JNK suppresses pulmonary fibroblast elastogenesis during alveolar development

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Abstract

Background: The formation of discrete elastin bands at the tips of secondary alveolar septa is important for normal alveolar development, but the mechanisms regulating the lung elastogenic program are incompletely understood. JNK suppress elastin synthesis in the aorta and is important in a host of developmental processes. We sought to determine whether JNK suppresses pulmonary fibroblast elastogenesis during lung development.

Methods: Alveolar size, elastin content, and mRNA of elastin-associated genes were quantitated in wild type and JNK-deficient mouse lungs, and expression profiles were validated in primary lung fibroblasts. Tropoelastin protein was quantitated by Western blot. Changes in lung JNK activity throughout development were quantitated, and pJNK was localized by confocal imaging and lineage tracing.

Results: By morphometry, alveolar diameters were increased by 7% and lung elastin content increased 2-fold in JNK-deficient mouse lungs compared to wild type. By Western blot, tropoelastin protein was increased 5-fold in JNK-deficient lungs. Postnatal day 14 (PND14) lung JNK activity was 11-fold higher and pJNK:JNK ratio 6-fold higher compared to PN 8 week lung. Lung tropoelastin, emilin-1, fibrillin-1, fibrillin-5, and lysyl oxidase mRNAs inversely correlated with lung JNK activity during alveolar development. Phosphorylated JNK localized to pulmonary lipofibroblasts. PND14 JNK-deficient mouse lungs contained 7-fold more tropoelastin, 2,000-fold more emilin-1, 800-fold more fibrillin-1, and 60-fold more fibulin-5 than PND14 wild type lungs. Primarily lung fibroblasts from wild type and JNK-deficient mice showed similar differences in elastogenic mRNAs.

Conclusions: JNK suppresses fibroblast elastogenesis during the alveolar stage of lung development.

Keywords: Lung development, Elastin, c-terminal Jun kinase, Rho kinase

Background

Beginning at approximately 35 weeks gestation and continuing for several years postnatally [1], large saccular air spaces in the lung are progressively divided into smaller ones by secondary alveolar septa leading to a twenty-fold increase in alveolar surface area [2]. This exponential increase in gas-exchange capacity is essential to meeting the bioenergetic needs of ex utero growth and development. Formation of a low-compliance elastin band at the leading edge of a growing secondary alveolar septal ridge likely plays critical mechano-developmental role in alveolar septation [3-6].

Elastin is necessary for structure and function of both vascular tissues and epithelial-containing organs. It is critical to the hysteretic properties of both arteries [7] and lung [8] and to the elasticity of other epithelial organs such as skin and intestine. In infants with impaired alveolar development [9] and in animal models of the same [10-12], elastin fiber density and organization is disrupted. Regulators of elastogenesis (formation of elastin fibers) such as transforming growth factor-β [13], fibroblast growth factor-2 [14], epidermal growth factor [15], and insulin like growth factor-1 [16] are well described. With regards to signaling in the fibroblast, pro-elastogenic cytokines generally activate SMAD-signaling pathways and anti-elastogenic activate mitogen-activated protein kinases 1&3 (erk 2&1) [17]. The spatial-temporal regulation of pro- and anti-elastogenic signals is likely of critical importance since the formation of discrete elastin bands is critical to both alveolar and vascular development, and elastin experiences virtually no turnover with a half live of 74 years [18] despite expression of pro-elastogenic cytokines.
throughout life [19]. In the arterial vasculature, JNK signaling has been shown to suppress elastogenesis [20], but no such role has been demonstrated in the lung. We hypothesized that activation of JNK suppresses elastogenesis during alveolar lung development contributing to the localization of elastin to alveolar septal tips. To test this hypothesis, we assessed JNK activation during lung development, localized phosphorylated JNK by confocal microscopy, quantitated alveolar development and elastin localization in JNK deficient mice, and quantitated mRNA levels of elastin-associated genes in JNK deficient mouse lungs and primary fibroblasts.

Methods

Animal use

C57BL/6J wild-type, B6.Mapk8tm1Flv/J (JNK1−/−), B6. Twiss2tm1.1(cre)Dor (Dermo-1-cre), and B6.Gt(Rosa) 26Sortm4(ACTB-tdTomato, EGFP)Luo/j (tomato) mice were obtained from Jackson Laboratories. Previously described mice with a floxed JNK1 allele (Mapk8tm2Flv/J, referred to as JNK1LoxP/LoxP) on a B6.Mapk9tm1FlvMapk10tm1Flv/J background (hereafter referred to as JNK2&3−/−) were utilized with permission from Roger Davis at the University of Massachusetts [21]. Since JNK3 is expressed only in the heart, testis and brain [22], single JNK2 or JNK3 knockout animals were not utilized. All animals were housed in a barrier facility with purified air and provided purified water and autoclaved food ad libitum. Animal use was approved by the Cincinnati Children’s Hospital Medical Center Animal Use and Care Committee.

To localize lung JNK activity, we utilized a mesenchymal cell lineage tracing strategy with immunofluorescent staining of pJNK. Dermo1 (a.k.a. twist2) is mesenchymal transcription factor expressed as early as E15.5 [23]. The “tomato” mouse constitutively expresses a membrane-bound td-tomato except in the presence of cre in which recombination excises td-tomato and eGFP is expressed. Crossing Dermo1-cre mice with “tomato” mice lineage traces lung mesenchymal cells with eGFP.

Genotyping

Mice were genotyped using the primers listed in Table 1.

Lung tissue processing and analysis

Tissue procurement and processing

Mice were sacrificed by intraperitoneal injection of ketamine, xylazine, and acepromazine (100, 6, and 2 mg/kg respectively) and severing of the left renal artery. After exsanguination, the trachea was cannulated with a 22 gauge blunt tip needle and lungs isovolumetrically inflated with 4% paraformaldehyde in PBS at a pressure of 25 cm H2O. Lung inflation was maintained by securing a silk ligature around the trachea, and then the chest was opened. The lungs were removed and then fixed overnight at 4°C. After fixation, the lung lobes were removed from the bronchi and dehydrated by serial passage into 70% ethanol and paraffinized. Five μm sections obtained at random angles through all five lobes.

Lung elastin staining and quantification

Lung sections were stained for elastin using the Hart’s Staining method (PolyScientific) and elastin quantitated by previously published methods [24]. Using a Zeiss Axio Imager A.2 three 20X images were obtained from each lobe yielding fifteen images per mouse. Using MatLab (MatWorks) these images were then separated into four components by cluster analysis after defining a color spectrum and intensity values for dense elastin (arterial walls and elastin bands at septal tips), diffuse elastin (elastin in alveolar walls and bases), non-elastin tissue, and airspace. Pixel quantitation for each component was performed for each image and average values per animal used for statistical analysis. For quantitation, the fraction of each component of total tissue (dense elastin + diffuse elastin + non-elastin tissue) was utilized.

Morphometry

Mean alveolar diameter was calculated utilizing a published morphometric MatLab program [8,25-28], and fractional airspace was calculated using previously described methods [29]. By defining secondary alveolar crests as invaginations into distal lung airspace by tissues associated with an

Table 1 Genotyping primers used

| Allele    | Common forward primer | Reverse primer                        |
|-----------|-----------------------|---------------------------------------|
| JNK1-null | CCAGGCTCCTCCTCATTTCA  | Wild Type-CAGGCTCATTCCCTCCTCATAG      |
|           |                       | Mutant-TCACCACATAAGCGGTATGC            |
| JNK2-null | GTTAGACAATCCACAGGTTGT  | Wild Type-CCAGCTCATTCCCTCCTCATAG      |
|           | GTGTTGAGGAACTGGGCTA   | Mutant-GGAGCCGGATAGATGAGTTACC         |
| JNK3-null | CGTAATCTGTACAGAAATCCCATA | Wild Type-CCAGACTGCTTGGGAAAA         |
|           |                       | Mutant-CTCTGCTTTCGAAAGCAACCCTT        |
| JNK1-floxed | GGATTATGCCCCTGTCCTGT | GAACACTGTGTCCTCAATTTCCATCC           |
elastin bundle, numbers of secondary alveolar septal crests were counted per 20X field.

**Immunofluorescence**

*Dermo1-cre-tomato* mice were sacrificed at PND5 and the lungs immunostained for pJNK using an AF647 secondary antibody. Confocal images were obtained using a Nikon AR1si inverted microscope.

**Tissue culture**

**Pulmonary fibroblast isolation**

Pulmonary fibroblasts were isolated from PND14 mice by previously described methods which yield >85% fibroblast purity [30,31]. Briefly, PND14 mice were sacrificed and their lungs inflated by gently instilling dispase (BD Biosciences) via the trachea and then plugging the trachea with 1% low melting point agarose. The lungs were incubated in dispase at 25°C for 45 minutes and lung tissue teased from bronchi and large vessels using sterile forceps. Dispase was neutralized using DMEM with 10% FBS and the fibroblasts were allowed to adhere in a 100 cm² tissue culture dish for 1 hour at 37°C and the plate rinsed with PBS. The fibroblasts were allowed to grow in DMEM with 10% FBS and 1% Penicillin/Streptomycin and used at passage 3.

**In vitro Cre-recombinase experiments**

*JNK2*&*3−/−* (which also has floxed *JNK1* alleles) pulmonary fibroblasts were infected at 50% confluence with 10⁶ plaque forming units of replication-deficient adenovirus expressing GFP or Cre (Vector BioLabs). Efficacy was assessed by Taqman qPCR for *JNK1* mRNA (Applied Biosystems, proprietary primers). RNA and cell culture media was collected at 48 hours.

**JNK activity, protein, RNA quantification**

**JNK activity assays**

For *in vivo* experiments, lung homogenate c-Jun phosphorylation was quantitated to measure JNK activity. To do so, JNK was immunoprecipitated from lung homogenates using an isofrom nonspecific JNK antibody (Santa Cruz) conjugated to agarose beads. Precipitated JNK was incubated with GST-labeled c-Jun (Michal Karin, University of California at San Diego) [32] and P-32 adenosine triphosphate, the kinase solution was separated on a polyacrylamide gel, transferred to a PVDF membrane, and radiolabeled c-Jun quantitated using a Storm 860 phosphorimager.

**Western blot**

Lungs from mice of at least two different litters were snap frozen and homogenized in RIPA buffer with protease inhibitor cocktail (Sigma) using a Qiagen TissueLyser II. Protein content was determined and the homogenates were electrophoretically separated and transferred to PVDF membrane. When not frozen, lysates were kept on ice until electrophoresis. Western blot for pJNK and JNK (both isofrom non-specific, 1:200 dilution, Santa Cruz) and tropoelastin (1:2000 dilution, Elastin Products Company) was performed and chemiluminescence detected using a General Electric LAS3000.

**PCR**

Lungs from mice aged E15.5 to 8 weeks were snap frozen and then homogenized in RLT buffer using a Qiagen TissueLyser II. For primary lung fibroblasts, passage two cells were seeded at 50% density and collected at 48 hours (with cells achieving 100% density). Tropoelastin, emilin-1, fibrillin-1, lysyl oxidase, fibrulin-5, surfactant protein B, CD31, and GAPDH mRNA was quantitated by Taqman PCR (Applied Biosystems proprietary primers).

**Soluble elastin quantification**

Pulmonary fibroblasts were cultured for 48-hours in DMEM, 10% FBS, 1% penicillin/streptomycin and the media analyzed for elastin content using the Fastin kit (Biocolor, UK) per manufacturer instructions.

**Statistical analysis**

Statistical comparisons between groups were performed using the Student two-tail t-test or one-way ANOVA using the Holm-Sidak method. p-values of <0.05 were considered significant.

**Results**

**JNK-deficient mice have enlargement of distal lung airspaces**

The lungs of PND14 *JNK1−/−* and *JNK2&3−/−* mice appeared to have larger airspaces with fewer alveoli as compared to wild type mice (Figure 1A-C). Mean alveolar diameter was increased by 2% in *JNK1−/−* mouse lung and by 7% in *JNK2&3−/−* lungs (Figure 1D.) Fractional airspace was increased by 14% in *JNK1−/−* mouse lung and by 19% in *JNK2&3−/−* lungs (Figure 1E). Both *JNK1−/−* and *JNK2&3−/−* mice had 20% fewer alveolar crests than wild type mice (Figure 1F). The enlarged distal airspaces and reduced numbers of secondary alveolar septal crests in JNK-deficient mice support a role for JNK in alveolar septation.

**JNK reduces lung elastin content**

To test whether JNK suppresses lung elastogenesis, elastin was morphometrically quantitated in PND14 wild type, *JNK1−/−* and *JNK2&3−/−* elastin-stained lung sections. Grossly, lung elastin content was increased in *JNK1−/−* and *JNK2&3−/−* lungs compared to wild type (Figure 2A-C). As described in methods, we quantitated the number of dense elastin, diffuse elastin, tissue, and...
airspace pixels in elastin-stained lung sections from at least three mice from two to three different litters (Figure 2D-G). Wild type lungs contained 7% more tissue than JNK-deficient lungs (not significant). Normalized to total tissue, JNK1−/− and JNK2&3−/− lungs had twice the amount of total elastin as wild type lungs (Figure 2H). While JNK1−/− lungs had a non-significant increase in dense elastin to total elastin ratio, JNK2&3−/− lungs had a 1.5-fold increase compared to wild type (Figure 2I). Both JNK1−/− and JNK2&3−/− lungs had 1.7 fold more diffuse elastin than wild type lungs (Figure 2J). By Western blot of PND14 lung homogenates, JNK1−/− and JNK2&3−/− lung contained 5- and 15-fold more tropoelastin than wild type lungs (Figure 2I). By Western blot of PND14 lung homogenates, JNK1−/− and JNK2&3−/− lung contained 5- and 15-fold more tropoelastin than wild type lungs homogenates respectively (Figure 2K). These data support the hypothesis that JNK suppresses elastogenesis during lung development and plays a role in the localization of elastin fibers to alveolar septal tip.

JNK activity Is increased during postnatal lung development

To determine whether JNK activity is dynamic over the course of lung development, we immunoprecipitated JNK from lung homogenates and quantitated conversion of c-Jun to phospho-c-Jun. Compared to PN 8 week lung, JNK activity was increased during the saccular and alveolar stages of lung development with the highest JNK activity in PND14 lung homogenates (Figure 3A). When confirmed in triplicate, JNK activity at PND14 was 11-fold higher than JNK activity at PN 8 weeks (Figure 3B). By Western blot, phosphorylation of the p54 isoform of JNK was increased in PND14 lung compared to PN 8 week without phosphorylation of the p46 isoform. There were no differences in total JNK protein content (Figure 3C-D). Among the molecules required for elastin fibril assembly are tropoelastin, lysyl oxidase, emilin-1, fibrillin-1, and fibulin-5. To identify an association between JNK activity and elastogenesis, mRNAs of these elastin-associated genes were quantitated at time points between E15.5 and postnatal week eight. The mRNAs of tropoelastin, lysyl oxidase, emilin-1, fibrillin-1, and fibulin-5 were all decreased during late alveolar development (PND14) compared to earlier time points (Figure 3E-F). These data demonstrate an association between decreased transcription of elastin-associated molecules and increased lung JNK activity during later alveolar development.

JNK activity is localized to pulmonary lipofibroblasts during alveolar development

We utilized both lineage tracing and immunofluorescent co-localization techniques to determine which cell type was experiencing JNK activation. We first narrowed the range of possible cell types by staining for pJNK in PND5 mesenchymal cell lineage traced lung sections. pJNK was present in a subset of mesenchymal cells and
was not activated in epithelial or hematopoietic cells (Figure 4A-E). Since pulmonary fibroblasts, endothelial cells, and pericytes are all derived from the lung mesenchyme, we performed immunostaining for desmin which is expressed in fibroblasts but not the other mesenchymal cell types. pJNK was present only desmin-positive cells (Figure 4F). Pulmonary fibroblasts can grossly be separated into the more fibrogenic myofibroblasts and less fibrogenic lipofibroblasts. Myofibroblasts express higher levels of α-smooth muscle actin (α-SMA). pJNK was not located within α-SMA positive cells (Figure 4G). Lipofibroblasts contain increased numbers of lipid droplets, produce less tropoelastin [31,33], and are typically located in alveolar corners [34]. In oil-red-o stained sections, pJNK localized to lipofibroblasts and not to oil-red-o positive macrophages in airways (Figure 4H). These data demonstrate that JNK activity during later alveolar development is restricted to pulmonary lipofibroblasts.

JNK suppresses pulmonary fibroblast elastogenesis

Since JNK suppresses elastogenesis in the aorta [20] and other MAP kinases regulate pulmonary fibroblast elastin synthesis [17], we tested whether JNK suppresses pulmonary fibroblast elastogenesis. Primary PND14 lung fibroblasts were cultured from wild-type, JNK1−/−, and JNK2&3−/− mice. JNK1−/− and JNK2&3−/− lung fibroblasts contained 8,000-fold more tropoelastin mRNA than wild type (Figure 5A), and the media from JNK1−/− and JNK2&3−/− lung fibroblast cultures contained 50% more soluble tropoelastin (Figure 5B). All three cell types were 50% confluent at seeding and 100% confluent at collection. To test whether JNK1 and JNK2 independently suppressed pulmonary fibroblast elastogenesis, JNK1 alleles were deleted from JNK2&3−/− lung fibroblasts using a cre-expressing adenovirus (Ad-Cre). JNK1 mRNA was reduced 35-fold in Ad-Cre infected fibroblasts compared to control (Ad-GFP) (Figure 5C) and >90% of Ad-GFP exposed fibroblasts expressed GFP. Deletion of JNK1...
increased tropoelastin mRNA three-fold (Figure 5D). To test whether JNK activation suppresses pulmonary fibroblast elastogenesis, we used the JNK activator anisomycin. Anisomycin reduced PND14 wild-type fibroblast tropoelastin mRNA in a dose-dependent manner that was not statistically significant (Figure 5E). Since epithelial or endothelial cell contamination of primary fibroblast cultures could account for the observed
differences, we quantitated surfactant protein B and CD31 (Pecam1) mRNA content. Neither marker could be detected in wild type fibroblast culture but the markers were abundant in wild type lung demonstrating that fibroblasts cultures did not have significant epithelial or endothelial cell contamination. Since JNK3 is not expressed in lung [22], these in vitro data demonstrate that JNK1 and JNK2 independently suppress pulmonary fibroblast elastogenesis.

**JNK suppresses mRNAs of elastin-associated genes**

Since the mRNAs of many elastin-associated molecules decreased in association with increased lung JNK activity, and since JNK suppressed elastogenesis *in vitro*, we tested whether JNK-deficient lungs had increased mRNA levels of elastin-associated molecules. JNK1−/− and JNK2&3−/− lungs contained 2,000-fold more emilin-1 (Figure 6A) and 700-fold more fibrillin-1 (Figure 6B) than wild type lungs. JNK1−/− lungs contained 10-fold and JNK2&3−/− 60-fold more fibulin-5 (Figure 6C), but differences in quantities of lysyl oxidase mRNA were small (Figure 6D). JNK1−/− and JNK2&3−/− lung contained 6–8 fold more tropoelastin mRNA than wild type (Figure 6E). To confirm that JNK-deficiency itself and not a secondary cell signaling alteration (such as altered mechanical strain or differences in matrix stiffness) was
Figure 5 JNK suppresses pulmonary fibroblast elastogenesis in vitro. (A) JNK1−/− and JNK2&3−/− PND14 lung fibroblasts contained more tropoelastin mRNA than wild type lung fibroblasts. (B) The media from JNK-deficient lung fibroblasts trended towards having more soluble tropoelastin than media from wild type lung fibroblasts. (C) Infection of PND14 JNK2&3−/− lung fibroblasts (which also contained floxed JNK1 alleles) with cre-expressing adenovirus (Ad-Cre) was effective at reducing JNK1 mRNA levels when compared to fibroblasts infected with a green fluorescent protein adenovirus (Ad-GFP). (D) Deletion of all three JNK isoforms from pulmonary fibroblasts resulted in increased tropoelastin mRNA compared to deletion of only JNK2&3. E The JNK activator anisomycin reduced wild type PND14 lung fibroblast tropoelastin mRNA in a dose-dependent manner. *p < 0.05, **p < 0.01, ***p < 0.001 by one-way ANOVA. All analyses were performed in triplicate.
responsible for these differences in elastin-related gene mRNA content, we quantitated these same mRNAs in primary lung fibroblasts. Neither emilin-1 nor fibulin-5 mRNA was detected in wild-type lung fibroblasts but it was present in both JNK1−/− and JNK2−/− lung fibroblasts (Figure 6F&H). Fibrillin-1 mRNA was detected in wild type lung fibroblasts, but mRNA content was several thousand-fold higher in JNK-deficient lung fibroblasts (Figure 6G). Lysyl oxidase mRNA was not statistically different between wild type and JNK-deficient lung fibroblasts (Figure 6I). Tropoelastin mRNA was assessed previously. A schematic depicting how emilin-1, fibrillin-1, fibulin-5, lysyl oxidase, and tropoelastin interact to form elastin fibers is shown in Figure 6J. These data demonstrate that JNK regulates many of the genes involved in elastin microfibril formation and support the concept that JNK suppresses the pulmonary fibroblast elastogenic program.

Discussion

For the first time, we have demonstrated a role for JNK in suppressing the elastogenic program during the alveolar stage of lung development. JNK activity increased during late alveolar development, and this increased JNK activity was negatively correlated with elastogenic gene mRNAs. JNK-deficiency increased lung elastin content and impaired alveolar septation. Both JNK-deficient lung and lung fibroblasts had increased mRNA levels of elastin-associated genes. Deletion of both JNK1 and JNK2 increased tropoelastin mRNA to a greater extent than either individually demonstrating that both isoforms independently regulate elastogenesis. Our observations are consistent with previous reports demonstrating JNK regulation of elastogenesis in the aorta [20] and AP-1 mediated suppression of tropoelastin expression [35]. Phosphorylated c-Jun, the readout of the JNK luciferase assay, is a component of the AP-1 transcriptional complex.

We utilized two methods to assess JNK activation during lung development. In addition to comparison of total JNK to phosphorylated JNK, we utilized a pull-down technique developed and validated at the University of California at San Diego in a variety of cell lines with increasing exposures to ultraviolet irradiation—a well described activator of JNK signaling [32]. This technique quantitates JNK phosphorylation of c-Jun. A drawback of this assay is the absence of a loading control which likely accounts for the increased variability in this assay compared to Western blot. Nonetheless, by both Western blot and JNK pull-down, we demonstrated a negative association between lung JNK activity and elastogenic mRNAs.

We have provided strong in vitro and in vivo loss of function data to support a role for JNK in regulating the elastogenic program in the lung. When assessed in isolation, these effects are clear and strong. However, multiple dynamic processes regulate elastogenesis which likely accounts for larger in vitro than in vivo effects. JNK activity was reduced in adult lung compared to developing lung. This discrepancy is likely due to reduced transcriptional activation of the elastogenic program in the adult compared to the juvenile. Conceptually, JNK activity may serve as a brake slowing elastogenesis during alveolarization which turns off when the elastogenic program is no longer activated. The localization of pJNK
to pulmonary lipofibroblasts is consistent with their reduced elastogenic and fibrogenic potential [31]; although, the fact that lipofibroblasts increase in number at an earlier age than JNK activity levels increase [36] makes a direct relationship between the two unlikely. The mechanism of how JNK suppresses elastogenesis needs to be determined in future studies.

The necessity of elastin band formation for secondary alveolar septation is well-established [4,16]. Both elastin [9] and the lysyl oxidase [37] content are increased in infants with chronic lung disease of prematurity. In mice, elastin haploinsufficiency leads to emphysematous changes [6], and Mamamoto et al. recently demonstrated that reduced matrix stiffness also impairs alveolarization [38]. Our data and these previously published data demonstrate that alveolar growth requires both a low compliance elastin band and high compliance alveolar walls. Therapies aimed at restoring alveolar growth should seek to normalize this relationship.

In conclusion, we report that JNK suppresses pulmonary fibroblast elastogenesis during the alveolar stage of lung development plays a role in alveolar septation.

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions
SL aided in project conceptualization, designed experiments, and performed experiments. HP designed morphometric programs and optimized programs for elastin quantitation. SY performed morphometry and other experiments. BV conceptualized the project, designed experiments, and analyzed data.

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