Endothelial-derived FGF2 contributes to the progression of pulmonary hypertension in humans and rodents.

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Endothelial-derived FGF2 contributes to the progression of pulmonary hypertension in humans and rodents

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Pulmonary hypertension (PH) is a progressive, lethal lung disease characterized by pulmonary artery SMC (PA-SMC) hyperplasia leading to right-sided heart failure. Molecular events originating in pulmonary ECs (P-ECs) may contribute to the PA-SMC hyperplasia in PH. Thus, we exposed cultured human PA-SMC to medium conditioned by P-EC from patients with idiopathic PH (IPH) or controls and found that IPH P-EC–conditioned medium increased PA-SMC proliferation more than control P-EC medium. Levels of FGF2 were increased in the medium of IPH P-ECs over controls, while there was no detectable difference in TGF-β1, PDGF-BB, or EGF levels. No difference in FGF2-induced proliferation or FGF receptor type 1 (FGFR1) mRNA levels was detected between IPH and control PA-SMCs. Knockdown of FGF2 in P-EC using siRNA reduced the PA-SMC growth-stimulating effects of IPH P-EC medium by 60% and control P-EC medium by 10%. In situ hybridization showed FGF2 overproduction predominantly in the remodeled vascular endothelium of lungs from patients with IPH. Repeated intravenous FGF2-siRNA administration abolished lung FGF2 production, both preventing and nearly reversing a rat model of PH. Similarly, pharmacological FGFR1 inhibition with SU5402 reversed established PH in the same model. Thus, endothelial FGF2 is overproduced in IPH and contributes to SMC hyperplasia in IPH, identifying FGF2 as a promising target for new treatments against PH.

Introduction

Pulmonary hypertension (PH) develops either as a complication of various disease states or as an idiopathic condition (1). Mutations in the bone morphogenetic protein receptor type 2 (BMPR2) have been linked to familial cases of idiopathic PH (IPH), but the molecular basis for common sporadic or associated forms is unknown. A major component of the pulmonary vessel remodeling process that leads to IPH is proliferation of pulmonary artery SMCs (PA-SMCs) (2), which may result either from inherent characteristics of PA-SMCs or from dysregulation of molecular events governing PA-SMC growth, such as signals originating in pulmonary ECs (P-ECs) (3).

One critical aspect of vessel development and maturation is the release of growth factors that recruit mural cells and stabilize the vessel wall (4). Thus, endothelium-derived growth factors that modulate SMC migration and proliferation during vessel maturation may be involved in abnormal vessel remodeling. Recent evidence indicates a role for dysregulation of these molecular pathways governing vessel maturation in the pulmonary vessel remodeling process that characterizes PH (1, 2). In previous studies, we found that culture medium from P-ECs induced proliferation of PA-SMCs and that this effect was considerably stronger when we used P-ECs from patients with IPH rather than from controls (3, 5). The mediators involved in this process were not identified, although part of the proliferative effect seemed related to overproduction of serotonin and endothelin-1 (ET-1) (3, 5). Moreover, we previously reported that the release of endothelium-derived growth factors capable of recruiting vessel-wall cells was influenced by the angiopoietin/Tie2 pathway (5). In the present study, we investigated whether growth factors known to participate in vessel development and EC/pericyte interactions were involved in this abnormal crosstalk between P-ECs and PA-SMCs during the progression of PH. We focused on FGF2 because, among the main growth factors expressed by ECs (including PDGF, TGF-β, EGF, and FGF2), only FGF2 was released in excessive amounts by P-ECs of patients with IPH. FGF2, a member of a large family of heparin-binding growth factors (6, 7), is synthesized by several cell types including tumor cells, fibroblasts, ECs, and macrophages (8–10). Moreover, FGF2 can be sequestered and stored as a complex in the extracellular matrix, then released by proteolytic processes to bind and activate cell targets, thereby promoting mitogenesis (11–14). FGF2 exerts its biological activity by binding to high-affinity tyrosine kinase FGF receptors (FGFRs), such as FGFR1, expressed on the surface of vascular cells (15–18).

Increased lung and circulating FGF2 levels have been reported in both experimental and human PH. Abnormally high levels of FGF2 were found in the blood of 51% and in the urine of 21% of patients with IPH (19) and in 2 animal models, a lamb model of PH developed by inserting an aortopulmonary vascular bypass graft (20) and the rat model of monocrotaline-induced (MCT-induced) PH (8). However, the importance of the FGF2 increase in initiating and perpetuating the disease remains unclear.
The objective of this study was to investigate the role for FGF2 in crosstalk between P-ECs and PA-SMCs in humans with IPH as well as in the development and perpetuation of experimental PH in rats. To this end, we developed siRNA to induce knockdown of human FGF2 in experiments designed to assess interactions between P-ECs and PA-SMCs from patients with IPH and from controls. We then conducted experiments in rats to determine whether repeated administration of siRNA directed against rat FGF2 and inhibition of FGFRs prevented and/or reversed MCT-induced pulmonary vascular remodeling and PH.

Results

Growth factors released by ECs from patients with IPH and controls. Levels of FGF2 protein were markedly increased in the medium of quiescent P-ECs from patients with IPH compared with those from controls (Figure 1A). In contrast, no differences were found between the 2 groups regarding the levels of TGF-β, PDGF, or EGF (Table 1). Measurements of FGF2 protein in cell culture medium showed greater FGF2 protein production by P-ECs than by PA-SMCs (Figure 1A). Moreover, adding exogenous human leukocyte elastase to cultured P-ECs resulted in significantly greater FGF2 release from the extracellular matrix of IPH P-ECs than that of control P-ECs (data not shown).

FGF2 expression in lungs from patients with IPH and controls. As shown in Figure 1B, whereas mRNA expression showed only a small nonsignificant increase, a marked increase in FGF2 protein was found in lung homogenates from patients with IPH compared with controls. In situ hybridization on lung sections showed that FGF2 mRNA was predominantly located in P-ECs of pulmonary arteries in both the patients with IPH (Figure 1F–J) and the controls (Figure 1E and H). In agreement with the PCR and ELISA results, in situ hybridization showed that P-ECs from patients with IPH generated a strong FGF2 signal, which predominated in remodeled pulmonary arteries. The FGF2 signal was weak in vessels from controls. RNA in situ hybridization with a sense probe produced little or no signal compared with that using an antisense probe (Figure 1K–M).

FGF2 expression by P-ECs and effect on proliferation of PA-SMCs. To evaluate the contribution of endothelium-derived FGF2 to the stimulation of PA-SMC growth, we treated cultured P-ECs from lungs of patients with IPH or controls with siRNA against human FGF2 (FGF2-siRNA) or with a scrambled sequence of this siRNA (control siRNA). We then added the treated P-EC medium to cultured PA-SMCs. PA-SMC growth responses were measured using [3H]thymidine incorporation. We first determined that P-EC transfection by FGF2-siRNA was effective in inhibiting FGF2 expression compared with transfection by the scrambled siRNA sequence. FGF2 mRNA levels were reduced by 10% in P-ECs from controls and by 55% in P-ECs from patients with IPH (Figure 2). Adding P-EC medium stimulated PA-SMC growth; however, the growth increase was larger when medium from IPH P-ECs was used compared with medium from control P-ECs (Figure 3A). PA-SMC growth responses were similar with medium of untreated P-ECs and medium of P-ECs treated with control siRNA (Figure 3, A and B). In contrast, as shown in Figure 3, B and C, PA-SMC growth was...
60% lower in IPH P-ECs treated with FGF2-siRNA than in IPH P-ECs treated with control siRNA. This difference was only 10% when P-ECs from controls were used.

Exogenous FGF2 induced a mitogenic effect that was similar in PA-SMCs from patients with IPH and from controls, and this effect was not altered by EGFR or PDGFR inhibitors. In contrast, pretreatment with the FGFR1 antagonist SU5402 in a dose of 10−8 M completely inhibited FGF2-induced proliferation (Figure 3D). Similarly, using real-time quantitative PCR, we found no significant difference in FGFR1 mRNA levels between IPH PA-SMCs and control PA-SMCs (2.75 ± 0.5 vs. 2.68 ± 0.4, respectively, n = 5; P = NS).

FGF2 expression in MCT-injected rats after treatment with FGF2-siRNA, scrambled siRNA sequence, or vehicle. To evaluate the effect of FGF2-siRNA in MCT-injected rats, FGF2 expression was measured in rat lungs at various times after MCT injection. FGF2 mRNA levels showed an early increase after 12 hours, which continued until day 5, then further increases on day 15 and day 21 after MCT administration. FGF2 mRNA levels were higher in MCT-injected rats than in control rats (Figure 4).

In pretreatment, the first FGF2-siRNA injection compared with control siRNA blunted the early increase in FGF2 mRNA and protein observed 12 hours after MCT injection. A larger FGF2 reduction was seen 120 hours after MCT administration, lung FGF2 mRNA and protein observed 12 hours after MCT injection. A larger FGF2-siRNA reduction was seen 120 hours after MCT administration.

Table 1

<table>
<thead>
<tr>
<th>Controls (n = 5)</th>
<th>IPH (n = 5)</th>
</tr>
</thead>
<tbody>
<tr>
<td>FGF2 (pg/ml)</td>
<td>2.10 ± 1.05</td>
</tr>
<tr>
<td>PDGF-BB (pg/ml)</td>
<td>2.40 ± 0.50</td>
</tr>
<tr>
<td>EGF (pg/ml)</td>
<td>0.75 ± 0.15</td>
</tr>
<tr>
<td>TGF-β1 (pg/ml)</td>
<td>163.3 ± 10.9</td>
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Growth factors released by P-ECs from controls and from patients with IPH


MCT-induced PH in rats is related to PA-SMC proliferation, collagen accumulation, and inflammation (21–24). Therefore, we evaluated the effect of FGF2 knockdown on these 3 events. Proliferating cell nuclear antigen (PCNA) immunohistochemistry results indicated that FGF2-siRNA markedly reduced SMC proliferation within the arterial wall (Figure 6, G–I). Moreover, FGF2-siRNA reduced both the accumulation of collagen fibers stained blue by Masson trichrome stain (Figure 6, J–L) and the infiltration of macrophages as assessed by immunohistochemistry using the macrophage-specific marker CD68.

Evaluation of the efficacy of FGF2-siRNA treatment in MCT-induced PH. In rats treated with 2 FGF2-siRNA injections on days 21 and 25 after the MCT injection, evaluation on day 42 showed marked decreases in PAP (Figure 7A), RV/(LV + S) (Figure 7B), and distal artery muscularization (Figure 7C) compared with rats given 2 injections of control siRNA. Systemic artery pressure and heart rate on day 42 were not significantly different from the values in control rats given saline instead of MCT (data not shown).

FGF2-siRNA injections decreased PA-SMC proliferation (Figure 7, F and G), collagen fiber accumulation (Figure 7, H and I), and macrophage infiltration (Figure 7, J and K) compared with injections of control siRNA.

Reversal of MCT-induced PH with the FGFR1 inhibitor SU5402. To confirm that the decreases in pulmonary vascular alterations and PH associated with FGF2-siRNA treatment were related to FGF2 knockdown, indicating a key role for FGF2 overproduction in PH, we investigated to determine whether the selective FGFR1 inhibitor SU5402 prevented and/or reversed PH induced by MCT in rats. In rats treated with SU5402 on days 21 to 42 after the MCT injection, evaluations on day 42 showed marked decreases in PAP, RV/(LV + S), and distal artery muscularization compared with rats treated with the vehicle (saline) (Figure 8).

Discussion

The present results show that dysregulated FGF2 signaling contributes to pulmonary vessel remodeling during the progression of PH. Overproduction by P-ECs of FGF2 that acts on PA-SMCs appears to be a critical component of the abnormal crosstalk between these
2 cell types in human IPH. Using siRNA against human FGF2, we showed that endothelial FGF2 promoted PA-SMC growth and that this effect was stronger with P-ECs from IPH patients than with those from controls. The crucial role for endothelial FGF2 on PH development was further supported by experiments in rats with MCT-induced PH. FGF2-siRNA not only prevented the development of PH but also reversed established PH. Treatment with FGF2-siRNA reduced PA-SMC proliferation in situ, extracellular matrix accumulation, and perivascular inflammatory-cell infiltrates. Consistent with these findings, FGF pathway inhibition via FGFR1 blockade by the selective inhibitor SU5402 reversed MCT-induced PH. Taken together, these results suggest that dysregulation of FGF2 signaling may be a treatment target in PH in keeping with data obtained in studies of several malignant disorders.

In previous studies (3, 5), we found that excessive release of serotonin and ET-1, 2 soluble growth-promoting factors acting on PA-SMCs, was an intrinsic abnormality of P-ECs from patients with IPH. Although increased release of these mediators contributes to the exaggerated stimulating effect of P-EC medium from patients with IPH on PA-SMC growth compared with P-EC medium from controls, it is not the only source of this paracrine effect. We measured growth factors in the medium of P-ECs from patients with IPH and from controls. Whereas the amounts of PDGF, EGF, and TGF-β did not differ, the amount of FGF2 released by P-ECs from patients was 10 times greater than the amount released by P-ECs from controls. This difference was even more pronounced when the cells were incubated with human leukocyte elastase (data not shown) because FGF2 can be sequestered and stored in the extracellular matrix, then released by proteolytic processes (11–14, 25, 26). Because the angiopoietin-1/Tie2 pathway is potentiated in IPH and contributes to PA-SMC hyperplasia via increased stimulation of serotonin and ET-1 synthesis by P-ECs (5), we investigated the effect of FGF2 on this pathway. Levels of mRNAs for ET-1 and Tie2 were not significantly different between IPH P-ECs stimulated with exogenous FGF2 and IPH P-ECs stimulated with the vehicle (data not shown), suggesting that FGF2 did not affect or modulate the contribution of the angiopoietin-1/Tie2 pathway to the aberrant PA-SMC proliferation that characterizes the pathogenesis of PH.

P-EC treatment with FGF2-siRNA induced 60% inhibition of the PA-SMC growth-promoting effect of P-EC medium from patients with IPH, whereas only 10% inhibition was observed with
Increased lung and circulating FGF2 levels have been reported in both experimental and human PH. Abnormally high levels of FGF2 have been reported in the blood of 51% and in the urine of 21% of patients with IPH (19) as well as in 2 animal models, a lamb model of PH developed by inserting an aortopulmonary vascular bypass graft (20) and the MCT-induced PH rat model (8). However, the importance of the FGF2 increase in initiating and perpetuating the disease was unclear. Increased FGF2 expression has also been documented in various malignancies in which FGF2 acts on the tumor cells via either autocrine or paracrine mechanisms (29–32). In our study of patients with IPH, increased FGF2 expression was found in the lung and the FGF2 signal was predominantly located in the endothelium of pulmonary vessels as well as in cultured P-ECs removed from their abnormal in vivo environment. These results are reminiscent of those obtained using embryonic or cancer stem cells maintained in long-term cultures, which overexpress endogenous FGF2 (7). Thus, our present results strongly suggest that FGF2 overproduction may be closely linked to the abnormal P-EC phenotype in IPH and, therefore, to the pathogenesis of IPH.

To assess the in vivo effect of FGF2 on the development of PH, we investigated the role of FGF2 in rats with experimental PH induced by MCT injection. When administered s.c., MCT is converted to its pneumotoxic metabolites in the liver; these are transported to the lungs, where they cause early injury to the pulmonary arterial endothelium and an inflammatory response before the onset of PA-SMC proliferation and the development of PH (21–24, 33, 34). Alterations in the structure of the pulmonary vessel walls include (a) thickening of the medial layer of the SMCs, (b) increased extracellular matrix synthesis and deposition, and (c) increased thickness of the adventitial layer because of edema and mononuclear inflammatory cell infiltrates composed mainly of macrophages (21–24). This increased pulmonary vascular remodeling causes an increase in PAP that eventually leads to RV hypertrophy. Morbidity and mortality in MCT-treated rats are known to be linked to vascular disease (24). Although the vascular changes that occur in response to MCT injection in rats do not resemble the complex pulmonary vascular lesions seen in patients with primary or advanced secondary PH, our results support a pathophysiological role for FGF2 in the development of PH, where FGF2 may serve as a link between altered EC signaling and pulmonary vascular remodeling. First, as previously described (8), we found that lung FGF2 expression increased markedly during the development of MCT-induced PH and that this increase was detected as early as 12 hours after injection at a time when PA-SMC proliferation was not yet detectable. Second, we found that pretreatment of MCT-injected rats with FGF2-siRNA before the MCT injection was followed by marked attenuation of PH development, as assessed on day 21 based on decreases in PAP, RV hypertrophy, and distal pulmonary artery muscularization, compared with pretreatment of MCT-injected rats with the scrambled siRNA sequence. These data constitute strong evidence that upregulation of lung FGF2 in response to MCT contributes to the initiation of PA-SMC proliferation and subsequent development of PH. Treatment of rats with FGF2-siRNA prior to MCT injection prevented the early rise in FGF2, and repeated administration of FGF2-siRNA 24 hours after MCT not only prevented the increase but also induced a major decrease in FGF2 mRNA and protein levels on day 5. In contrast, in the heart and liver, FGF2-siRNA did not significantly affect FGF2 mRNA or protein levels compared with control siRNA (data not shown). Third, we found that treatment with FGF2-siRNA not only prevented but also almost completely reversed established PH, suggesting that a sustained increase in lung FGF2 expression may be necessary for both the progression and the maintenance of MCT-induced PH. Consistent with this finding, blocking FGF2 signaling by the selective FGFR inhibitor SU5402 reversed established MCT-induced PH in rats, decreasing the values of PAP, RV hypertrophy, and peripheral artery muscularization to the levels observed in vehicle-treated rats.
Because PA-SMC hyperplasia is among the main pathological changes in patients with PH, we focused on the growth-promoting effect of FGF2 on PA-SMCs and found strong inhibition of SMC proliferation by FGF pathway blockade with either FGF2-siRNA or SU5402. However, we cannot exclude the possibility that FGF blockade acts via a synergistic effect on apoptosis. Pardo et al. reported that FGF2 increased the expression of the antiapoptotic proteins XIAP and Bcl-XL through the PKCε signaling pathway to inhibit apoptosis of lung cancer cells (35). Moreover, Xiao et al. reported that FGF2 inhibited apoptosis in human small-cell lung cancer cells via upregulation of the survivin protein and decreased release of Smac from mitochondria to the cytoplasm (36). Moreover, the multifunctional role for the FGF signaling pathway may not be confined to direct effects on PA-SMCs during PH progression. FGF2 not only increases PA-SMC proliferation but also directly modulates proteolytic processes via effects on the MMPs and urokinase-type plasminogen activator/plasmin systems, e.g., via regulation of the expression of collagenase (37), matrilysin (38), stromelysin-1 (39), and the urokinase-type plasminogen activator (40, 41), which contribute both to pulmonary vascular remodeling and to the release of growth factors such as FGF2 from the extracellular matrix. FGF2 also stimulates the expression of adhesion molecules on ECs, thereby promoting the recruitment of inflammatory cells and potentiating the inflammatory process (42). In addition to FGFR, activation of 2 other tyrosine kinase receptors, EGFR and PDGFRβ, have been implicated in the pathogenesis of PH (43, 44): Merklinger et al. reported that selective blockade of EGFR signaling induces regression of medial hypertrophy through SMC apoptosis and reverses progressive PH (43), while Schermuly et al. reported that the PDGF

Figure 5
Validation of acute and chronic lung FGF2 knockdown by siRNA. (A) FGF2 mRNA and protein levels in rat lung 12 hours and 120 hours after MCT injection. Rats were injected with FGF2-siRNA or the scrambled siRNA sequence either once, 3 days before MCT injection (12-hour group), or twice, 3 days before and 1 day after MCT injection (120-hour group). (B) FGF2 mRNA and protein levels in rat lungs 21 days after MCT injection. Rats were injected with FGF2-siRNA or the scrambled siRNA sequence (2 [2iv] or 4 [4iv] injections). FGF2 mRNA levels were assessed using real-time quantitative PCR and protein levels with FGF2 immunoblotting (the lanes were run on the same gel but were not contiguous). All values are mean ± SEM from at least 5 animals in each group. *P < 0.05; **P < 0.01; ***P < 0.001 vs. rats injected with saline instead of MCT or vs. rats treated with the scrambled siRNA sequence.
Figure 6

Evaluation of the efficacy of FGF2-siRNA pretreatment in MCT-induced PH. Results obtained in rats treated with FGF2-siRNA or the scrambled siRNA sequence before the MCT injection and studied on day 21. Compared with the scrambled siRNA sequence, FGF2-siRNA significantly prevented the development of PH, as assessed by (A) PAP; (B) RV hypertrophy reflected by the RV/(LV + S) weight ratio; and (C) percentages of NM, PM, FM, and FM+ intra-acinar vessels. (D–F) Hematoxylin-phloxine-saffron stain; (G–I) PCNA immunostaining; (J–L) Masson trichrome stain; and (M–O) CD68 immunostaining of rat lungs. *P < 0.05; **P < 0.01; ***P < 0.001 vs. rats injected with saline instead of MCT or vs. rats treated with scrambled siRNA sequence. Scale bars: 25 μm.
Figure 7
Evaluation of the efficacy of FGF2-siRNA treatment in MCT-induced PH. Results obtained in rats treated with FGF2-siRNA or the scrambled siRNA sequence (2 injections) on day 21 and day 25 after MCT injection (treatment) and studied on day 42. Compared with the scrambled siRNA sequence, FGF2-siRNA significantly reversed the development of PH, as assessed by (A) PAP; (B) RV hypertrophy reflected by the RV/(LV + S) weight ratio; and (C) percentages of NM, PM, FM, and FM+ intra-acinar vessels. (D and E) Hematoxylin-phloxine-saffron stain; (F and G) PCNA immunostaining; (H and I) Masson trichrome stain; and (J and K) CD68 immunostaining of rat lungs. *P < 0.05; **P < 0.01; ***P < 0.001 vs. rats injected with saline instead of MCT or vs. rats treated with scrambled siRNA sequence. Scale bars: 25 μm.
receptor antagonist imatinib mesylate reverses pulmonary vascular remodeling in 2 different animal models of PH (44). In the present study, we did not observe a direct relationship between FGF2 and the expression or activation of EGFR or PDGFRβ, but it is possible that these key elements affect common critical mediators of vascular remodeling in similar ways.

Methods

Patients with pulmonary arterial hypertension and controls. We studied lung specimens obtained during lung transplantation in 8 patients with IPH and during lobectomy or pneumonectomy for localized lung cancer in 8 controls. Age (mean ± SD) was 34 ± 10 years in the patients with IPH and 56 ± 12 in the controls. Mean PAP in the patients with IPH was 66 ± 17 mmHg. Preoperative echocardiography was performed in the controls to rule out PH, and the lung specimens from the controls were collected at a distance from tumor foci. This study was approved by the local ethics committee (CPP Ile-de-France VII, Le Kremlin-Bicêtre, France), and patients provided informed consent prior to their contribution to the study.

Isolation and culture of P-ECs and PA-SMCs. Human P-ECs and PA-SMCs were isolated and cultured as previously described (3, 45). To characterize the endothelial cell phenotype, P-ECs were labeled with acetylated low-density lipoprotein coupled to a fluorescent carbocyanine dye (1,1′-diocadecyl-3,3′,3′-tetramethylindocarbocyanine perchlorate [DiI-Ac-LDL]) and stained with antibodies against the endothelial cell–specific Ulex europaeus agglutinin-1 (UEA-1) Cells positive for DiI-Ac-LDL and UEA-1 and negative for desmin and vimentin were classified as P-ECs and contributed more than 95% of the cell-culture population. To identify PA-SMCs, we used between passages 3 and 6. To determine whether FGF2 promotes SMC proliferation assessed by (3H)thymidine incorporation, we examined cultured cells for expression of muscle-specific contractile and cytoskeletal proteins including smooth-muscle α-actin, desmin, and vinculin. Cells were used between passages 3 and 6.

Measurements of production of FGF2 and other growth factors by P-ECs. To quantify SMC growth factors in P-EC culture medium, we seeded P-ECs from controls and patients with IPH on 6-well plates at a density of 3 × 10⁶ cells/well and allowed the cells to adhere. After 24 hours, the cells were incubated in serum-free MCDB131 for 24 hours. PDGF-BB, TGF-β1, FGF2, and EGF were measured in the P-EC culture medium using ELISAs (R&D Systems).

FGF2 knockdown in P-ECs by siRNA. To determine whether FGF2 produced by P-ECs affected PA-SMC proliferation, we seeded PA-ECs from controls and patients with IPH, then grew them to 70% confluence in serum-free medium. The media were collected for treatment of PA-SMCs, whereas the cells were harvested for measurement of FGF2 mRNA. The siRNA duplex sequences used to target FGF2 (FGF2-siRNA) were as follows: sense, 5′-GCAAGAAGCAAUCGGGCGAU-A-3′; and antisense, 5′-UAUACUGCGCAUGCUGUUCAGUGC-3′. The scramble sequences were as follows: sense, 5′-GCAAGAAGCAAUCGGCGAUCAUA-3′; and antisense 5′-AUGAUUCCCGCGAUUCCUUGC-3′.

SMC proliferation assessed by (3H)thymidine incorporation. PA-SMCs in DMEM supplemented with 15% FCS were seeded in 24-well plates at a density of 5 × 10⁴ cells/well and allowed to adhere. The cells were subjected to 48 hours of growth arrest in serum-free medium, then treated with 1 ml of conditioned P-EC medium. We also tested the effect of exogenous PDGF (10 ng/ml) and FGF2 (10 ng/ml) on PA-SMC proliferation with or without imatinib (10⁻⁸ M), EGF antagonist (10⁻⁸ M; 324674, Calbiochem), and SU5402 (10⁻⁵ M; 572630, Calbiochem). Under each condition, (3H)thymidine (1 μCi/ml) was added to each well. After incubation for
24 hours, the cells were washed twice with PBS, treated with ice-cold 10% trichloroacetic acid, and dissolved in 0.1 N NaOH (0.5 ml/well). The incorporated radioactivity was counted and reported as cpm/well.

Real-time quantitative RT-PCR for measurement of FGF2 and FGFR1 mRNA levels. FGF2 expression was examined in lungs and P-ECs from patients with IPH and from controls. Levels of FGF2 mRNA were measured also in rat lungs at various times between day 1 and day 21 after the MCT injection. Finally, FGFR1 expression was assessed in PA-SMCs from patients with IPH and from controls.

Total RNA was extracted using the Qiagen RNeasy Mini Kit (Qiagen) according to the manufacturer’s instructions. RNA concentrations were determined using standard spectrophotometric techniques, and RNA integrity was assessed by visual inspection of ethidium bromide-stained agarose gels.

Reverse transcription was performed using random hexamer primers and reverse transcriptase (Invitrogen). For PCR, we used Primer Express Software, version 2.0 (Applied Biosystems) to design a sense primer (5’-ACGCGTCCGGAAGA-3’) and an antisense primer (5’-ACATCTGGTGGAACAAT-3’) for FGF2 and a sense primer (5’-TCTGGTCCGGAACAC-3’) and an antisense primer (5’-TTG-GGGAAGCTAATCGAGA-3’) for FGFR1. To avoid inappropriate amplification of residual genomic DNA, intron-spanning primers were selected and internal control 18S rRNA primers provided. For each sample, the amplification reaction was performed in duplicate using SYBR Green Mix (Applied Biosystems) and specific primers. Signal detection and result analysis were achieved using ABI PRISM 7000 sequence detection software (Applied Biosystems). Expression of the gene of interest was computed relative to the expression of the internal standard r18S, as follows: mRNA = 1/2^ΔΔCt gene of interest - Ct r18S).

In situ hybridization. In situ hybridization was performed on 5-µm thick sections of lungs from patients with IPH and from controls, using 3’-[35S]-labeled oligonucleotide probes specific for human FGF2. Terminal deoxynucleotidyl transferase (TdT) was used to label 1 pmol of oligonucleotide probe in 1 x TdT buffer (GE Healthcare) with a 20-fold excess of [35S]dATP (PerkinElmer) for 1 hour at 37°C. Probes were purified with the QiAquick Nucleotide Removal Kit (Qiagen). Sections were equilibrated at room temperature, then covered with 200 µl of hybridization mixture containing the [35S]-labeled probe (106 cpm/µl) diluted in hybridization buffer. Hybridization was then allowed to occur overnight in a moist chamber at 42°C. Slides were rinsed in sodium chloride/sodium citrate (150 mM NaCl, 15 mM Na2HPO4, pH 7.0), dehydrated, and dried, as described elsewhere (46). Sections hybridized to the sense probe served as the negative control. Sections were exposed to film as described above and developed after 7–14 days.

The sections were finally dipped in Kodak NTB2 nuclear track emulsion at 2. In the treatment, the rats were left untreated for 21 days, then divided randomly 50 rats into 5 groups of 10 animals. In the pretreatment protocol, 1 group received 2 i.v. FGF2-siRNA injections according to protocol 1, 1 group received 4 injections according to protocol 2, and 1 group received control siRNA injections at the same times as in protocol 2. In the treatment, the rats were left untreated for 21 days, then divided randomly into 2 groups, of which one received FGF2-siRNA and the other received control siRNA. The FGF2-siRNA injections were given on day 21 and day 25 after the MCT injection.

Effect of treatment with SU5402 on established MCT-IPH. To assess the potential effects of FGF2 knockdown used before and after the development of PH induced by MCT (60 mg/kg s.c.), we randomly divided 50 rats into 5 groups of 10 animals. In the pretreatment protocol, 1 group received 2 i.v. FGF2-siRNA injections according to protocol 1, 1 group received 4 injections according to protocol 2, and 1 group received control siRNA injections at the same times as in protocol 2. In the treatment, the rats were left untreated for 21 days, then divided randomly into 2 groups, of which one received FGF2-siRNA and the other received control siRNA. The FGF2-siRNA injections were given on day 21 and day 25 after the MCT injection.

Effect of treatment with SU5402 on established MCT PH. To assess the potential effects of the FGFR1 inhibitor SU5402 on established PH, adult male Wistar rats (200–250 g) were given MCT (60 mg/kg s.c.), left untreated for 21 days, then randomly divided into 2 groups (10 animals in each group), of which one was treated with SU5402 (25 mg/kg/day) and the other given the vehicle, from day 21 to day 42. All treatments were given once a day by s.c. injection (47).

In situ SMC proliferation, matrix accumulation, and inflammation. To determine the mechanism by which FGF2-siRNA diminished PH induced by MCT, PCNA labeling, collagen accumulation, and infiltrating macrophages were measured in rat lung sections. Tissue sections were deparaffinized in xylene, then treated with a graded series of alcohol washes, rehydrated in PBS (pH 7.5), and incubated with target retrieval solution (Dako) in a water...
bath at 90 °C for 20 minutes. Endogenous peroxidase activity was blocked with H2O2 in PBS (3%, v/v) for 5 minutes. Slides were washed with PBS, incubated for 30 minutes in a protein-blocking solution, and incubated for 30 minutes with anti-PCNA mouse monoclonal antibody (PC-10, 1:200; Dako) or anti-CD68 antibody (Hyclut Biotechnology) diluted 1:200. Then the slides were processed using the alkaline phosphatase LSAB+ system horseradish peroxidase detection kit (Dako). Brown color was generated by a diaminobenzidine substrate, and nuclei were counterstained with hematoxylin. Slides were stained with the collagen-specific Masson trichrome dye (48). Image analysis was performed with a charge-coupled device Iris camera (CDD Iris; Sony) coupled with a light microscope (Laborlux).

Statistics. Data are expressed as mean ± SEM. The nonparametric Mann-Whitney test was used for comparisons between 2 groups. Comparisons of data obtained at various times after MCT injection or in various treatment groups were performed using the nonparametric Kruskal-Wallis test followed, if significant, by Dunn’s test. To compare the degree of pulmonary arterial muscularization between groups, we used the nonparametric Mann-Whitney or Kruskal-Wallis test after ordinal classification of the vessels as NM, PM, FM, or FM+. The effect of siRNA injection on FGF2 expression in the lung at various times after MCT injection was evaluated by 2-way ANOVA, testing for treatment and time effects. When a time-by-treatment interaction was found, vehicle and active treatment were compared using the nonparametric Mann-Whitney test.

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