation. To test the effect of LPAR1^{M318R} on the LPA-induced MAP kinase-signaling cascade, we isolated primary rat embryonic fibroblasts (REFs) that endogenously express LPAR1 and tested activation of the MAP kinase-signaling cascade in these cells. For this, serum-starved homozygous mutant and wild-type REFs were stimulated with LPA and ERK1/2 phosphorylation was measured. LPA treatment resulted in a rapid and transient ERK1/2 phosphorylation response (Figure 3F). In contrast, homozygous mutant REFs show an attenuated LPA-induced ERK1/2 phosphorylation response, demonstrating that M318R in LPAR1 diminishes LPA signaling in primary REFs. These results demonstrate that the LPAR1^{M318R} mutation results in a hypomorphic phenotype in rats.

Effects of LPAR1^{M318R} mutation and Ki16425 treatment on bodyweight and survival

On day 10, body weight was comparable in Wistar control and LPAR1 mutant rats kept in room air (19-20 g; Figure 4A) and in 100% oxygen (12 g). Administration of Ki16425 or its solvent DMSO to Wistar control rats for 10 days had no adverse effects on mean body weight in room air controls and oxygen-exposed pups. Exposure to hyperoxia resulted in a 40% survival on day 10 in non-treated and 60% in DMSO-treated Wistar rats and was not affected by administration of Ki16425 (Figure 4B). In LPAR1 mutant rats with experimental BPD survival was twofold higher than in non-treated O₂-exposed Wistar control rats (80%, p < 0.05). Room air-exposed pups showed no morbidity or mortality during the experimental period of 10 days.

Effects of LPAR1^{M318R} mutation and Ki16425 treatment on lung airway development and inflammation

Ten days after birth, LPAR1 deficiency or administration of Ki16425 or DMSO did not have adverse effects in the lung (Figure 5) on the number of alveolar crests (Figure 6A), pulmonary vessel density (Figure 6B), arterial medial wall thickness (Figure 6C), alveolar septal thickness (Figure 6D), and influx of macrophages (Figure 6E) and neutrophilic granulocytes (Figure 6F). Oxygen exposure for 10 days resulted in lung edema, a heterogeneous distribution of enlarged air-spaces (Figure 5B) with a decreased number of alveolar crests (2.4-fold, p < 0.001; Figure 6A) surrounded by septa with increased thickness (1.9-fold, p < 0.001; Figures 5B and 6D), reduced pulmonary vessel density (2.1-fold, p < 0.001; Figures 5B and 6D). Exposure to hyperoxia also led to an inflammatory response, characterized by an overwhelming influx of macrophages (6.7-fold, p < 0.001 in LPAR1 deficient rats and 6.0-fold, p < 0.001 in Ki16425 treated rats, Figures 5J and 6E) and neutrophils (5.7-fold, p < 0.001 in LPAR1 deficient experiments and 4.5-fold, p < 0.001 in Ki16425 experiments, Figures 5N and 6F), compared to room air-exposed controls.

In LPAR1 deficient rat pups reduced alveolar septal thickness (1.4-fold, p < 0.001; Figures 5C and 6D) and influx of macrophages (2.2-fold, p < 0.01; Figures 5K and) and neutrophils (3.8-fold, p < 0.01; Figures 5O and 6F) was observed compared to oxygen exposed controls, but beneficial effects on hyperoxia-induced inhibition of alveolarization and angiogenesis, and increased arterial medial wall thickness were absent. Administration of Ki16425 reduced medial wall thickness 1.3-fold (p < 0.05; Figure 5H and 6C), alveolar septal thickness (1.4-fold, p < 0.01; Figures 5D and 6D) and the influx of macrophages (2.3-fold, p < 0.01; Figures 5L and 6E) compared to oxygen-exposed controls. Ki16425 had no beneficial effects on hyperoxia-induced inhibition of alveolarization and angiogenesis, nor on the influx of neutrophils.

Effects of LPAR1^{M318R} mutation and Ki16425 treatment on lung collagen deposition and elastin expression

In all rat pups kept under normoxia collagen III was only present at high levels in the perivasculature of large and small blood vessels (Figure 7A). Expression was low or absent in alveolar septa. In lungs of pups exposed to hyperoxia for 10 days, collagen III deposition increased (7.3-fold, *p* < 0.001; LPAR1 experiment and 7.8-fold, *p* < 0.001; Ki16425 experiment (Figure 7M)), and was present in the perivasculature of blood vessels and in thick alveolar septa (Figure 7B). LPAR1 deficiency or treatment of experimental BPD with Ki16425 for 10 days reduced collagen III expression by 55% (p < 0.001; Figure 7M) in the thin alveolar septa (Figures 7C and 7D). In all rat pups kept under normoxia, elastin was predominantly present on the septal tips in the alveoli and in the wall of blood vessels (Figure 7E, F, I and J). Under normoxia, elastin expression increased 1.4-fold (p < 0.05; Figure 7N) in LPAR1 deficient rats compared to Wistar controls, which can be explained by an increase in elastin expression in the small arterioles (Figure 7F). In wistar rats kept under hyperoxia, elastin expression decreased 1.4-fold compared to normoxia-exposed controls after DMSO administration (p < 0.01; Figure 7N), and was predominantly present in the alveolar walls rather than on septal tips and in the walls of small blood vessels (Figures 7G and 7K). LPAR1 deficient rats showed a 1.8-fold decrease in elastin expression after exposure to hyperoxia (p < 0.001; Figure 7N). LPAR1 deficiency or treatment with Ki16425 for 10 days had no effect on elastin expression in blood vessels and on the hyperoxiainduced irregular elastin expression in alveolar walls in pups with hyperoxia-induced lung injury (Figures 7H and 7L).

Effects of LPAR1^{M318R} mutation and Ki16425 treatment on coagulation, vascular leakage and CINC1 expression in BALF

Pulmonary fibrin deposition was studied in homogenates as a read-out for lung damage (Figure 8A). Fibrin deposition was at reference levels in all rat pups kept under normoxia for 10 days (< 20 ng fibrin mg⁻¹ tissue), and increased (29-fold, p < 0.01 in LPAR1 experiment and 5-fold, p < 0.01 in Ki16425 experiment) in lungs of pups exposed to 100% oxygen for 10 days. LPAR1 deficiency or Ki16425 administration did not reduce hyperoxia-induced fibrin deposition. Total protein concentration in bronchoalveolar lavage fluid (BALF) was determined to establish the effect of pulmonary edema by capillary-alveolar leakage (Figure 8B). After exposure to hyperoxia for 10 days the protein concentration increased (19-fold, p < 0.01; LPAR1 experiment and 11-fold, p < 0.05; Ki16425 experiment) and was not

affected significantly in LPAR1 deficient or Ki16425 treated rat pups with experimental BPD, although Ki16425-treated pups with experimental BPD showed a tendency towards lower levels. CINC1 expression in BALF increased after exposure to hyperoxia for 10 days (13-fold, p < 0.001; LPAR1 experiment and 5.0-fold, p < 0.01; Ki16425 experiment) and was not affected in rat pups kept under normoxia (Figure 8C). In LPAR1 deficient rat pups CINC1 expression decreased 2.6-fold (p < 0.05) compared to oxygen exposed controls. CINC1 expression was not affected significantly after Ki16425 treatment of experimental BPD.

mRNA expression in lung tissue

LPAR1-6 expression during neonatal lung development and experimental

BPD—Directly after birth on neonatal day 1 the mRNA of LPAR1-6 is expressed in the lung (Figure 9A-F). During early neonatal lung development expression of LPAR2 (B), and LPAR5 (E) and LPAR6 (F) increased, whereas LPAR3 (C) decreased. This resulted in a 2.0fold (P < 0.001), 1.7-fold (P < 0.05) and 1.4-fold (P < 0.01) increase in mRNA expression for LPAR2, LPAR5 and LPAR6, respectively, on day 10 compared with day 1. During normal lung development LPAR2 expression was increased 1.7-fold (P < 0.01) on day 3, and LPAR3 was decreased 1.7-fold (P < 0.01) on day 3 and 2.0-fold on day 6 (P < 0.001) compared with day 1. In adult lung mRNA expression increased 3.6-fold (P < 0.001) for LPAR3 compared with day 10 and decreased for LPAR1, LPAR4 (D) and LPAR6 compared with day 10: 2.6-fold (P < 0.001) for LPAR1, 5.2-fold (P < 0.001) for LPAR4 and 1.8-fold (P < 0.001) for LPAR6. Exposure to 100% oxygen for 10 days resulted in a decrease in mRNA expression for LPAR1 (1.3-fold, P < 0.001), LPAR2 (2.1-fold, P < 0.001), LPAR3 (1.5-fold, *P* < 0.01), LPAR4 (1.5-fold, *P* < 0.001) and LPAR6 (2.9-fold, *P* < 0.001). Exposure to 100% oxygen for 6 days resulted in an increase in mRNA expression for LPAR1 (1.2-fold, P < 0.01) and LPAR3 (1.7-fold, P < 0.001) and a decrease in expression for LPAR6 (2.9-fold, P < 0.001), whereas after 3 days of hyperoxia LPAR2 expression was decreased 1.4-fold (P < 0.05) on day 3.

Effects of LPAR1^{M318R} mutation and Ki16425 treatment on mRNA expression

in lung tissue—In all rat pups kept under normoxia low levels of mRNA expression (Figure 10) of the pro-inflammatory factor chemokine-induced neutrophilic chemoattractant-1 (CINC1; Figure 10A), the pro-coagulant factor tissue factor (TF; Figure 10B) and anti-fibrinolytic protein plasminogen activator inhibitor 1 (PAI-1; Figure 10C), and relatively high levels of mRNA expression of LPAR1 (Figure 10D), LPAR2 (Figure 10E) and LPAR3 (Figure 10F) were observed. Ten days of oxygen exposure resulted in an increase in mRNA expression of CINC-1 (8.3-fold, p < 0.001 in LPAR1 control and 11.8-fold, p < 0.001 in Ki16425 control), TF (3.0-fold, p < 0.001 in LPAR1 control and 2.8-fold, p < 0.001 in Ki16425 control) and PAI-1 (40.3-fold, p < 0.001 in LPAR1 control and 51.9-fold, p < 0.001 in LPAR1 control and p < 0.001 in Ki16425 control). The sequence in mRNA expression of LPAR1 (1.6-fold, p < 0.01 in LPAR1 control and p < 0.001 in Ki16425 control). The sequence in mRNA expression of LPAR1 (1.7-fold, p < 0.01). Exposure of LPAR1 deficient rat pups to hyperoxia resulted in an increase in mRNA expression of CINC1 (1.8-fold, p < 0.05) and a decrease in LPAR1 (1.7-fold, p < 0.01; Figure 10D) and LPAR3 (1.6-fold, p < 0.05; Figure 10F) compared to oxygen-exposed controls. Treatment of oxygen-

exposed pups with Ki16425 for 10 days resulted in an increase in mRNA expression of TF (1.3-fold, p < 0.05) compared to DMSO-treated oxygen-exposed pups.

Effects of LPAR1^{M318R} mutation and Ki16425 treatment on right ventricular hypertrophy

Because the increase in elastin staining in small arterioles in LPAR1 deficient rats may be associated with vascular remodeling and pulmonary arterial hypertension (PAH), and Ki16425 treatment of experimental BPD reduced arteriolar medial wall thickness we investigated right ventricular hypertrophy (RVH) on haematoxylin and eosin heart sections (Figures 11A and 11B). On neonatal day 10 LPAR1 deficient rat pups developed RVH under normoxia compared to room air Wistar controls, affected by an increase in the ratio RV/LV free wall thickness (1.4-fold, p < 0.01; Figure 11C), which could be explained by an increase in right ventricular free wall thickness (1.6-fold, p < 0.001; Figures 11A and 11D). In adult LPAR1 mutant rats the ratio RV/LV free wall thickness was similar to wild type controls, demonstrating that neonatal RVH in LPAR1 mutant rats is transient. Exposure to hyperoxia for 10 days resulted in an 1.4-fold increase in the ratio RV/LV free wall thickness in Wistar control pups compared to room air controls (p < 0.05), but did not aggravate RVH in LPAR1 deficient rat pups compared to LPAR1-deficient controls kept in room air. Administration of Ki16425 for 10 days during normal neonatal development had no adverse effect on the heart (Figure 11B). Exposure to hyperoxia for 10 days resulted in RVH (Figure 11B), affected by an 1.3-fold increase in the ratio RV/LV free wall thickness (Figure 11C) compared to room air controls (p < 0.05). Administration of Ki16425 prevented RVH, demonstrated by a normalization in relative RV/LV free wall thickness (p < 0.01) compared to oxygen-exposed pups. Hyperoxia and treatment with Ki16425 only effected the right ventricle (Figure 11D), but not the left ventricle (Figure 11E). Because transient RVH in LPAR1 deficient rats on day 10, but not in adult rats, can not be explained by pulmonary hypertension, we investigated whether RVH could be explained by a relatively late closure of the ductus arteriosus. From day 7 onward the ductus arteriosus was rudimentary in both wild type (Figure 12A) and LPAR1 mutant rat pups (Figure 12B). Closure of the ductus arteriosus was demonstrated in haematoxylin and eosin stained sections at the level of entrance of the ductus into the aorta in wild type (Figure 12C and D) and LPAR1 mutant rat pups on day 7 (Figure 12E and F). This suggests that neonatal RVH can not be explained by a relatively late closure of the ductus arteriosus in LPAR1 mutant rats.

Discussion

Mutant homozygous LPAR1^{M318R/M318R} rats have an N-ethyl-N-nitrosourea (ENU)induced missense mutation in helix 8 of LPAR1 that disrupts a highly conserved hydrophobic interaction between the transmembrane 7 domain and helix 8. This causes a reduced and punctate LPAR1 expression at the cell surface, impaired LPA-induced signaling, as shown in primary rat embryonic fibroblasts, an apparent loss-of-function phenotype, which was characterized by a craniofacial disorder (this study and van Boxtel *et al.* 2011), previously observed in LPAR1 knockout mice (Contos *et al.* 2000). In neonatal LPAR1^{M318R/M318R} mutant rat pups prolonged exposure to hyperoxia, an *in vivo* model for experimental BPD (Wagenaar *et al.*, 2004), survival was prolonged and pulmonary injury was reduced, demonstrated by a decrease in alveolar septal thickness, inflammation and

extravascular collagen deposition in the lung. LPAR1 deficiency had no beneficial effects on lung alveolarization, vascularization, arterial medial wall thickness (pulmonary arterial hypertension; PAH), capillary alveolar leakage, and fibrin deposition and no adverse effects on normal lung development. Similar beneficial effects on inflammation and extravascular collagen deposition were observed after Ki16425 treatment. In addition Ki16425 treatment of experimental BPD reduced PAH and prevented right ventricular hypertrophy (RVH), demonstrating that LPAR1 and/or -3 antagonists may be suitable candidates to reduce lung inflammation, fibrosis and PAH-induced RVH in preterm infants with severe BPD.

The receptors LPAR1-6 and autotaxin, the enzyme that is essential for the local production of the ligand LPA, are differentially expressed during lung development and/or in hyperoxia-induced lung injury, suggesting a role for LPA-LPA receptor signaling in the pathophysiology of severe experimental BPD, in which arrested alveolarization, inflammation and pulmonary hypertension play a pivotal role. However, the role of LPA-LPA receptor signaling in experimental BPD is still unclear. The relatively high expression of LPAR1 in the neonatal lung and the adaptive response in mRNA expression in the neonatal lung towards hyperoxia, may contribute to the beneficial pulmonary effects that we observed in LPAR1 deficient rats and after treatment with the LPAR1 and -3 antagonist Ki16425 of rat pups with experimental BPD in this study.

Similar to adult rodents exposure of neonatal rats to hyperoxia for 1 week increases ERK 1/2phosphorylation in lung tissue homogenates (Porzionato A, 2015, Lang Y, 2010 and Jiang J, 2011). When extrapolating our data on reduced ERK 1/2 phosphorylation in embryonic fibroblasts from LPAR1 mutant rats to neonatal lung, we expect reduced LPA-induced ERK 1/2 phosphorylation in neonatal experimental BPD lungs from LPAR1 mutant rats and from rats in which LPAR1 and -3 is pharmacologically blocked compared to oxygen-exposed Wistar control rats. This suggests that reduced ERK activation induced by LPAR1 deficiency or blocking may contribute to the beneficial effects we observed in experimental BPD. Because LPA-induced LPAR1 stimulation and exposure to hyperoxia activate many signal transduction pathways, including ERK and P38 MAPK (Choi 2010 and Porzionato A, 2015), which may be complicated by redundancy in LPA-LPAR-dependent signaling as demonstrated in the differential expression of LPAR1-6 during normal neonatal lung development and in experimental BPD (Figure 9), additional experiments are needed to establish the potential beneficial role of reduced LPAR1 and/or -3-induced ERKphosphorylation in experimental BPD. Differences in response towards hyperoxia-induced neonatal lung injury between the two experimental approaches (LPAR1 mutant rats, deficient in LPAR1 and pharmacological inhibition of LPAR1 and -3) are neutrophil influx and CINC1 expression, right ventricular hypertrophy, and survival. These differences in response towards experimental BPD may be explained by (1) administration of fluid, resulting in a less severe phenotype in the Ki16425 experiments, (2) potential adverse effects of DMSO (10%) or compound, (3) a relatively low compound concentration to avoid potential toxic effects of DMSO, (4) inhibition of both LPAR1 and -3 in the Ki16425 experiments and (5) the continuous absence of LPAR1 during embryonic, fetal and postnatal development in the LPAR1 mutant rats. The absence of a beneficial effect on survival and the influx of neutrophils in the Ki16425 experiments may be explained by the daily administration of fluid. This results in less severe experimental BPD with reduced

inflammation (influx of macrophages and neutrophils and CINC1 expression in BALF), vascular leakage, fibrin deposition and mortality in the DMSO treated control groups, thereby obscuring the potential beneficial effect of Ki16425 on neutrophil influx and survival in experimental BPD, but adverse effects of solvent (10% DMSO) or compound (Ki16425) can not be excluded. Furthermore, the additional blocking of LPAR3 by Ki16425 and the relatively low concentration of Ki16425 we used in this study with neonatal rats (5 mg kg⁻¹ day⁻¹) compared to adult rodents (up to 20 mg kg⁻¹; Zhao *et al.*, 2015) may also contribute to the differences in response between both experimental approaches. The absence of a beneficial effect on medial wall thickness and RVH in LPAR1 mutant rats may be explained by the development of RVH in LPAR1 mutant rats during neonatal development in room air.

Inflammation is an important contributor to the pathogenesis of BPD, because it may result in severe tissue damage and fibrosis, and treatment with anti-inflammatory agents provides protection against hyperoxia-induced neonatal lung disease or experimental BPD (de Visser et al. 2012, Yi et al. 2004). LPAR1 deficiency and inhibition protected against hyperoxiainduced neonatal lung injury in rats by reducing fibrosis, as demonstrated by reduced extravascular collagen III deposition, and inflammation, as demonstrated by a reduced pulmonary influx of macrophages and neutrophils in LPAR1 deficient rats, and macrophages after Ki16425 treatment. This resulted in reduced mortality among LPAR1 deficient rats, but not in Ki16425 treated rats. This anti-inflammatory effect was confirmed by the reduction of the neutrophilic attractant CINC1 in BALF in mutant rats, but was not accompanied by the observed expression of the mRNAs of the inflammatory marker CINC1, suggesting the involvement of other chemokines and cytokines, other mechanisms involved in inflammation, including cell adhesion and migration or regulation of gene expression at a post-transcriptional level. Our data are supported by observations in mice deficient in LPAR1 or autotaxin, and mice treated with an LPAR1 antagonist, which are protected against bleomycin-induced fibrosis and mortality (Oikonomou et al. 2012, Swaney et al. 2010, Tager et al. 2008), in allergic asthmatic mice with reduced autotaxin expression or enzymatic activity, which resulted in an attenuated inflammatory response (Park et al. 2013) and in Ki16425 treated mice which are protected against LPS-induced peritoneal sepsis (Zhao et al. 2015). Pro-inflammatory properties of LPA have been demonstrated in human lung epithelial cells by the secretion of IL8 and activation of NFkappa B, P38 MAPK and AP1 (Cummings et al. 2004, Saatian et al. 2006). Blocking of LPAR1 attenuates the LPSinduced proinflammatory response by reducing IL-6 production in vivo and in vitro in MLE12 mouse lung epithelial cells, probably via inhibition of LPS-induced p38 MAPK phosphorylation and NFkappa B activation through a direct interaction of CD14 and LPAR1 (Zhao et al. 2011).

LPAR1 deficiency in mice is associated with increased perinatal mortality and altered suckling behavior that may be related to a craniofacial deformation and may lead to a reduction in food intake, bodyweight and survival in the perinatal period (Contos 2000). Also LPAR1 mutant rats have a craniofacial deformation (this study and van Boxtel *et al.* 2011), but these rats grow normally, suggesting normal suckling behavior and milk intake, and are protected against neonatal mortality in hyperoxia compared to Wistar control pups,

which may be explained by species specific differences between mice and rats. We only observed increased survival of experimental BPD in LPAR1 deficient rat pups, but not after Ki16425 treatment. This discrepancy in survival between both experiments may be explained by dehydration of LPAR1 deficient rats and their controls or by the lack of selectivity of Ki16425 for LPAR1 and -3. Although we used the same Wistar rat strain as a control, only the pups in the Ki16425 experiments received daily 100 µl of fluid, thereby reducing potential dehydration and subsequent pathology, as demonstrated by a decreased inflammatory response and mortality in the hyperoxia-exposed control pups in the Ki16425 experiments. The higher survival rate in the oxygen-exposed controls of the Ki16425 experiments may obscure a potential beneficial effect on survival of Ki16425 treatment in experimental BPD. Alternatively, the discrepancy in survival between both experiments may be explained by differences in the efficacy to reduce the pulmonary influx of neutrophilic granulocytes to the lung, which was confirmed by CINC1 expression in BALF, or by a direct effect of aberrant LPAR1 signaling on embryonic development in LPAR1

LPAR1 deficiency and inhibition did not reduce vascular leakage. This finding is supported by our fibrin deposition data, in which extravascular fibrin deposition depends on leakage of the plasma protein fibrinogen into the alveolar lumen and its local conversion into fibrin by thrombin (Wagenaar *et al.* 2004). These data are in agreement with the absence of a beneficial effect of Ki16425 on LPS-induced protein accumulation in bronchoalveolar lavage fluid (Zhao *et al.* 2011), but in contrast with beneficial effects of LPAR1 deficiency or pharmacological inhibition of bleomycin-induced vascular leakage in experimental lung fibrosis (Swaney *et al.* 2010, Tager *et al.* 2008). These differences in the efficacy to reduce pulmonary vascular leakage may be explained, at least in part, by differences in animal models, species, receptor antagonist, and the onset and progression of tissue damage in these models of experimental lung disease.

A role for LPA in the cardiovasculature is demonstrated in atherosclerosis (Schober & Siess 2012) and the regulation of blood pressure in vivo, both as a vasopressor and a vasodilator depending on the species studied and experimental setup (Tokumura et al. 1978, 1995). In mice, LPA is a potent endothelial nitric oxide synthase (eNOS)-dependent vasodilator of the intact thoracic aorta (Ruisanchez et al. 2014). This LPA-induced vasodilation response of intact thoracic aortas is LPAR1-dependent, because it can be abolished by pharmacological inhibition of LPAR1 (Ruisanchez et al. 2014). Based on these experimental data, we expected an adverse effect of LPAR1 blocking or deficiency on pulmonary arterial hypertension (PAH). However, LPAR1 and -3 blocking reduced medial wall thickness of small arterioles, which was confirmed by a reduction in RVH, whereas LPAR1 deficiency had no beneficial effect on PAH, thereby confirming the data observed in LPAR1 knockout mice (Cheng et al. 2012). The discrepancy in response towards PAH-induced RVH in LPAR1 mutant and Ki16425 treated rat pups with BPD may be explained by: (1) a direct effect of aberrant LPAR1 signaling on embryonic lung and heart development in LPAR1 mutant rats only, resulting in neonatal RVH or (2) a beneficial effect of LPAR1 and -3 inhibition on PAH in Ki16425-treated rat pups. LPAR1 and -2 double knockout mice suffered from lung vascular remodeling and RVH (Cheng et al. 2012). Interestingly, we

observed an increase in elastin expression in small arterioles in LPAR1 deficient rats, which was not associated with an increase in our marker for PAH, i.e. increased medial wall thickness, but was associated with RVH. LPAR1 has a wide spread tissue distribution and is expressed in many cell types, including endothelial and (vascular) smooth muscle cells in blood vessels. We speculate that LPA-induced LPAR1 activation on the endothelium may lead to vasodilation, whereas LPAR1 activation on vascular smooth muscle cells may lead to vasoconstriction (Ruisanchez et al. 2014). In the cardiovasculature this behavior is also observed for other receptors, including the apelin receptor APJ, bradykinin B1 and B2 receptors, and muscarinic receptors, and may be complicated by receptor expression on inflammatory cells and cardiomyocytes (Felipe et al. 2007, Japp & Newby 2008, Obi et al. 1994). Because we observed transient RVH in LPAR1 deficient rats on day 10, but not in adult rats, which can not be explained by pulmonary hypertension, we investigated whether RVH could be explained by a delayed adaptation of the fetal circulation directly after birth by studying relatively late closure of the ductus arteriosus. The ductus arteriosus was closed in both wild type and LPAR1 mutant rat pups from day 7 onward. Therefore, it is unlikely that a delayed closure of the ductus arteriosus contributes to neonatal RVH and may be explained by other developmental differences in fetal and early neonatal heart development, including a persisting foramen ovale in LPAR1 mutant neonatal rats, for which additional research is needed. In addition, no right ventricular outflow obstructions or ventricular septal defects were observed in neonatal and adult hearts that could explain the transient RVH in LPAR1 mutant rats.

When the absence of adverse effects of treatment with LPAR1 and/or -3 blockers in rats is confirmed in patients, extrapolation of the beneficial effects of LPAR1 deficiency and treatment with blockers of LPAR1 or LPAR1 and -3 in rat pups with experimental BPD to preterm infants with respiratory failure, may result in a beneficial effect on lung inflammation, pulmonary arterial hypertension and fibrosis which are major reasons for mortality or morbidity in preterm infants with severe BPD.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Abbreviations

ASMA a smooth muscle actin

BALF	bronchoalveolar lavage fluid	
BPD	bronchopulmonary dysplasia	
CINC1	chemokine-induced neutrophilic chemoattractant-1	
CLD	chronic lung disease	
DMSO	dimethyl sulfoxide	
ENU	N-ethyl-N-nitrosourea	
HA	hemagglutinin	
IL	interleukin	
IVS	interventricular septum	
LPA	lysophosphatidic acid	
LPAR	LPA receptor	
LV	left ventricle	
MCP1	monocyte chemoattractant protein 1	
MPO	myeloperoxidase	
O ₂	oxygen	
PAH	Pulmonary arterial hypertension	
PAI-1	plasminogen activator inhibitor 1	
RA	room air	
RT-PCR	reverse transcriptase polymerase chain reaction	
RVH	right ventricular hypertrophy	
TF	tissue factor	
vWF	von Willebrand factor	

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Figure 1.

Pilot experiment to find the optimal dose of Ki16425 for treatment of experimental BPD by determining the pulmonary influx of ED-1-positive monocytes and macrophages in tissue sections in room air (RA), pups injected daily with DMSO (10%, open bar) and O₂-exposed pups (O₂) injected daily with 10% DMSO or Ki16425 (shaded bars): 1, 2.5 and 5 mg kg⁻¹ day⁻¹. Data are expressed as mean \pm SEM. ***p < 0.001 versus RA control. ^{††}p < 0.01 and ^{†††}p < 0.001 versus age-matched DMSO-treated O₂-exposed control; n=4-8 per group.



Figure 2.

Representative lung sections stained for lysophosphatidic acid receptor 1 (LPAR1 or endothelial differentiation gene 2 (EDG2); A-H) of wild type (A, B, E and F) and LPAR1 mutant rat pups (C, D, G and H) kept in room-air (RA; A, C, E and G) or 100% O_2 (B, D, F and H) until 10 days of age. Boxed areas in panels A, B, C and D are represented in panels E, F, G and H, respectively. a = alveolus; ar = arteriole; br = bronchus. Arrow in panel B indicates LPAR1-positive inflammatory cells.



Figure 3.

Nonsynonymous mutation in the LPAR1 gene results in a hypomorphic phenotype. (A) Alignment of amino acid sequence of the 8th helices of different class A GPCRs. Indicated in red is the NPxxY(x)_{5,6}F motif of which the tyrosine in transmembrane (TM) 7 domain and the phenylalanine in helix 8 form a hydrophobic interaction in the inactive state. In LPAR1-3 the phenylalanine is replaced by a methionine, conserving hydrophobicity. The arrowhead indicates the location of the amino acid change caused by the mutation. Indicated in blue are the basic amino acids that are commonly found in helix 8. (B) LPAR1 mutant rats have a craniofacial malformation with a shortening of the eye to nose tip distance and an increase in the distance between the eyes compared to wild type Wistar rats. (C) Decreased membrane expression of overexpressed LPAR1^{M318R} compared to overexpressed wild-type or mutant LPAR1 were incubated with an anti-HA antibody. The antibody can only bind if the cell expresses the receptor in the membrane, because only then the HA-tag is localized outside the cell. (D) Western blot analysis of COS-7 cells overexpressing either HA-LPAR1^{WT} or HA-LPAR1^{M318R}. (E) The size of the cell surface expressing receptor pool as

percentage of the total receptor pool is decreased in cells expressing LPAR1^{M318R}. Graph indicates the pool of cell surface expressed receptor as a percentage of total HA-tagged expressed receptors measured by cell surface ELISA. (F) LPAR1^{M318R/M318R} rat embryonic fibroblasts (REFs) display a hypomorphic ERK1/2 phosphorylation pattern. Western blot analysis of MAP kinase pathway activation in wild type and mutant REFs after 1 mM LPA treatment.

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Figure 4.

Growth (A) and survival (B) on day 10 in room air (RA) and age-matched O₂-exposed wild type or LPAR1 mutant rat pups (O₂). Wild type Wistar rats were injected daily with DMSO (10%) or Ki16425 (5 mg kg⁻¹ day⁻¹) for 10 days. Open bars: RA and RA-LPAR1-mutant or Ki16425-treated pups, shaded bars: O₂-control, O₂-LPAR1 mutant or O₂-Ki16425 treatment. Data are expressed as mean \pm SEM. *p < 0.05 and ***p < 0.001 versus own room air controls. [†]p < 0.05 versus age-matched O₂-exposed control. LPAR1^M = LPAR1^{M318R/M318R} mutant rat; Ki = Ki16425. - : daily administration of solvent (DMSO) in Ki16425 experiments or wild type rat in LPAR1 mutant rat experiments. +: daily treatment with 5 mg kg⁻¹ of Ki16425 or LPAR1 mutant rat in 4-7 independent experiments per group.



Figure 5.

Representative lung sections stained for von Willebrand Factor (vWF; A-D), α smooth muscle actin (ASMA; E-H), the monocyte and macrophage marker ED1 (I-L) or myeloperoxidase (MPO) as a marker for neutrophilic granulocytes (M-P) of wild type (A, B, D, E, F, H, I, J, L, M, N and P) and LPAR1 mutant rat pups (C, G, K and O) kept in roomair (RA; A, E, I, and M) or 100% O₂ (B-D, F-H, J-L and N-P) injected daily with 10% DMSO (A, B, E, F, I, J, M and N) or 5 mg kg⁻¹ day⁻¹ of Ki16425 (D, H, L and P) until 10 days of age. a = alveolus. Arrows in panels A-D indicate vWF-positive blood vessels.

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Figure 6.

Lung morphometry, including the quantifications of alveolar crests (A), number of pulmonary vessels (B), arterial medial wall thickness (C), septal thickness (D) and influx of macrophages (E) and neutrophilic granulocytes (F) was determined on paraffin sections in LPAR1 mutant and wild type Wistar rats on day 10. In the LPAR1 experiments pups did not receive treatment. Wild type pups served as controls in RA (open bar) and hyperoxia (shaded bar) for LPAR1 mutant rat pups kept in RA (open bar) or hyperoxia (shaded bar). In the Ki16425 experiments, RA pups were injected daily with DMSO or Ki16425 (open bars) and O₂ pups were injected daily with DMSO or Ki16425 (shaded bars): 5 mg kg⁻¹ day⁻¹ until 10 days of age. Values are expressed as mean \pm SEM. *p < 0.05, **p < 0.01 and ***p < 0.001 versus own room air controls. $^{\dagger}p < 0.05$, $^{\dagger\dagger}p < 0.01$ and $^{\dagger\dagger\dagger}p < 0.001$ versus agematched O₂-exposed controls. LPAR1^M = LPAR1^{M318R/M318R} mutant rat; Ki = Ki16425. -: daily administration of solvent (DMSO) in Ki16425 experiments or wild type rat in LPAR1 mutant rat, n=6-10 per group. Two independent experiments were performed with LPAR1 mutant rats and three for the Ki16425 intervention studies.



Figure 7.

Representative lung sections stained for collagen III (A-D) and elastin (E-L) of wild type (A, B, D, E, G, I-L) and LPAR1 mutant rat pups (C, F and H) kept in room-air (RA; A, E, F, I and J) or 100% O₂ (B-D, G, H, K and L) injected daily with 10% DMSO (A, B, I and K) or 5 mg kg⁻¹ day⁻¹ of Ki16425 (D, J and L) until 10 days of age. Quantification of collagen III deposition (M) and elastin expression (N) on lung tissue paraffin sections on day 10 in LPAR1 mutant and wild type Wistar rat pups. In the LPAR1 experiments pups did not receive treatment. Wild type pups served as controls in RA (open bar) and hyperoxia (shaded bar) for LPAR1 mutant rat pups kept in RA (open bar) or hyperoxia (shaded bar). In the Ki16425 experiments, room air pups were injected daily with DMSO or Ki16425 (open bars) and O₂ pups were injected daily with DMSO or Ki16425 (shaded bars): 5 mg kg⁻¹ day⁻¹ until 10 days of age. Values are expressed as mean ± SEM. ***p* < 0.01 and ****p* < 0.001 versus own room air controls. *p* <0.05 versus RA Wistar control. ^{†††}*p* < 0.001 versus age-matched O₂-exposed control. LPAR1^M = LPAR1^{M318R/M318R} mutant rat; Ki = Ki16425. -: daily administration of solvent (DMSO) in Ki16425 experiments or wild type rat in LPAR1 mutant rat experiments. +: daily treatment with 5 mg kg⁻¹ of Ki16425 or

LPAR1 mutant rat, n=6-10 per group. Two independent experiments were performed with LPAR1 mutant rats and three for the Ki16425 intervention studies.

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Figure 8.

Quantification of extravascular fibrin deposition in lung homogenates (A), and total protein concentration in bronchoalveolar lavage fluid (BALF; B) and CINC1 expression in BALF (C) on day 10 in LPAR1 mutant and wild type Wistar rat pups. In the LPAR1 experiments pups did not receive treatment. Wild type pups served as controls in RA (open bar) and hyperoxia (shaded bar) for LPAR1 mutant rat pups kept in RA (open bar) or hyperoxia (shaded bar). In the Ki16425 experiments, room air pups (open bars) were injected daily with DMSO or Ki16425 and O₂ pups (shaded bars) were injected daily with DMSO or Ki16425 and O₂ pups (shaded bars) were injected daily with DMSO or Ki16425 and O₂ pups (shaded bars) were injected daily with DMSO or Ki16425 in day⁻¹ until 10 days of age. Values are expressed as mean \pm SEM. **p* < 0.05, ***p* < 0.01 and ****p* < 0.001 versus own RA controls. †*p* < 0.05 versus age-matched O₂-exposed control. LPAR1^M = LPAR1^{M318R/M318R} mutant rat; Ki = Ki16425. -: daily administration of solvent (DMSO) in Ki16425 experiments or wild type rat in LPAR1 mutant rat, n=8-12 per group For both experimental models three independent experiments were performed for fibrin deposition, total protein measurement in BALF as a marker for vascular leakage and CINC1 expression in BALF.

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Figure 9.

Relative mRNA expression of Lysophosphatidic acid receptor (LPAR) 1 (A), LPAR2 (B), LPAR3 (C), LPAR4 (D), LPAR5 (E) and LPAR 6 (F) in pups on days 1, 3, 6 and 10 after birth (N=12) and in adults (N=5) during normal development (white bars) and after exposure to 100% oxygen (shaded bars). Values are expressed as mean \pm SEM. *p < 0.05, **p < 0.01 and ***p < 0.001 versus own age-matched RA-exposed controls. p < 0.05, p < 0.01 and p < 0.001 versus RA control on day 1. *p < 0.05 and ***p < 0.001 versus RA control on day 1. *p < 0.05 and ***p < 0.001 versus RA control on day 1.

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Figure 10.

Relative mRNA expression by quantitative real time RT-PCR in lung homogenates (A-F) of chemokine-induced neutrophilic chemoattractant-1 (CINC1; A), tissue factor (TF; B), plasminogen activator inhibitor 1 (PAI-1; C), LPAR1 (D), LPAR2 (E) and LPAR3 (F) on day 10 in LPAR1 mutant and wild type Wistar rat pups. Levels of mRNA expression are normalized to that of β -actin. In the LPAR1 experiments pups did not receive treatment. Wild type pups served as controls in RA (open bar) and hyperoxia (shaded bar) for LPAR1 deficient rat pups kept in RA (open bar) or hyperoxia (shaded bar). In the Ki16425 experiments, RA pups (open bar) were injected daily with DMSO or Ki16425 (open bars) and O₂ pups (shaded bar) were injected daily with DMSO or Ki16425 (shaded bars): 5 mg kg⁻¹ day⁻¹ until 10 days of age. Values are expressed as mean \pm SEM. *p < 0.05, **p <0.01 and ***p < 0.001 versus own RA controls. $^{\dagger}p < 0.05$ and $^{\dagger\dagger}p < 0.01$ versus agematched O₂-exposed controls. LPAR1^M = LPAR1^{M318R/M318R} mutant rat; Ki = Ki16425. -: daily administration of solvent (DMSO) in Ki16425 experiments or wild type rat in LPAR1 mutant rat experiments. +: daily treatment with 5 mg kg⁻¹ of Ki16425 or LPAR1 mutant rat, n=8-10 per group. Three independent experiments were performed for both experimental models.

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Figure 11.

Right ventricular hypertrophy (RVH) was determined in adult rats and in rat pups on day 10 in paraffin sections stained with haematoxylin and eosin (A, B) of LPAR1 mutant and Wistar control rat pups kept in room-air (RA) or 100% O₂ (A) and Wistar rat pups kept in room-air (RA) or 100% O₂ injected daily with 10% DMSO or 5 mg kg⁻¹ day⁻¹ of Ki16425 (B) until 10 days of age. RVH was depicted as RV/LV free wall thickness ratio (C) after measuring RV (D) and LV (E) free wall thickness in room air (RA) and age-matched O₂exposed pups (O₂). Wild type pups served as controls in RA (open bar) and hyperoxia (shaded bar) for LPAR1 mutant rat pups kept in RA (open bar) or hyperoxia (shaded bar). In the Ki16425 experiments RA pups (open bar) were injected daily with DMSO or Ki16425 and O₂ pups (shaded bar) were injected daily with DMSO or Ki16425: 5 mg kg⁻¹ day⁻¹ until 10 days of age. Data are expressed as mean \pm SEM. *p < 0.05 and ***p < 0.001 versus own room air controls. p < 0.01 versus RA Wistar control, $\dagger \dagger p < 0.01$ versus age-matched O₂-exposed controls. LPAR1^M = LPAR1^{M318R/M318R} mutant rat; Ki = Ki16425. -: daily administration of solvent (DMSO) in Ki16425 experiments or wild type rat in LPAR1 mutant rat experiments. +: daily treatment with 5 mg kg⁻¹ of Ki16425 or LPAR1 mutant rat, n=8-12 pups per group and n=5 in adult rats. Three independent experiments were performed for both experimental models.

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Figure 12.

Representative pictures of neonatal arterial pole of wild type (A,C,D) and LPAR1 mutant (B,E,F) rat pups 7 days after birth, showing macroscopically a whitish functionally closed ductus arteriosus (A,B). Microscopically (C,D and E,F) no vascular lumen in the ductus arteriosus could be detected indicating the irreversible process of anatomical closure has occurred in both the WT (C and D) and the LPAR1 mutant (E and F). Boxed areas in panels C and E are depicted in panels D and F, respectively. Ao = aorta, DA = ductus arteriosus, PT = pulmonary trunk, RV = right ventricle.

Table 1

Sequences of oligonucleotides for forward and reverse primers for real-time RT-PCR

Gene Product	Forward Primer	Reverse Primer
CINC1	5'-GCACCCAAACCGAAGTCATA-3'	5'-GGGGACACCCTTTAGCATCT-3'
LPAR1	5'-TGTGCTGGGTGCCTTTATTG-3'	5'-GGCAACACACATCGAGCAGTA-3'
LPAR2	5'-CACTGCCTCTGTGACTTGGA-3'	5'AGACAAGCAGGCTGGATAGG 3'
LPAR3	5'-ACACGAGTGGCTCCATCAG-3'	5'-CCACGAAGGCTCCTAAGACA-3'
LPAR4	5'-ACCATCAGGACCAGGAGGAAT-3'	5'-CCACTGAGGACTAGGATCCAGACT-3'
LPAR5	5'-TCCTACTGGCCAACCTCATC-3'	5'-GGCCTGAAGGCTGTTCTTTA-3'
LPAR6	5'-CTCTGCATTGCTGTCTCCAA-3'	5'-TGAGTTCTGGATCGTGTCTGA-3'
MCP1	5'-ATGCAGTTAATGCCCCAGTCA-3'	5'-TTCTCCAGCCGACTCATTGG-3'
PAI-1	5'-AGCTGGGCATGACTGACATCT-3'	5'-GCTGCTCTTGGTCGGAAAGA-3'
TF	5'-CCCAGAAAGCATCACCAAGTG-3'	5'-TGCTCCACAATGATGAGTGTT-3'
β-actin	5'-TTCAACACCCCAGCCATGT-3'	5'-AGTGGTACGACCAGAGGCATACA-3'