Septation of the gas-exchange saccules of the morphologically immature mouse lung requires regulated timing, spatial direction, and dosage of transforming growth factor (TGF)-β signaling. We found that neonatal hyperoxia acutely initially diminished saccular TGF-β signaling coincident with alveolar simplification. However, sustained hyperoxia resulted in a biphasic response and subsequent up-regulation of TGF-β signaling, ultimately resulting in bronchopulmonary dysplasia. Significantly, we found that the TGF-β-induced matricellular protein (TGFBI) was similarly biphasically altered in response to hyperoxia. Moreover, genetic ablation revealed that TGFBI was required for normal alveolar structure and function. Although the phenotype was not neonatal lethal, Tgfbi-deficient lungs were morphologically abnormal. Mutant septal tips were stunted, lacked elastin-positive tips, exhibited reduced proliferation, and contained abnormally persistent alveolar α-smooth muscle actin myofibroblasts. In addition, Tgfbi-deficient lungs misexpressed TGF-β-responsive follistatin and serpine 1, and transiently suppressed myofibroblast platelet-derived growth factor α differentiation marker. Finally, despite normal lung volume, Tgfbi-null lungs displayed diminished elastic recoil and gas exchange efficiency. Combined, these data demonstrate that initial suppression of the TGF-β signaling apparatus, as well as loss of key TGF-β effectors (like TGFBI), underlies early alveolar structural defects, as well as long-lasting functional deficits routinely observed in chronic lung disease of infancy patients. These studies underline the complex (and often contradictory) role of TGF-β and indicate a need to design studies to associate alterations with initial appearance of phenotypical changes suggestive of bronchopulmonary dysplasia. (Am J Pathol 2016, 186: 777–793; http://dx.doi.org/10.1016/j.ajpath.2015.11.024)
lifetimé.5–8 Although essential for life, exposing the developing lung to relative high oxygen concentration, as occurs with extreme preterm birth, is thought to adversely inhibit saccular lung development and secondary septation,9–7 resulting in bronchopulmonary dysplasia (BPD). Because protective therapeutic strategies have thus far been ineffective in reducing BPD and most infants born extremely premature continue to develop BPD,9 it is critical to identify novel therapeutic targets for translational approaches, prevention, and/or correction of BPD in patients.

BPD is primarily a disease of alveolar-capillary membrane simplification, resulting from injury to the developing saccular lung.10 Alveolar-capillary membrane simplification is linked to abnormalities in the lung mesenchyme and its associated ECM proteins and growth factors.11,12 Of the growth factors known to influence mesenchymal homeostasis, one that has received considerable interest is transforming growth factor (TGF)-β. TGF-β promotes ECM formation by increasing production of collagens and fibronectin13 and preventing ECM degradation by inhibiting proteases and up-regulating protease inhibitors.14 TGF-β can also induce expression of tropoelastin from lung fibroblasts,15 which is essential to distal lung morphogenesis.16 Signaling is accomplished when one of three ligands (TGFβ-1, TGFβ-2, or TGFβ-3) binds to common type II receptor (TBRII), which then recruits and phosphorylates one of several type I receptors.17 After activation by TBRII, type I receptors phosphorylate intracellular Smad2 and/or Smad3, which complex with co-Smad4 and translocate to the nucleus to stimulate downstream target expression. TGF-β ligands are secreted in an inactive state, bound to ECM latent TGF-β binding proteins; thus, the matrix serves as a TGF-β sink. All three ligands, their receptors, and Smad2/3 are spatiotemporally regulated throughout saccular and alveolar stages,18 suggesting they are actively involved in normal lung development. However, the downstream mediators of TGF-β signaling essential to accomplish normal alveolar septation are relatively unknown.

Exposure of the neonatal rodent lung to hyperoxia during postnatal saccular and early alveolar lung development [which occurs in the mouse approximately between birth and postnatal (P) day 7] reliably inhibits secondary septation and results in bronchopulmonary dysplasia (BPD). Because protective therapeutic strategies have thus far been ineffective in reducing BPD and most infants born extremely premature continue to develop BPD,9 it is critical to identify novel therapeutic targets for translational approaches, prevention, and/or correction of BPD in patients.

To generate TgfbilacZ knockin mice, we screened a 129SvJ mouse embryonic stem cell genomic DNA library (Genome System Inc., St. Louis, MO) by PCR and identified a bacterial artificial chromosome clone that contains the genomic DNA that codes for the Tgfb1 gene. Bacterial artificial chromosome DNA was isolated using a large-construct purification kit (Qiagen, Valencia, CA), and Southern blots were used to confirm the identity and integrity of the clone. A replacement-targeting vector was constructed where the bacterial β-galactosidase gene was knocked into the Tgfb1 gene locus. The 5′ homology arm, a 3-kb DNA fragment that contains all of the 5′ untranslated region and promoter elements, was PCR amplified using a high-fidelity Thermal Ace DNA polymerase kit (Invitrogen, Carlsbad, CA). In the PCR, these primers were engineered to produce 5′ SalI and 3′ XhoI sites, which enabled us to directionally clone the insert in frame with a promoterless IRES-lacZ-neo-loxP cassette.24 The resultant vector expressed the lacZ reporter under the control of the endogenous Tgfb1 promoter. A 3.5-kb SalI-EcoRI 3′ homology arm was cloned in frame into the 3′ end of the floxed IRES-lacZ-neo-loxP cassette. This resulted in a targeting vector where the translation start site and all of the first exon were deleted. After sequencing verification, the complete targeting construct was cloned into the thymidine kinase-negative selection vector, as described.24 Southern hybridization using both 3′ external probe and 5′ internal probes on BamHI-digested DNA was used for both primary and secondary screening. The 3′ external probe was the primary screen where the wild-type allele (8.3 kb) can be easily distinguished from the larger targeted allele (approximately 16 kb). Five correctly targeted embryonic stem clones were identified, and used to generate chimeras that were subsequently bred with C57BL/6J mice for six generations. Southern and PCR genotyping (primers 5′-TGTCCC-GTGCTTAAAGTGTGAG-3′ and 5′-CAGCAGCAGACCATTCTCA-3′ to detect null allele approximately 399 bp; 5′-TCAACAGCCACAGTGAAGG-3′ and 5′-GCCTGT-AACCATTGCCACT-3′ to detect wild-type allele approximately 520 bp) was used to identify TgfbilacZ heterozygotes, which were bred to a Cre splicer mouse (El1a-Cre) to remove the neomycin resistance cassette. Mice were backcrossed to C57BL/6J mice to remove the Cre allele, maintained on a C57BL/6J background, and interbred to generate homozygotes.

For the hyperoxia studies, two or more litters of wild-type C57BL/6J (The Jackson Laboratory, Bar Harbor, ME) mice born within 6 hours of one another were pooled and separated into two equal groups of six to eight mice by 12 hours of age. Half of the pups were maintained in a 30 × 20 × 20-inch polypropylene chamber (BioSpherix, Lacona, NY) in which the
oxygen concentration was tightly controlled at 85% O2, and the other half were maintained in room air (21% O2), as described.22 Nursing dams were rotated between groups every 48 hours to prevent oxygen toxicity to the dams, and pups were continuously exposed until harvested for analysis at P2, P4, P7, and P14. All procedures were performed with the approval of the Institutional Animal Care and Use Committee of Indiana University School of Medicine (Indianapolis, IN; protocol 10107).

Lung Histology and Morphometric Analysis

After euthanization and aortic transection, the right ventricle was injected with cold phosphate-buffered saline until the lungs were visually clear of blood. A tracheotomy was performed, and a 22-gauge syringe adapter was inserted and secured with suture. With chest closed to avoid overdistension, lungs were inflation fixed with 4% paraformaldehyde at a constant pressure of 25 cm H2O for 45 minutes, then the heart and lungs were removed en bloc and fixed in 4% paraformaldehyde overnight at 4°C, as described.22 The entire left lobe of each animal was paraffin embedded and divided into serial sections (7 μm thick) in entirety. Serial, equidistant sections through the entire left lung were hematoxylin and eosin stained, and random fields containing distal airspaces were photographed with a 20× objective on a Zeiss Axioskop2Plus Microscope (Carl Zeiss Inc., Thornwood, NY) at 1292 × 968 pixels. To assess distal airspace maturation, computer-aided morphometric analysis was performed using ImageJ software version 1.44 (NIH, Bethesda, MD) to measure mean linear intercept, nodal density, and septal thickness, as previously described.22 For all morphometric analyses, 12 nonoverlapping fields from at least six slides per animal were obtained, and at least four to eight animals per condition at each time point were examined. Only fields consisting of alveolar septa were used for analysis; fields containing large vessels, conducting airways, or sectioning artifacts were avoided.

Transmission Electron Microscopy

Transmission electron microscopy sample isolation, fixation, and processing were performed as previously described.24 Briefly, P4 lung lobes were fixed with 4% paraformaldehyde and 2% glutaraldehyde solution buffered at pH 7.4 with 0.1 mol/L sodium cacodylate. Specimens were then washed and dehydrated in an ethanol series, embedded in Polybed 812 (Hillsboro, OR). Specimens were then washed and 2% glutaraldehyde solution buffered at pH 7.4 with lead citrate and viewed with a FEI-Philips SM12 STEM (Hillsboro, OR).

Assessment of DFco and in Vivo Lung Compliance

After anesthesia and tracheotomy, an 18-gauge cannula was inserted and mice were mechanically ventilated with a flexiVent (SCIREQ, Montreal, ON, Canada) using 100% O2, a tidal volume of 10 mL/kg at a rate of 150 breaths/minute, and a positive end-expiratory pressure of 3 cm H2O. The pulmonary elastic properties were evaluated from quasistatic pressure-volume curves obtained in vivo. Mechanical ventilation of the anesthetized animals was interrupted, and the pulmonary system was cycled in step-wise increments between the end-expiratory pressure of 3 cm H2O and a maximal airway pressure of 20 cm H2O. To control for varying degrees of atelectasis and total lung capacity between animals, pressure-volume curves were normalized to the maximal inspiratory lung volume at 20 cm H2O, and only the expiratory portion of the curve was examined to determine elastic properties. The operation was performed in triplicate and averaged for each animal. Immediately after assessment of lung compliance, the diffusing factor for carbon monoxide (DFco) was assessed, as previously described.23 Briefly, lungs were inflated via a tracheal catheter containing 0.8 mL of test gas (0.5% Ne, 0.5% CO, 20% O2, and balanced nitrogen) and held for 6 seconds. Then, the exhaled gas was diluted with room air (RA) to a final volume of 2 mL, and concentrations of neon and carbon monoxide were immediately determined using a 3000Micro GC bench-top gas chromatographer (INFICON, East Syracuse, NY). DFco was calculated, as described,23 and measurements were obtained in triplicate, with each measurement separated by a 150-second period of mechanical ventilation to ensure disappearance of carbon monoxide and neon from the previous measurement. For each animal, DFco was expressed as the average of the three measurements.

Immunohistochemistry, in Situ Hybridization, and X-Gal Analysis

Immunohistochemical staining used the Vectastain Elite ABC kit (Vector Labs, Burlingame, CA) with diamobenzidine (Vector Labs) and hydrogen peroxide as chromogens. Dilution of primary antibodies was 1:100 for rabbit anti-TGFBI (ProteinTech, Chicago, IL), 1:500 for rabbit anti-follistatin (ProteinTech), 1:500 for rabbit anti-TGFBI (ProteinTech, Chicago, IL), 1:500 for rat anti-smooth muscle actin (e-SMA; Sigma-Aldrich, St. Louis, MO), 1:5000 for rat anti—Ki-67 (Abcam, Cambridge, MA), 1:25 for rat anti—Ki-67 (Abcam), 1:200 for platelet endothelial cell adhesion molecule-1 (BD Biosciences Pharmingen, San Jose, CA), and 1:5000 for rat anti—α smooth muscle actin (α-SMA; Sigma-Aldrich, St. Louis, MO). Confocal imaging of α-SMA, Pro-SPC, lacZ, and DAPI nuclear marker colocalization was performed as described.26 Proliferation and cellularity were assessed on Ki-67—stained lung sections. The proliferative index was calculated as the percentage of Ki-67+ nuclei per 20× field. Cellularity was determined as the total number of nuclei per 20× field, normalized to the percentage tissue density, and expressed as nuclei per millimeter.2 Terminal deoxyribonucleotidyl transferase-mediated dUTP nick-end labeling staining was detected in paraffin sections using the In Situ Cell Death Detection kit (Roche, Indianapolis, IN), and
elastin fiber deposition was identified via Weigert’s Resorcin Fuchsin stain (Electron Microscopy Sciences, Hatfield, PA). Wild-type littersmates were always used as age-matched controls. Sense and anti-sense [35S]UTP-radiolabeled Tgfb1,27 Tgfb1,2,3,28 Pdgf receptor-α,29 Clara cell secretory protein, and serpine 1 RNA probes were transcribed for in situ hybridization, as described.30 Specific signal was observed only with the anti-sense probe and detected in at least three consecutive sections. Isolation of tissues, fixation, processing, and whole-mount staining for β-galactosidase were performed as described.24 Subsequently, X-Gal—stained lungs were fixed, wax embedded, divided into serial sections (10 μm thick), and counterstained with eosin. For all histological analyses, 12 nonoverlapping fields from at least three slides per animal were obtained, and at least four to eight animals per condition at each time point were examined. Only fields consisting of alveolar septa were used for analysis.

Western Blot Analysis

Western blot analysis was performed as described.24,25 Proteins isolated from P2, P4, P7, and P14 left lungs of wild-type, Tgfb1lacZ, and 85% O2 hyperoxic mice were probed with TGFBI (1:250; R&D Systems, Minneapolis, MN), phospho-Smad1/5/8 (1:1000; Cell Signaling, Danvers, MA), phospho-Smad3 (1:1000; Epitomics, Burlingame, CA), phospho-Smad2 (1:1000; Cell Signaling), total Smad2,3,26 total Smad3 (1:1000; Epitomics), α-SMA (1:75,000; Sigma-Aldrich), and loading control α-tubulin (1:10,000; Sigma-Aldrich). Signal was detected via ECL-plus (Amersham, Pittsburgh, PA), and densitometric quantification of Western blot data (n = 3 to 6 lungs per genotype) was analyzed using ImageJ software version 1.44 (NIH, Bethesda, MD; http://imagej.nih.org/ij/).

qPCR Analysis

To examine Tgfβ1, Tgfβ2, and Tgfβ3 ligand expression temporal levels within neonatal wild-type and 85% O2 hyperoxic lungs, we used quantitative PCR (qPCR) on cDNA synthesized from P2, P4, P7, and P14 lungs (n = 4 lungs per exposure per time point) using a Superscript-II kit (Invitrogen). cDNA was amplified within the linear range using primers (Tgfβ1 PPM02991B-200, Tgfβ2 PPM02992A-200, and Tgfβ3 PPM02993A-200; Qiagen) and normalized to GAPDH (PPM02946E-200), using RT2 SYBR Green qPCR Mastermix Kit (Qiagen), as described.25 To examine Tgfb1, follistatin, and elastin expression in response to 85% O2 hyperoxia and within Tgfb1 nulls, we used qPCR on cDNA synthesized from wild-type, 85% O2-exposed, and Tgfb1-null P4, P7, and P14 lungs (n = 4 lungs per exposure per time point) using a Superscript-II kit. cDNA was amplified within the linear range using primers (Tgfb1, Fst, and Elastin; Qiagen) and normalized with GAPDH, as above. To assess TGF-β superfamily expression profiles in wild-type, Tgfb1lacZ, and 85% O2 hyperoxic P4 lungs, we used qPCR on cDNA synthesized from left lung isolates using a Superscript-II kit. mRNA from P4 left lungs (n = 4 pooled for each genotype) was isolated using RNAEasy (Qiagen) kit and reverse transcribed using SuperScript II Reverse Transcriptase (Thermo Fisher Scientific, Waltham, MA). Two samples per genotype were used in the Mouse TGF-β/Bmp Signaling Pathway RT2 Profiler PCR Array (Qiagen), and qPCR was performed in technical triplicate for each sample, as described.25 All qPCR data are representative of three separate experiments (n = 3 plates per genotype), and ΔACt-based fold-changes were calculated using GAPDH as a standard.

Statistical Analysis

Statistical analysis was performed with Prism software version 5.02 (Graphpad Software, San Diego, CA). Comparisons between experimental groups (RA versus O2 or wild type versus Tgfb1lacZ nulls) were made using unpaired t-test. P < 0.05 was considered statistically significant. All data are presented as means ± SEM. For statistical analysis of quantitative PCR data, cDNAs from wild-type, Tgfb1lacZ, and 85% O2 hyperoxic P4, P7, and P14 lungs (n = 4 pooled left lung samples for each genotype) were analyzed in triplicate for each transcript.

Results

Atypical TGF-β Signaling in Response to Hyperoxia Exposure during a Critical Period of Early Alveolar Development

We previously demonstrated that the TGF-β-responsive matricellular protein, periostin, was robustly expressed in the developing distal airspaces of the mouse saccular lung, and that continuous neonatal hyperoxia exposure from postnatal birth onward resulted in dynamic biphasic mis-expression.22 Initially, periostin was suppressed during early lung alveolar septation but was subsequently up-regulated at P14, concomitant with a fibrotic response.22 Furthermore, we found exposure of neonatal mouse lungs to hyperoxia restricted to as little as 4 days (from P0 to P4) resulted in persistent simplification of distal lung structure and inhibited adult lung function.22,23 Because it was unexpected that suppressed TGF-β signaling would be associated with defective alveolar septation, and to better define the contributions of the extracellular, intracellular, and downstream products of TGF-β signaling during both normal and hyperoxia-inhibited alveolar development, we used qPCR to examine the relative mRNA expression levels of all three TGF-β ligands during early alveolar development (P2 to P7) (Figure 1A). Although Tgfβ1 was the predominate isoform at P2, at P4 there was down-regulation of Tgfβ1 with relative persistence of Tgfβ2, resulting in equal expression of the two isoforms. By P7, during peak alveolar development, Tgfβ2 persisted as the dominate isoform. Although
Smad-mediated signaling is suppressed in P4 lungs (represented as means ± SEM). Morphometric quantitation (C) of Western measurement of phosphoplated Smad2 and Smad3 protein levels (D) confirms that intracellular TGF-β canonical Smad-mediated signaling is suppressed in P4 lungs (asterisk) exposed to 85% O₂, when compared with room air (RA) P4 littermate lungs and total Smad2 and Smad3 levels. E: qPCR mRNA profiling of 84 TGF-β-responsive genes reveals that, in addition to Tgfβ1 ligands, two downstream targets are statistically misexpressed (asterisk) at P4 and P14 after 85% O₂ exposure. Specifically, Tgfβ3 mRNA levels are suppressed, whereas follistatin (Fst) mRNA levels increase via 85% O₂ when compared with RA controls at P4. However, both Tgfβ1 and Fst are elevated via 85% O₂ when compared with RA controls at P14. F–I: In situ hybridization of altered Tgfβ2 (F and H) and Tgfβ1 (G and I) at P4 and P14 mRNA expression within alveolar tissue sections in animals continuously exposed to 85% O₂. Note the more punctate Tgfβ2 saccular wall expression when compared with Tgfβ2. J–M: In situ hybridization verification of normal Tgfβ2 (J and L) and Tgfβ1 (K and M) at P4 and P14 in RA control animals. Enlarged insets (F, H, J, and L) show expression at developing septal tips. N–Q: Immunohistochemical analysis confirms that the reduction in mRNA expression translates to a reduction in TGFβ1 protein expression in P4 lungs from animals continuously exposed to 85% O₂ when compared with RA. Arrows highlight expression at developing septal tips (Q and O). R: Western analysis of phosphorylated Smad2 levels in P2 to P14 lungs reveals TGF-β1 canonical Smad signaling is subsequently elevated in P14 85% O₂ exposed lungs (asterisk). Data are represented as means ± SEM (A–C, E, and R). N = 4 lungs from separate animals per group per time point (A–R). *P < 0.05 by t-test (RA versus hyperoxia at each time point; A–C, E, and R). Scale bars: 50 μm (F, G, N, and P); 10 μm (Q). Original magnifications: ×10 (N and P); ×40 (D and Q). GAPDH, glyceraldehyde-3-phosphate dehydrogenase.

Figure 1  Hyperoxia acutely diminishes saccular lung transforming growth factor (TGF)-β superfamily signaling and misexpression of down-stream targets. A: Quantitative PCR (qPCR) analysis of relative whole lung transforming growth factor (Tgf) β1, Tgfβ2, and Tgfβ3 ligand mRNA expression levels during normal postnatal saccular (P2, P4) and early alveolar (P7) lung development in room air (RA) control mice. B: qPCR assessment of Tgfβ1, Tgfβ2, and Tgfβ3 mRNA expression reveals all three ligands are suppressed in P4 lungs exposed continuously to 85% hyperoxia (oxygen) when compared with normal RA.

P4 hyperoxia significantly reduced the pSmad3:total-Smad2,3 ratio, indicating that pSmad3-mediated intracellular TGF-β signaling is statistically suppressed at these early stages by 85% O₂ exposure (Figure 1, C and D).

Moreover, to examine the broader extent of suppressed TGF-β signaling and identify TGF-β–pathway downstream targets that are altered by hyperoxia, we used expression profiling in RA and hyperoxia-exposed P4 lungs. Quantitative PCR using a custom array of 84 genes related to TGF-β/bone morphogenetic protein–mediated signal transduction, including SMAD, SMAD target genes, adhesion, and
extracellular molecules and transcription factors, revealed significant changes in transcript levels (>1.5-fold) of two downstream signaling products within hyperoxia-exposed lungs (Figure 1E), namely Tgfbi and follistatin (Fst). Fst was significantly up-regulated at P4 (Figure 1E) and is not thought to be normally expressed within the neonatal lung; it is inhibin-icin up-regulated at P4 (Figure 1E) and is not thought to be significantly down-regulated at P4 (Figure 1E), namely response to neonatal hyperoxia.14,32 Because Tgfb ligand at the mRNA level does not always cause changes in intracellular signal activity, we examined pSmad2 protein levels P2 to P14. In agreement with published data,14,32 elevated intracellular TGF-β canonical Smad-mediated signaling was evident only with prolonged, continuous, hyperoxia exposure through P14 (Figure 1R). Taken together, hyperoxia acutely diminishes saccular lung TGF-β superfamily signaling and misexpression of downstream TGF-β–responsive targets during early neonatal lung morphogenesis, coincident with physiological alveolar simplification. However, continued hyperoxia exposure results in a biphasic response and subsequent up-regulation of TGF-β signaling and downstream responder expression.

Figure 2 Expression of transforming growth factor-β–induced matricellular protein (TGFBI) during lung development. A–F: In situ hybridization analysis of Tgfbi spatiotemporal mRNA expression in distal lung sections throughout the pseudoglandular (E15; A), canalicular (E17; B), saccular (postnatal day P) 0 (C) and P4 (D)), and alveolar (P7 (E) and P14 (F)) stages of lung development. Note widespread Tgfbi expression by discrete cells throughout the saccular and alveolar walls that peaks P4 to P7 but is dramatically reduced by P14. G: Representative immunoblot of duplicate lung TGFBI protein expression (top panel) from P0, P4, P7, and P14 whole lung homogenates with α-tubulin loading control (bottom panel). H: Densitometry quantification confirms peak lung TGFBI expression coincides with alveolar-septal development at P4 to P7, followed by significant down-regulation by P14. Densitometry normalized with α-tubulin. *P < 0.05 versus P7, by one-way analysis of variance with Tukey’s post test. N = 4 lungs per group (A–H). Scale bars: 50 μm (A and B); 20 μm (C); 10 μm (D–F).

TGFBI Spatiotemporal Expression Coincides with Alveolar Septal Development

Given the close correlation of Tgfbi expression in P4 septal tips and significant down-regulation in response to hyperoxia, we examined Tgfbi mRNA throughout the major stages of lung development. In situ hybridization localized Tgfbi mRNA to the mesenchyme immediately beneath the developing epithelial tubules at embryonic day (E)14.5 during the pseudoglandular period (Figure 2A) and at E16 during the canalicular period (Figure 2B). At birth, in the early postnatal saccular stage, Tgfbi expression was absent from the large airways and concentrated in scattered cells within the developing saccules (Figure 2C). Expression intensified, expanded, and concentrated at developing septal tips during the transition from the saccular to alveolar stage at P4 (Figure 2D) and P7 (Figure 2E). However, in contrast to earlier stages, by P14 when bulk alveolar development is nearly complete, Tgfbi mRNA was largely...
undetectable in alveolar regions (Figure 2F) and restricted within the chondrogenic framework around the bronchus and bronchi airways. Whole lung analysis demonstrated peak expression of a single approximately 50-kDa TGFBI protein between P4 and P7, which coincides with saccular maturation and bulk alveolar septation (Figure 2, G and H). Western detection of TGFBI expression at P14 was similar to P0 levels and significantly decreased. Thus, peak lung TGFBI expression correlates with maximal alveolar-septal development from P4 to P7 during normal mouse lung morphogenesis, with negligible levels in adult lungs.

Genetic Ablation of Tgfbi Pinpoints TGFBI as a Required Component of Normal Alveolar Structure and Function

Because both TGFBI spatiotemporal expression pattern and its down-regulation after hyperoxia exposure appeared coincident with alveolar simplification, we hypothesized that Tgfbi is necessary for normal alveolar septal development. To test its requirement in distal lung development, we generated TgfbilacZ knockin mice in which TGFBI protein translation was abolished by knocking in a β-galactosidase reporter and floxed-neo selection cassette to replace the ATG start site, most of exon 1, and all of exon 2 of the Tgfbi allele (Figure 3A). After Cre-mediated removal of the floxed-neo selection cassette, we found that homozygous null pups were 100% viable. The TgfbilacZ knockout mice are fertile, are indistinguishable from wild-type littermates at P7 (Figure 3E), and continue to have normal growth and outward development through adulthood (Figure 3F). Western blot analysis of neonatal lung homogenates (Figure 3G) and immunohistochemistry (Figure 3, H and I) of Tgfbi-null and wild-type fetal lungs both confirmed the absence of TGFBI protein within TgfbilacZ/lacZ mutants and generation of a null allele.

Using our β-galactosidase reporter to map Tgfbi gene expression, we confirmed that lacZ expression in heterozygous TgfbilacZ animals (Figure 3, A–D) faithfully recapitulated wild-type endogenous Tgfbi expression previously observed by in situ hybridization, especially within the chondrogenic elements of the skeleton and heart. Within the lungs, lacZ expression is initially detected in E13 lung buds, restricted to the developing airways at E15 (Figure 3D) and E17 (Figure 4A), widespread throughout the peripheral lung at P7 (Figure 4B), but confined to chondrogenic cells around the mature airways and absent from the peripheral airspaces from P14 onward (Figure 4C). Although lacZ expression was similar in both heterozygotes and homozygous TgfbilacZ-null lungs at E17 (Figure 4, A and D) and P7 (Figure 4, B

Figure 3  Systemic loss of Tgfbi does not affect postnatal viability. A: Strategy for generation of TgfbilacZ mice. B: PCR genotyping detection of wild-type (WT; top panels) and null TgfbilacZ (bottom panels) alleles in Tgfbi null (−/−), heterozygous (+/−), and WT (+/+) animals. C and D: Whole-mount X-Gal staining demonstrating TgfbilacZ reporter expression in heterozygote E12 (C) and E15 embryos (arrow indicates lacZ within the developing lung buds; D). E and F: Null pups (left panel) are grossly indistinguishable from wild-type pups (right panel) at P7 and exhibit similar growth characteristics from P4 through to adulthood (Ad). G: Western blot analysis verifies the absence of transforming growth factor-β–induced matricellular protein (TGFBI) from within duplicate null P4 lungs when compared with age-matched WT lungs and α-tubulin loading control. H and I: Immunohistochemical staining for TGFBI in E15 lung sections from WT (H) and null (I) littermate embryos confirms in utero TGFBI absence in nulls. Data are represented as means ± SEM (F). N = 5 to 9 animals per group per time point (F). Scale bar = 50 μm (H and I).
and E), P14 null lungs exhibited ectopic, persistent expression in the peripheral distal lung (Figure 4F). Closer examination of cellular localization with control heterozygous lacZ-stained lung sections indicated that at E15, expression was absent from both the mesenchymal cells surrounding the terminal tubules and the peripheral mesenchyme, but confined to a band of mesenchymal cells immediately beneath the proximal epithelial tubules (Figure 4G). There was progressive expansion of cells away from the control large airways and into single cells in the surrounding mesenchyme from E17, with widespread expression concentrated in the septal tips at P7 (Figure 4H), but differed at P14 in that expression persisted in discrete cells within the septal tips (Figure 4H). The ectopic pattern of staining was reminiscent of the persistence of α-SMA+ myofibroblasts previously observed in response to chronic hyperoxia.22 Cooﬁning of Tgfbi−/− lungs with α-SMA (Figure 4I), surfactant protein-C (Figure 4J), and platelet endothelial cell adhesion molecule (Figure 4K) antibodies revealed that most lacZ+ cells were α-SMA+ (Figure 4I), indicating that Tgfbi-expressing cells are α-SMA+ myofibroblasts and not alveolar type II cells. Confocal immunofluorescence staining conﬁrmed that α-SMA is tightly associated with Tgfbi−/−-expressing cells (Figure 4L), indicating that TGFBI is expressed within the myofibroblast lineage. Furthermore, single-cell analysis of E18.5 lungs supports our data that the predominant cell type expressing Tgfbi is the myofibroblast.33

To determine the role of Tgfbi in conducting airway development and alveolar septation, we examined Clara cell secretary protein expression and measured distal airspace size (via mean linear intercept) and complexity (via nodal point density) throughout alveolar development. Clara cell secretary protein mRNA was equally expressed within columnar epithelial cells lining E17, P4, and P7, wild-type and null bronchi/bronchioles. Similarly, gross comparison of in utero wild-type and null lungs did not reveal any differences. Moreover, no differences between heterozygote and wild-type lungs were observed by histological analysis or by the analyses of speciﬁc markers described below. However, representative lung sections from wild-type (Figure 4M) and Tgfbi-null (Figure 4N) mice indicated that, although saccular structure was similar at P4, loss of Tgfbi resulted in persistent distal airspace simplification from P7 onward (Figure 4, M and N). The terminal air sacs are subdivided into smaller units via alveolarization in wild types, but in Tgfbi nulls, alveolar airspace enlargement was evident from P7 to P28, but normalized in 12-month adult lungs (Figure 4O and Table 1). However, Tgfbi-null nodal density, a measure of distal airspace complexity, was decreased from P7 through to adulthood (Figure 4P and Table 1). Signiﬁcantly, although the persistent structural differences in airspace complexity were of small magnitude, functional analysis of gas exchange, via measurement of the DFco, indicated that adult lungs lacking Tgfbi have a significant decrease in gas exchange efficiency (Figure 4Q). In addition, although total lung volume was similar in both wild-type and Tgfbi knockouts (Figure 4R), lungs of Tgfbi nulls displayed diminished elastic recoil (Figure 4S). Thus, although Tgfbi is neither required for in utero lung development nor lung airway morphogenesis, it is necessary for normal neonatal lung saccular remodeling, specifically during the initial alveolar septation stage. Moreover, Tgfbi is also required for subsequent normal lung elastic properties and gas exchange function.

Structural Analysis of Tgfbi-Null Lungs Suggests Normal Vascular Development but Ectopic Localization of Key Matrix Proteins

To begin to realize the mechanisms underlying abnormal alveolar development in Tgfbi nulls, we examined key mesenchymal components known to be vital for alveolar septation, including elastin deposition,16 vascular development,14 and α-SMA+—myofibroblast localization.35–37 At P7, when elastin is normally located in the tips of developing alveolar septa in wild-type animals (Figure 5A), in Tgfbi nulls, elastin formed bands within the alveolar walls and there was a lack of elastin1+ tips (Figure 5B). Because qPCR analysis of elastin mRNA levels did not reveal any alteration in P4 or P7 null lungs, the reduction of elastin deposition at the septal tips most likely represents either a defect in microfibril formation or a deﬁciency in other accessory proteins required for elastin ﬁber assembly. As screened by staining for the endothelial marker,

Figure 4  Phenotypic analysis of Tgfbi knockout (KO) lungs. A–F: Whole-mount X-Gal staining indicating Tgfbi lacZ reporter expression in heterozygous (+/−) lungs at E17 (A), postnatal day (P) 7 (B), and P14 (arrow indicates lacZ restricted around large airways; C) compared with Tgfbi−/− nulls (+/−) at E17 (D) and P7 (E) and P14 (F). F: Although there is similar lacZ spatiotemporal expression at E17 and P7, there is widespread ectopic lacZ reporter expression within the P14-null distal airspaces. G and H: Histological analysis conﬁrms whole-mount reporter patterns and ectopic Tgfbi lacZ expression in developing septal tips of P14 nulls (H) compared with heterozygous controls (G). I–K: Immunohistochemical analysis of X-Gal−/−stained sections demonstrates colocalization of α-SMA-smooth muscle actin (α-SMA) protein (I, marker of alveolar myofibroblasts) with Tgfbi lacZ-expressing cells (arrows) but not with Pro-SPC (marker of alveolar type II cells; J) or platelet endothelial cell adhesion molecule (PECAM) marker (K); L: Confocal immunofluorescence localization of Tgfbi-driven lacZ (gray) with α-SMA (red), Pro-SPC (green), and nuclear DAPI (blue) markers. There is colocalization of lacZ with α-SMA (asterisk), but not with Pro-SPC (arrowheads). M and N: Hematoxylin and eosin—stained inﬁltered lung sections throughout alveolar lung development in wild-type (M) and Tgfbi nulls (N) from P4 through 12 months/adulthood (Ad). O and P: Quantiﬁcation of distal airspace development of sections in M and N via mean linear intercept (O) and nodal point density (P) in wild-type (WT) and Tgfbi nulls reveals that during the period of rapid alveolar septation, loss of Tgfbi retards alveolar septation, resulting in larger and/or simpler distal airspaces from P7 onwards. Q: Functional analysis by way of measurement of the diffusing factor for carbon monoxide (DFco) in adult WT and Tgfbi nulls reveals a persistent functional insufﬁciency. R: However, left lung volume, as assessed by water displacement in adult WT and Tgfbi nulls, is equivalent. S: Assessment of lung elastance by pressure-volume loops in adult WT (broken line) and Tgfbi nulls (solid line) confi rms nulls exhibit decreased elastic recoil. Data are represented as means ± SEM (O–R). N = 4 to 9 separate animals per group per time point (A–S). *P < 0.05 by t-test (WT versus Tgfbi null at each time point). Scale bars: 50 μm (G, H, M, and N); 5 μm (I and L); 10 μm (J); 20 μm (K).
von Willebrand factor, normal vascular development was maintained in Tgfbi-null lungs (Figure 5, C and D). Although α-SMA was widely expressed at P7 in both groups, the intensity of α-SMA staining was modestly increased in Tgfbi nulls and localization of α-SMA protein within the alveolar septa was not consistently at the tips (Figure 5, E and F). Notably, at P14, when α-SMA protein expression is typically absent from the alveolar walls, Tgfbi-null lungs had persistence of alveolar α-SMA−myofibroblasts (Figure 5, G and H). Likewise, Western blot analysis confirmed α-SMA protein levels were significantly elevated within both P7 and P14 null lungs (Figure 5I).

Coincident with the first indication of diminished alveolar septation (Figure 4, M–P), Tgfbi-null lungs had statistically significant reduced cellularity at P4 and P7 (Figure 5J), which may be explained partly by a reduced proliferation index in null distal lung cells at P4 (Figure 5K). At P14, Tgfbi nulls displayed similar cellularity (Figure 5J) in association with a statistically significantly increased null proliferation index (Figure 5K). Because terminal deoxynucleotidyl transferase-mediated dUTP nick-end labeling staining at P4, P7, or P14 did not detect any up-regulation within null lungs compared with wild type, this suggests that a transient reduction of proliferation likely accounts for the observed declining differences in airspace size as the Tgfbi-null lungs mature into adulthood. Transmission electron microscopy of septal tips in wild-type (Figure 5, L and M) and Tgfbi-null (Figure 5, N and O) lungs indicated that key ultrastructural components, including collagen microfilaments and elastin (present as both amorphous elastin and microfibrils), were present in wild-type and null septa but that elastin staining was diminished at tips of stunted null septa (Figure 5N). Moreover, the septal tips of Tgfbi-null lungs lacked myofibroblasts, because the myofibroblast nuclei were instead mostly localized to the bases of underdeveloped Tgfbi-null septa (Figure 5N). Combined with the histomorphometric results (Figure 4) and the spatiotemporal expression of TGFBI (Figure 3), these structural data indicate that the absence of Tgfbi results in diminished alveolar septation and interference with normal septal tip reorganization that is necessary during secondary septa morphogenesis.

Genetic Loss of Tgfbi Results in an Imbalance of Lung Fibroblast Populations during a Critical Period of Alveolar Septal Development

Expansion via proliferation and spreading of platelet-derived growth factor receptor (PDGFR) α+ mesenchymal progenitor cells to the distal airspaces is essential for alveolar septal growth.35–38 Given the diminished proliferation and alveolar septation observed, as well as the diminished elastin and mislocalization of α-SMA in Tgfbi-null lungs, we examined Pdgfra expression (Figure 6, A–D). At P4, when distal airspace development was similar to wild types, Tgfbi-null lungs displayed normal expression of Pdgfra mRNA (Figure 6, A and C). However, unlike wild-type lungs, which continued to have widespread Pdgfra expression (Figure 6B), P7 Tgfbi nulls exhibited a paucity of Pdgfra+ cells (Figure 6D) coincident with distal airspace simplification (Figure 4, M–P). Using the same targeted TGF-β1 signaling array that identified Tgfbi as being significantly down-regulated by hyperoxia (Figure 1), two other components of TGF-β signaling, follistatin and serpine 1, were found to be up-regulated in P7 Tgfbi isolated lungs when compared with age-matched control littermate lungs. More important, Fst and Serpinel1 are associated with abnormal maturation and reduced proliferation of lung fibroblasts,26,39 both of which we observed in developing Tgfbi-null lungs. At P7, striking up-regulation and ectopic expression of FST protein were observed within the alveolar septa of Tgfbi-null lungs (Figure 6G) when compared to wild type (Figure 6E), and Fst was also found to be up-regulated in response to physiological hyperoxia-mediated alveolar inhibition (Figure 1E), suggesting that this negative regulator of TGF-β signaling may play a conserved role within both physiological and genetic mouse models of neonatal chronic lung disease. Although not as robust, serpine 1 mRNA was also up-regulated and ectopically expressed in P7 Tgfbi-null lungs (Figure 6H), when compared to wild type (Figure 6F). Thus, inhibition of alveolar septation and mislocalization of α-SMA/elastin in Tgfbi-null lungs are associated with decreased lung proliferation and misexpression of TGF-β-pathway genes related to lung fibroblast development. Moreover, this imbalance within alveolar septal fibroblast populations may underlie the observed decreased adult lung elastic recoil and poor gas exchange function.

**Discussion**

Initial Postnatal TGF-β Signaling Suppression Contributes to BPD Pathogenesis in Mice

Human saccular maturation, vital for establishment of postnatal distal lung complexity, is designed to occur in the protected, low-oxygen, uterine environment. Although

<table>
<thead>
<tr>
<th>Postnatal age (N)</th>
<th>Weight (g)</th>
<th>MLI (μm)</th>
<th>Nodal density (points per mm²)</th>
</tr>
</thead>
<tbody>
<tr>
<td>P4-WT (6)</td>
<td>2.8 ± 0.3</td>
<td>68.82 ± 1.70</td>
<td>1120 ± 35</td>
</tr>
<tr>
<td>P4-KO (5)</td>
<td>2.6 ± 0.2</td>
<td>71.41 ± 1.80</td>
<td>1069 ± 46</td>
</tr>
<tr>
<td>P7-WT (6)</td>
<td>4.4 ± 0.3</td>
<td>65.03 ± 1.07</td>
<td>1393 ± 69</td>
</tr>
<tr>
<td>P7-KO (5)</td>
<td>4.0 ± 0.3</td>
<td>73.46 ± 0.98*</td>
<td>1066 ± 61*</td>
</tr>
<tr>
<td>P14-WT (7)</td>
<td>10.0 ± 0.7</td>
<td>54.40 ± 0.40</td>
<td>2105 ± 39</td>
</tr>
<tr>
<td>P14-KO (8)</td>
<td>9.7 ± 1.0</td>
<td>57.28 ± 0.91*</td>
<td>1898 ± 56*</td>
</tr>
<tr>
<td>P28-WT (7)</td>
<td>16.9 ± 1.2</td>
<td>52.61 ± 1.24</td>
<td>2153 ± 77</td>
</tr>
<tr>
<td>P28-KO (6)</td>
<td>14.9 ± 0.9</td>
<td>57.47 ± 1.33*</td>
<td>1932 ± 30*</td>
</tr>
<tr>
<td>Ad-WT (4)</td>
<td>29.9 ± 0.3</td>
<td>49.22 ± 0.87</td>
<td>2522 ± 41</td>
</tr>
<tr>
<td>Ad-KO (4)</td>
<td>29.2 ± 1.7</td>
<td>51.17 ± 0.95</td>
<td>2339 ± 33*</td>
</tr>
</tbody>
</table>

Values are means ± SEM.

*K0*, knockout; MLI, mean linear intercept; WT, wild-type littermate.

Table 1 Growth and Distal Lung Development in the Presence and Absence of Tgfbi

The American Journal of Pathology

ajp.amjpathol.org ■ The American Journal of Pathology
little is known about the factors that control it, the process of saccular maturation is likely complex and dependent on intricate molecular networks that establish and orchestrate timely remodeling of lung structure and function in vivo. Although essential for life, prematurely exposing the immature lung to relative hyperoxia clearly inhibits saccular lung development and secondary septation by uncoupling normal molecular pathways.\textsuperscript{6,7,10} Significantly, TGF-β signaling is one of the key growth factor conduits known to influence injury, mesenchymal

![Figure 5](image-url)

Figure 5  Structural and ultrastructural examination of neonatal lung alveolar septation defects in Tgfbi-null mice. A–F: Immunohistochemical analysis of elastin (A and B), von Willebrand factor (vWF; C and D), and α-smooth muscle actin (α-SMA; E and F) localization in postnatal day (P) 7 wild-type (WT; +/+ ) and Tgfbi null (−/− ) lungs. There is a significant lack of elastin-positive P7 null tips (septal tips indicated via arrows in A +/+ and B −/− ), but vWF and α-SMA are similarly distributed in WT (C and E) and null (D and F) P7 tips. G and H: Compared with WT animals (G), there is ectopic expression of α-SMA protein in alveolar septal tips of P14 Tgfbi nulls (H). I: Western verification of elevated α-SMA levels in P7 and P14 Tgfbi null lungs. J and K: Assessment of alveolar tissue cellularity (J) and proliferation index (K) in WT and Tgfbi nulls at P4, P7, and P14. L–O: Transmission electron micrographs of septal tips at P4 in WT (L, enlarged in M) and Tgfbi null (N, enlarged in O) lungs. Normal collections of collagen microfibrils (col) and parallel collagen bundles (asterisk) are observed within both WT and blunted Tgfbi null septal tips; however, amorphous elastin (ela) deposition is diminished in stunted Tgfbi null septal tips (N, arrowhead) when compared with WT (L, arrowhead). Data are represented as means ± SEM (I–K). N = 5 to 6 separate animals per group per time point (A–O). *P < 0.05 by t-test (WT versus Tgfbi null at each time point). Scale bars: 50 μm (A–H); 2 μm (K and M); 500 nm (L and N). KO, knockout.
homeostasis, and lung development. Although mechanically ventilated preterm infants with developing lung disease have elevated TGF-β levels in tracheal aspirates, animal studies linking altered TGF-β signaling to arrested alveolar development are conflicted. Chronic hyperoxia impairs secondary septation and results in alveolar simplification associated with up-regulation of TGF-β. However, although most models associate elevated signaling after chronic exposure to moderate (85%) or severe (>95%) oxygen well into the alveolar stage, they may not accurately recapitulate the human condition, which is associated with relatively brief exposure to modest hyperoxia during the more sensitive saccular and early alveolar stages. Forced expression of ectopic TGF-β1 in postnatal mice throughout saccular and alveolar development inhibits alveolarization, but with significantly more fibrosis and inflammation than that seen in other models of inhibited alveolar septation. On the basis of these data, many have speculated that excessive TGF-β disrupts septation and should be targeted as a therapy to promote alveolarization. However, inhibiting TGF-β during development via genetic ablation of the Tgfβ-II receptor, Smad3, or latent TGF-β binding proteins results in normal lungs at birth, followed by a subsequent age-related increase in the size of alveolar spaces, challenging the assumption that TGF-β

Figure 6 Dynamic expression deviations in myofibroblast progenitor and downstream transforming growth factor (TGF)-β family signaling markers in Tgfbi null lungs during alveolar septal development. A–D: In situ hybridization detection of platelet-derived growth factor receptor α (Pdgfra) mRNA expression in developing alveoli in wild-type (A and B) and Tgfbi nulls (C and D), at postnatal day (P) 4 and P7, reveals suppression of Pdgfra in P7 null lungs. E and G: Immunohistochemical localization of follistatin (FST) protein within P7 wild-type (E) and Tgfbi-null (G) lungs indicates excessive FST deposition in null alveoli. F and H: In situ hybridization of serpine mRNA expression in developing alveoli in P7 wild-type (F) and Tgfbi-null (H) detects up-regulation within the null lung periphery. Scale bars: 50 μm (A–F); 20 μm (G and H).
up-regulation is a primary effector inhibiting initial alveolar septation.

We previously demonstrated that the TGF-β–responsive matricellular protein, periostin, is an early marker of inhibited neonatal murine lung alveolar septation, because its deposition is acutely suppressed via hyperoxia before subsequent up-regulation coincident with fibrosis. TGF-β–responsive matricellular protein, periostin, is an early marker of up-regulation is a primary effector inhibiting initial alveolar lung development. Forced genetic overexpression of TGFβ1 alone likely results in non-physiologic absolute and relative expression levels of TGF-β ligands and, therefore, may not accurately recapitulate the physiological responses to neonatal hyperoxia. Our present data, combined with the previously mentioned published works, underline the complex, often contradictory, role of TGF-β (and its isoforms) and indicate a need to carefully design studies that focus on clinically appropriate stages of alveolar lung development, as well as use strategies capable of determining the possible benefits of ligand-specific supplementation at physiological levels to protect against neonatal hyperoxic lung injury. Only then will the contribution of altered TGF-β signaling to alveolar development be fully elucidated and appropriate treatments to modulate TGF-β be designed.

Loss of the TGF-β Effector, TGFBI, Underlies Early Alveolar Structural Defects

We have identified TGFBI as a widely expressed, previously unrecognized, critical component for normal secondary alveolar septation. Not only did we correlate hyperoxia-mediated acute suppression of several stages of the TGF-β signaling apparatus and of the TGF-β–responsive Tgfβ1 gene, our expression data demonstrated that TGFBI was dynamically expressed within the septal tips (Figures 1 and 4) during alveolar lung development and was then down-regulated at the conclusion of bulk alveolarization. Moreover, Tgfβ1-null neonates exhibit both histological and functional deficits in secondary septal morphogenesis. Structurally, the genetic Tgfβ1-null lung phenotype is similar to the physiological hyperoxia-induced arrested alveolar septation phenotype at P4 to P14, with stunted septa and simplification of distal airspaces (Figures 1 and 4). In addition, both the hyperoxia-induced and Tgfβ1-null neonatal lungs exhibited diminished proliferation at P4 (Figure 5), altered α-SMA myofibroblast marker expression, and ectopic expression of FST (Figures 1 and 6). Our finding that Pdgfrα (a known marker of lung fibroblast precursors and regulator of myofibroblast differentiation in vivo and in vitro expression data suggested that the additive effect of hyperoxia and up-regulated TGF-β signaling during hyperoxia was protective of lung development. Our study differs from these reports in that we used lower concentrations of oxygen (85% versus 100%) and documented not only early down-regulation of TGFβ1, but also TGFβ2 and TGFβ3. It is probable that a fine balance of absolute and/or relative concentrations of TGF-β ligand expression is required for maintenance of normal alveolar lung development.
associated with impaired alveolar septation in mouse mutants lacking TGFBI. The prolonged and ectopic expression of both Tgfbi \(^{+/+}\) reporter and \(\alpha\)-SMA protein within the null P14 lungs is consistent with the notion that, in the absence of TGFBI, lung fibroblast differentiation is hindered. In addition, these neonatal Tgfbi null structural defects are associated with impaired gas diffusion and decreased elastic recoil in adult mice. Combined, these data demonstrate that TGFBI is transiently required for both normal alveolar remodeling and subsequent normal long-term murine lung function.

During perinatal lung development, the interstitial fibroblasts are initially exposed to TGF-\(\beta\), a stimulus for myofibroblastic differentiation.\(^{58}\) Although signaling through PDGFR\(\alpha\) is a critical event for alveolarization, because it is necessary for the appearance of alveolar myofibroblasts,\(^{35–38}\) down-regulation of signaling during later stages of alveolar development is required for septal maturation.\(^{59}\) In addition, during realveolarization after pneumonectomy, Pdgfr\(\alpha\)-expressing cells reexpand into the alveolar space, proliferate, and then down-regulate Pdgfr\(\alpha\) expression as they increase \(\alpha\)-SMA expression. Interference with temporal PDGFR\(\alpha\) signaling impairs the balance between structural and contractile alveolar fibroblasts, resulting in impaired alveolar septation.\(^{37,38,59,60}\)

Similar to our Tgfbi-null expression data (Figure 6), tracheal aspirate mesenchymal stromal cells from premature infants developing BPD express lower levels of PDGFR\(\alpha\),\(^{61}\) suggesting that defective PDGFR signaling is a primary feature of alveolar septation defects. Our results revealed that both reduced TGF-\(\beta\) signaling mediated by hyperoxia and reduced PDGFR signaling mediated by the absence of TGF-\(\beta\)-responsive TGFBI are both associated with alveolar septal defects. It is not yet clear whether TGF-\(\beta\) and PDGFR signaling are coregulated, reciprocally interact, or regulated by one another within myofibroblasts. However, there are pharmacological data that suggest imatinib and nilotinib, two BCR-ABL tyrosine kinase inhibitors, can suppress both pathways simultaneously\(^{62}\) to reduce lung fibroblast proliferation. Moreover, TGF-\(\beta\) can signal via Smad-independent pathways through c-AbI, and imatinib has been shown to suppress TGF-\(\beta\)-downstream signaling via inhibition of c-AbI.\(^{63}\) It is clear, however, that when TGF-\(\beta\) signaling is down-regulated via canonical pSmad-dependent\(^{47}\) or noncanonical Smad-independent pathways during early alveolarization,\(^{57}\) secondary septal morphogenesis is perturbed. It is established that TGF-\(\beta\) signaling is exquisitely regulated at both the extracellular and intracellular levels, and that TGF-\(\beta\) downstream effectors are not only compartment specific but also temporally regulated within the epithelial, mesenchymal, and immune components of the lung. Our multilayered data support the premise that initial suppression of TGF-\(\beta\) ligand signaling via intracellular Smads, as well as loss of key downstream TGF-\(\beta\) effectors (eg, TGFBI), underlie murine early lung alveolar septation defects. Although beyond the scope of our studies because of the lack of a suitable myofibroblast-restricted driver and inability to specifically test TGF-\(\beta2\) function, our results suggest that supplementation of TGF-\(\beta\) (or more likely a downstream effector like TGFBI) from birth to P7 may possibly amend initial hyperoxic lung deficits. Thus, therapeutic manipulation of TGF-\(\beta\) signaling to ameliorate or prevent BPD in patients remains a tempting, but extremely challenging, prospect.

**TGFBI Is Required for Proper Lung Elastin Morphogenesis and Normal Lung Function**

Given the structural phenotypes and expression profiling similarities resulting from neonatal hyperoxia and Tgfbi deficiency, in addition to the four FAS1 domains, TGFBI includes a tripeptide Arg-Gly-Asp (RGD) motif.\(^{64}\) More important, TGFBI has been shown to influence cell-cell and cell-substratum interactions, including an ability to bind integrins in lung fibroblasts and vascular smooth muscle cells, as well as laminins, fibronectin, type I/II/IV collagens, and elastin. These influence cell proliferation, adhesion, and migration.\(^{65–71}\) all of which are important for lung development. Elastic fibers are composite structures (containing both amorphous elastin and microfibrils), and generation of a functional fiber occurs in a well-defined temporal sequence during lung tissue development.\(^{72}\) Collagen-elastin scaffolds and the elastic fiber-microfibril interface are thought to influence elastic fiber assembly. Previous work in mechanically ventilated mouse pups and preterm lambs has implicated elastin scaffold assembly as a critical component of alveolar septation.\(^{73,74}\) Although it is not yet known whether TGFBI plays a scaffold function during alveolarization, its coassociation with elastin microfibrils and collagens,\(^{68}\) its modulation of both cell-substratum and cell-cell interactions,\(^{70,71,75}\) and its spatial localization at the septal tips make this a likely role. The fact that we did not find any alteration of elastin mRNA levels in Tgfbi nulls, but detected a lack of elastin fibrils at the septal tips, suggests that an absence of TGFBI most likely compromises microfibril formation and normal deposition. Moreover, because TGFBI shares significant homology with periostin, which plays a scaffolding function during cardiovascular and skin tissue morphogenesis,\(^{28,76}\) Tgfbi deficiency could result in a failure of accessory proteins for elastin fiber assembly localizing to nascent microfibrils. Finally, the decreased elastance we observed in lungs of adult Tgfbi-null mice suggests that TGFBI may be important for proper lung elastin development and function.

**TGFBI Nulls Exhibit Additional TGF-\(\beta\)–Pathway Gene Alterations Related to Lung Fibroblast Maldevelopment**

Significantly, we saw up-regulation of FST gene and protein expression coincident with reduced expression of TGFBI, both secondary to hyperoxia and within Tgfbi nulls (Figures 1 and 6). FST contains a heparin-binding
ternary complexes with collagen VI.78 Although a direct proteoglycans, biglycan and decorin, in the ECM to form of lung development and results in long-standing de-in alveolar lung structure and function.5,81,82 Attempts at Drs. Wei Li and Weinian Shou (Indiana University School Transgenic and Knockout Mouse Core for assistance and Electron Microscopy Research Core and Indiana University We thank both the Indiana University School of Medicine) for help with targeting vector design and generation.

References

Conclusions
Most infants born extremely preterm will develop BPD because of inhibited saccular and alveolar lung development.7,9,10 Although relatively brief, lung injury incurred during the initial hospitalization affects a critical window of lung development and results in long-standing deficits in alveolar lung structure and function.5,81,82 Attempts at preventing BPD have been largely unsuccessful, in part, because of an inadequate understanding of BPD pathogenesis. Identifying novel effectors of saccular and early alveolar development, as well as carefully understanding the developmentally appropriate context of aberrant signaling pathways after preterm birth, is critical to design therapies that promote normal alveolar lung development and function.

Acknowledgments
We thank both the Indiana University School of Medicine Electron Microscopy Research Core and Indiana University Transgenic and Knockout Mouse Core for assistance and Drs. Wei Li and Weinian Shou (Indiana University School of Medicine) for help with targeting vector design and generation.
Ahlfeld et al


32. Simmons O, Bolanis EM, Wang J, Conway SJ: In situ hybridization (both radioactive and nonradioactive) and spatiotemporal gene expression analysis. Methods Mol Biol 2014, 1194:225—244


63. Reinboth B, Joshi A, Håkansson E, Gibson MA: Beta ig-h3 interacts directly with biglycan and decorin, promotes collagen VI aggregation, and participates in ternary complexing with these macromolecules. J Biol Chem 2006, 281:7816–7824