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## **Characterization of the mid-foregut transcriptome identifies genes regulated during lung bud induction**

**Guetchyn Millien**1, **Jennifer Beane**1,4, **Marc Lenburg**3, **Po-Nien Tsao**1, **Jining Lu**1, **Avrum Spira**1,2,4, and **Maria I. Ramirez.**1,2,\*

1 *Pulmonary Center, Department of Medicine, Boston University School of Medicine, Boston, MA 02118*

2 *Department of Pathology, and Laboratory Medicine, Boston University School of Medicine, Boston, MA 02118*

3 *Depatment of Genetics and Genomics, Boston University School of Medicine, Boston, MA 02118*

4 *Bioinformatics Program, Boston University College of Engineering, Boston, MA 02118*

### **Abstract**

To identify genes expressed during initiation of lung organogenesis, we generated transcriptional profiles of the prospective lung region of the mouse foregut (mid-foregut) microdissected from embryos at three developmental stages between embryonic day 8.5 (E8.5) and E9.5. This period spans from lung specification of foregut cells to the emergence of the primary lung buds. We identified a number of known and novel genes that are temporally regulated as the lung bud forms. Genes that regulate transcription, including DNA binding factors, co-factors, and chromatin remodeling genes, are the main functional groups that change during lung bud formation. Members of key developmental transcription and growth factor families, not previously described to participate in lung organogenesis, are expressed in the mid-foregut during lung bud induction. These studies also show early expression in the mid-foregut of genes that participate in later stages of lung development. This characterization of the mid-foregut transcriptome provides new insights into molecular events leading to lung organogenesis.

### **Keywords**

Lung; development; organogenesis; foregut; endoderm; embryo; mouse; microarray; RNA amplification; gene expression; real time PCR; laser capture microdissection; transcription factors; chromatin remodeling; Fox; Notch

> Between E7.5 and E9.5 of mouse development, remarkable morphogenetic changes take place in the ventral foregut resulting in formation of distinct organs including pancreas, liver, thyroid, and lung (Grapin-Botton and Melton, 2000; Wells and Melton, 1999). Organ-specific genes such as Pdx1 (pancreas) (Murtaugh and Melton, 2003), albumin (liver) (Jung et al., 1999), Hhex and Pax8 (thyroid) (Parlato et al., 2004) and thyroid transcription factor 1 (thyroid and lung) (DeFelice et al., 2003; Desai et al., 2004; Kimura et al., 1999) demarcate different regions

<sup>\*</sup>Corresponding author. Boston University School of Medicine., Evans Biomedical Research Center, 650 Albany St., X-440, Boston, MA 02118., E-mail address: mramirez@bu.edu, Fax: +1-617-638-7530.

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of the foregut as development progresses. By E9 of development, embryos containing between 16 and 20 somites already show thyroid, liver and pancreas primordia but the mid-foregut region, between the pharyngeal arches and the liver and pancreas buds, is still a tube that shows no sign of lung bud formation. In this environment, the trachea and the lung evaginate from the ventral and ventrolateral foregut respectively at approximately E9.5 (Cardoso and Lu, 2006; Warburton et al., 2005). These developmental events are regulated by fibroblast growth factors, bone morphogenetic proteins, retinoic acid, and sonic hedgehog, and their receptors among other factors (Bellusci et al., 1997; Desai et al., 2004; Jung et al., 1999; Litingtung et al., 1998; Rossi et al., 2001; Sakiyama et al., 2003; Sekine et al., 1999; Warburton et al., 2005; Weaver et al., 2000). These signaling networks activate downstream effectors in both foregut mesoderm and endoderm to induce organ specific gene expression (Cleaver and Krieg, 2001; Horb, 2000; Jung et al., 1999; Kumar et al., 2003; Matsumoto et al., 2001; Wells and Melton, 2000).

Compared to the detailed understanding of the programs leading to formation of the liver and pancreas (Bort et al., 2006; Lee et al., 2005; Lemaigre and Zaret, 2004; Murtaugh and Melton, 2003; Serls et al., 2005; Tremblay and Zaret, 2005), the lung morphogenetic program has not been explored in detail, in large part due to the paucity of marker molecules known to be expressed in the presumptive lung region of the foregut. Microarray studies have been used to characterize profiles of gene expression in embryonic tissues including preimplantation mouse embryos (Sherwood et al., 2007; Zeng et al., 2004), pre-pancreas and early pancreatic endodermal cells (Gu et al., 2004). The latter database has been particularly important for the identification of genes involved in acquisition of pancreatic cell fate and has also established the transcriptional profiles of endodermal precursors. To date, global profiles of gene expression in the developing lung have been focused on the processes of branching morphogenesis and on perinatal lung development (Banerjee et al., 2004; Bonner et al., 2003; Lu et al., 2005; Lu et al., 2004a), but there are not yet similar data on molecular changes that accompany the initiation of lung organogenesis.

We report herein the characterization of the transcriptome of the lung region of the foregut (referred as mid-foregut) prior and during initiation of lung organogenesis, including trachea and primary bud formation. We have identified genes expressed in mid-foregut cells, and genes whose expression levels change during lung primordium formation.

### **1. RESULTS AND DISCUSSION**

### **1.1 Developmental lung genes are expressed in the prospective lung region of the foregut**

We studied temporal differences in gene expression in mid-foregut tissues containing endoderm and mesoderm cells isolated by microdissection at three developmental stages prior and after lung bud formation (16–20, 21–25 and 26–30 somite stages) (Figure 1A, B). We determined by real time RT-PCR (QRT-PCR) whether the tissues to be used in the microarray studies expressed lung and liver marker genes. Twenty four hours before lung budding, at E8.5, the mid-foregut already expresses lung developmental genes. Thyroid transcription factor 1 (Titf-1, TTF-1, Nkx2.1), a transcription factor critical for lung cell differentiation (Kimura et al., 1999), was detected in the mid-foregut tissue at all three developmental stages, and the level of expression was increased as the primary lung buds formed (Figure 1C); although Titf-1 is also expressed during thyroid organogenesis, the thyroid was excluded from the mid-foregut samples analyzed, as shown in Figure 1A–B. Fibroblast growth factor 10 (FGF10) expressed in the foregut mesenchyme can be detected at the three developmental stages, and significantly increases in 26–30 somite vs. 21–25 somite foreguts (Figure 1C). Surfactant protein C (SP-C) mRNA, a lung-specific gene (Khoor et al., 1994; Wang et al., 1994), expressed very early in lung development and a downstream target of Titf-1, was not detected in 16–20, 21–25 or 26– 30 somite embryos by QRT-PCR (data not shown) but can be detected later in the mid-foregut

endoderm of embryos containing more than 30 somites (Figure S1A–B, supplementary material). Albumin mRNA was not detected in the mid-foregut by real time PCR (data not shown), but is highly detected in the posterior foregut endoderm (liver), isolated by laser capture microdissection from embryos containing more than 30 somites (Figure S1A–B, supplementary material). Microarray analyses of amplified mid-foregut RNA show expression of the lung expressed genes T1α and caveolin-1 (Ramirez et al., 2002; Williams et al., 1996), and marked reduction of the liver genes  $\alpha$ -fetoprotein and transthyretin (Jung et al., 1999; Lee et al., 2005), as development proceeds (Table S3, supplementary material). Pax8, a thyroid gene expressed also in the branchial arches (Trueba et al., 2005) was detected at low levels in the 16–20 somite samples, and its level remains low and unchanged during lung bud formation (Table S3, supplementary material). Overall, these data confirm the specific nature of the regions selected to study. The exact timing of lung cell specification in the multipotent foregut has not been clearly determined, but detection of Titf-1 and FGF10 in the mid-foregut at E8.5 indicates certain commitment of the cells to start lung formation (Desai et al., 2004).

### **1.2 Statistical analysis and functional classification of the microarray data**

Three independent mRNA samples, at each of the developmental stages selected (16–20s, 21– 25s, and 26–30s), were amplified and analyzed by microarrays (total of 9 arrays). Hierarchical clustering analysis of genes significantly up-regulated or down regulated across all nine samples, with ANOVA and Student's t-test  $p \le 0.05$  and detection  $p \le 0.05$  are shown in Figures 2A and 2B. The result illustrates that the general premise of this study is correct, i.e. that significant changes in gene expression accompany the induction of the lung as it emerges from the foregut at the 26–30 somite stage.

A number of comparisons are possible to analyze the data. We chose to identify differences between early and late foreguts as representing the absence and the presence of lung buds. Relative to the earliest group, 104 genes are up-regulated more than 1.8 fold and 119 genes down-regulated more than 1.8 fold in the late group undergoing lung organogenesis. Although a change higher than 2-fold has been conventionally used to consider genes as functionally important, we have opted to use 1.8 fold as cut-off level due to the high number of genes showing a statistically significant change between 1.8 and 2-fold. A number of previous studies also opted to use lower cut-off levels (1.2–1.5) when the changes are highly significant (Jeong et al., 2005; Lu et al., 2004b; McReynolds et al., 2005). Enriched biological themes within the up-regulated genes and the down-regulated genes were identified using appropriate computer algorithms (Figure 2C–D). The most significantly overrepresented, up-regulated biological process (Ease score=  $3.35 \times 10^{-3}$ ) was regulation of transcription (Figure 2C). That group contains largely, but not solely, transcription factors; many have not been previously shown to be related to lung development. Some of these transcription factors were further analyzed by QRT-PCR and whole mount in situ hybridization. Within the down-regulated genes, macromolecule/protein biosynthesis was the most significantly overrepresented biological process (Ease score=  $3.23 \times 10^{-9}$ ) (Figure 2D). Some transcriptional regulatory genes were also down-regulated. Genes categorized as functioning in proliferation, transport, and organization and biogenesis are represented in both the up- and down-regulated groups.

To find genes enriched in each of the three time points studied, especially in the intermediate time point (21–25 somite stage), a parametric ANOVA was performed. 1076 probe sets with a p-value less than 0.05 were selected for further analysis by K-means clustering (Figure 3). Five clusters (A–E) were generated in which genes decline from 16–20 to 21–25 somites or to 26–30 somites, increase from the 16–20 somite stage with a peak at 26–30 somites, or peak in the middle at the 21–25 stage, or increase from 21–25 to 26–30 somites. We listed in Table 1 genes that are enriched in each developmental time point identified by ANOVA and/or Student's t tests, and are involved in transcriptional regulation. These genes include factors

that bind directly to cis-elements in the DNA, co-factors, and genes that form chromatin modifying complexes. For a complete list of genes see Tables S4–S6 in supplementary material.

### **1.3 Real time RT-PCR confirmation of temporal changes in gene expression**

Forty three genes shown to change level of expression during lung induction were analyzed by QRT-PCR using non-amplified mid-foregut RNA. mRNA levels in 26–30 somite samples normalized to GAPDH mRNA levels were expressed relative to the 16–20 somite samples set at a relative value of one. A similar semi-quantitative comparison of the same genes was done for the microarray data. As shown in Figure 4A–B and Figure S3A–B, 19 out of 43 genes tested showed statistically significant changes in expression by QRT-PCR correlating well to the microarray data, although the absolute fold-differences do not match precisely as would be expected due the different nature of the methods and the analysis of different samples. In addition, the trend in expression patterns of other 12 genes (e.g. Cbfa2t1, Cbx4, Six1) was confirmed by QRT-PCR, although the fold change observed by this method was not statistically significant ( $p > 0.05$ ) (Table S7A). The remaining 12 genes tested did not match the microarray data (Table S7B).

### **1.4 Transcription factors and co-factors are developmentally regulated as the lung buds form**

The largest change demonstrated by QRT-PCR analysis is in the LIM domain transcription factor Isl1 (Thor et al., 1991). Isl1 is necessary for proliferation and survival of cells in the foregut mesoderm and dorsal pancreatic mesenchyme, and is linked to cell fate decision in motor neurons (Pfaff et al., 1996), cardiac cells (Cai et al., 2003), and pancreatic islets (Ahlgren et al., 1997) but has not been previously implicated in lung development. Isl1 null mice [Isl1 (−/−)] are developmentally arrested soon after E9.5, show abnormal organization of the vascular endothelium, and severely abnormal heart development with reduction of the amount of atrial tissue. Expression of FGF10, Bmp4 and Bmp7 is highly downregulated in pharyngeal endoderm and splanchnic mesoderm in the absence of Isl1 at E8.5–9.0, likely as a result of direct or indirect regulation of these growth factors by Isl1 and/or viability of the foregut cells. A recent publication indicates a putative cis-element for Isl1 in the FGF10 promoter (Ohuchi et al., 2005). We now show that Isl1 is transiently expressed in the early stages of lung morphogenesis and its expression is down regulated as the lung begins branching morphogenesis. Early lethality of the Isl1 null mutation precludes to study the role of Isl1 in lung budding.

Other transcription factors that directly bind to DNA validated by QRT-PCR include HoxA4 (Packer et al., 2000), NFIB (Chaudhry et al., 1997), and Foxf2 (Aitola et al., 2000). We have identified by microarrays expression in the mid-foregut region of other Fox genes that were not previously linked to the process of lung bud formation. Most of the Fox genes were detected at high levels and did not change over the time period studied. Fox transcription factors play important roles in development (Lee et al., 2005), and during induction of the endoderm derived organs (Carlsson and Mahlapuu, 2002). Foxa2 and Foxa1 are critical for initial steps in foregut tube closing and viability of endodermal cells (Ang and Rossant, 1994). They also regulate expression of several lung genes including surfactant protein genes (Costa et al., 2001). Other Fox factors, such as Foxf1, Foxp1, and Foxb1, are regulators of later events in lung development (Costa et al., 2001).

Changes in level of expression of the Dach2 co-factor, that belongs to the Eya/Six/Dach transcriptional complex involved in cell fate decisions in other organs (Davis et al., 2001), was validated by QRT-PCR. Three members of these complexes, sine-oculis 1 (Six1) and -5 (Six5) and Daschund 2 (Dach2) change their level of expression as the lung buds; Six1 is down regulated, and Six5 and Dach 2 are up-regulated. Eya/Six/Dach complexes can switch between

gene activation and repression of gene expression depending on the members of the family that form these complexes.

Four additional genes showing a large increase in 26–30 somite samples compared to 16–20 somite samples in the microarray analysis were measured by QRT-PCR of non-amplified mRNA and confirmed the original findings (Figure S3A–B, supplementary material). The putative functions of these molecules are diverse and include the enzymes alpha-2,8 sialyltransferase 8A (Yoshida et al., 1995) and 3-oxoacid CoA transferase 1 (Ganapathi et al., 1987), the apoptosis-related gene caspase 7 (Lakhani et al., 2006), and syndecan 2 (David et al., 1993) among others.

### **1.5 Expression of selected transcription factor and signaling gene families during lung bud formation**

We evaluated microarray data for expression of members of the Fox, Hox, Tbx, GATA transcription factor families (Table 2) and genes of selected signaling pathways that participate in lung development including FGFs, Bmps, Shh, Retinoic Acid, and Notch pathways (Table 3) (Cardoso, 1995;Cardoso and Lu, 2006;Warburton et al., 2005). We have detected Foxa1, Foxa2, Hox A4, A5, B4, B5, B8; Tbx 1, 2; GATA 5, 6; FGF7, FGF1, FGFRII and IV; Bmp4, and BmpR1a; Notch1 and Dll3 mRNAs among others. QRT-PCR confirmed that HoxA4 and Foxf2 (Figure 4A–B) are significantly increased in 26–30 somite samples. We anticipated finding expression of FGF10 and Titf-1 (Nkx2.1) but these factors were not detected in the microarray analysis, possibly because they are of very low abundance, are poorly amplified prior to hybridization, and/or the oligos on the microarray are 5′ to the amplified message region as the amplification method is 3′bias (Table S1); however these mRNAs can be readily detected by QRT-PCR in non-amplified mid-foregut samples (Figure 1C). Microarray analysis of amplified or non-amplified samples can always produce false positive or negative results. Therefore, validation by other methods is necessary. In this study we have analyzed selected genes by QRT-PCR and whole mount in situ hybridization.

The representation of members of the Fox family is notable as shown in Table 2. Of the 26 Fox family members represented on the microarrays used in the study, 12 are present in the mid-foregut (detection  $p \le$  value 0.05). Foxa1 and Foxa2 display the highest detection levels. However, with the exception of Foxp1 (Shu et al., 2001), Foxm1 (Kim et al., 2005) and Foxf2 (Wang et al., 2003) as shown in Tables 1 and 2, mRNA expression levels of other Fox genes do not change significantly during initiation of the lung budding. Among the signaling pathways, it is notable the number of Notch pathway related genes present in the mid-foregut as the lung forms (24 present in the mid-foregut out off 114 probe sets, Table 3). Changes in Notch 1 (Taichman et al., 2002), Delta-like 3 (Ladi et al., 2005), and presenilin enhancer 2 (Francis et al., 2002) are statistical significant. As the Notch pathway has been shown to regulate early cell fate decisions in other tissues (Louvi and Artavanis-Tsakonas, 2006), its role in lung induction will entail further evaluation.

### **1.6 Chromatin remodeling and DNA methylation genes change their level of expression during initiation of lung organogenesis**

Among several chromatin remodeling genes identified in Table 1, smarce1/BAF57 and Ing3 were confirmed to increase their level of expression significantly as the lung buds. Smarce1/ BAF57 is one of the genes that form part of the ATP-dependent chromatin remodeling SWI/ SNF complex (Chen and Archer, 2005;Domingos et al., 2002) which modify nucleosomes changing the accessibility of transcription factors to their binding sites on the DNA (Sudarsanam and Winston, 2000). Ing3 is a component of the NuA4 histone acetyltransferase (HAT) complex (Doyon et al., 2004). These proteins are generally linked to gene activation.

Some polycomb family genes (PcG) such as Suz12, Cbx4, and Rnf2 (Ringrose and Paro, 2004) and the DNA methylation genes Dnmt3a and Mbd2 (Li, 2002) linked to gene silencing are up-regulated in the mid-foregut region as the lung bud forms. Many of these genes are ubiquitously expressed but an increase in the level of expression during lung induction highlights the importance of chromatin remodeling and DNA methylation in lung organogenesis (Lee et al., 2006).

### **1.7 Anterior-posterior patterns of gene expression in the foregut endoderm**

The presence of distinct molecular fields within the anterior-posterior axis of the foregut endoderm was shown by QRT-PCR analysis of laser capture microdissected foregut epithelium that is free of adjacent mesenchymal cells (Figure 5A). This procedure allows the collection of regional epithelial samples representing anterior, mid, and posterior areas of the foregut endoderm. In 21–25 somite samples the patterns of expression of three transcription factors differ along the anterior-posterior foregut axis (Figure 5B). Irx5, a member of the Iroquois homeobox gene family expressed in the E9.5–10.5 foregut (Cohen et al., 2000), is expressed at similar levels in the three regions. In contrast Hlxb9 mRNA is highly expressed in the most posterior sample with low to marginally detectable levels in the mid and anterior samples. This is consistent with its essential role in specification of gut epithelial cells to a pancreatic fate (Li et al., 1999). FoxO1, a homeobox transcription factor shown to inhibit pancreas-specific genes (Kitamura et al., 2002), is expressed in a linear gradient opposite to that of Hlxb9 with the highest levels anteriorly. In 26–30 somite samples, the patterns of expression of three genes identified in the microarray analysis, Isl1, Dach2, and smarce1/BAF57, also differ along the anterior-posterior foregut axis (Figure 5C). Isl1 is detected in the mid-and posterior foregut endoderm at similar levels, but higher than in the anterior endoderm. Dach2, is higher in the mid-foregut endoderm than in neighboring regions, while smarce1 is higher in the posterior region, although it is expressed in other regions of the foregut.

### **1.8 Patterns of expression of genes identified in the microarray analysis**

To further validate the microarray data we analyzed by whole mount in situ hybridization the pattern of expression of a number of genes involved in regulation of transcription. The transcription factor Isl1 is significantly up-regulated in the mid-foregut at the time of lung bud formation by microarray and QRT-PCR analyses. Whole mount in situ hybridization performed in 26–30 somite embryos (E9.5) (Figure 6A and 6B) showed expression of Isl1 mRNA in primary lung-bud mesenchyme. Although at this stage Isl1 is no longer expressed in cardiac cells, it can be detected in the sinus venosus (Figure 6A). Expression is also high in the stomach region (Figure 6B). Isl1 is essential for motor neuron differentiation and normal development of the heart, pancreas, and splanchnic mesenchyme, but its importance in lung development is not known due to the early lethal phenotype of the Isl1 null mutant mice (Ahlgren et al., 1997; Cai et al., 2003; Pfaff et al., 1996; Thaler et al., 2004). Assessment of Isl1 expression by QRT-PCR in total lung at different developmental time points shows that Isl1 expression is transient, since the mRNA level peaks on embryonic day E9.5 and decreases at E11.5 to that of E8.5 (Figure 6C). This is followed by a gradual decline during later development to levels that are about 10–15% of the peak value. Isl1 continue to be detected in adult lung although the levels are lower than the ones observed during lung budding.

Expression of the forkhead transcription factor FoxG1 was localized in the mesenchyma of the primary lung buds at E9.5 and in the lung mesenchyme at E11.5 (Figure 6D–E). FoxG1 is involved in morphogenesis of the telencephalon by controlling proliferation and differentiation of precursor cells (Martynoga et al., 2005). It participates in different signaling pathways. FoxG1 interferes with TGFβ pathway by association with Smad-interacting proteins (Seoane et al., 2004); it also interacts with the delta/notch/hes pathway by combining with Hes homodimers to repress transcription (Yao et al., 2001).

Two genes involved in chromatin modification identified in the microarray analysis, Carm1 and Cbx4, were barely detected by whole mount in situ hybridization in the primary buds at E9.5 (data not shown). Their expression, though, is highly increased by E11.5. Carm 1 is concentrated in mesenchymal cells at the tips of the lung branches (Figure 6F), while Cbx4 is in the epithelium (Figure 6I). Carm1 (Coactivator associated arginine methyltransferase) is a transcriptional activator that interacts with the p160 family of nuclear receptor-associated factors and methylates histone 3 at arginine 17. Methylation of arginines by Carm1 occurs along with acetylation of histones to remodel chromatin and recruit RNA polymerase II (Teyssier et al., 2002). Cbx4 (Chromobox homolog 4) is one of the five mouse Polycomb homologs that act as transcriptional repressors. Cbx4 binds to chromatin preferentially to histone 3 trimethylated in lysine 9 (H3K9me3) (Bernstein et al., 2006) and promotes SUMOylation of transcriptional repressors such as the DNA methyltransferase Dnmt3a (Li et al., 2007). Three other transctiption factors Zfp26 (Chowdhury et al., 1988), musculin (Lu et al., 1999), and Rfx5 (Yong et al., 2007) that are up-regulated in the mid-foregut during lung budding (Table 1) can also be detected by whole mount in situ hybridization at E9.5 in a very distinctive pattern. Zfp26, a zinc-finger transcription factor is detected in the sub-epithelial mesenchyme of the primary buds (Figure 6G). At E10.5 it is barely detected with some expression remaining at the tips of the forming branches (Figure 6H). Musculin, or MyoR, a repressor of muscle differentiation that competes with the myogenic factor MyoD, is expressed in the mid-region of the foregut, in particular in the esophagus, but it is not expressed in the primary lung buds (Figure 6J). Expression can be detected also in the branchial arches and the stomach. The branchial arches continue to be positive at E10.5 and the lung is negative (Figure 6K). In contrast Rfx5 is expressed in mesenchymal cells of the branchial arches, primary lung buds and stomach in the sub-epithelial layer (Figure 6L). Expression of Zbtb1, Ing3, Rbbp4 and Smarce 1 was detected in the mid-region of the foregut and/or whole embryos at E9.5 by whole mount in situ hybridization but they are ubiquitously expressed (data not shown).

### **1.9 Concluding Remarks**

In this study we have characterized the transcriptome of the mid-region of the mouse foregut during the period between lung specification of foregut cells and appearance of the initial lung buds. We have identified a substantial number of genes present in the prospective lung region, and genes that change their level of expression as the lung bud forms. We selected, for further studies, those involved in regulation of transcription since they were overrepresented among the genes that change as the lung forms, and because these factors likely activate the developmental program that sustains lung formation. The information generated, therefore, expands the list of genes to study new and known pathways driving lung progenitor cell differentiation and lung morphogenesis.

### **2. EXPERIMENTAL PROCEDURES**

### **2.1 Isolation of mid-foregut tissue**

We selected based on morphological features, three developmental stages to study gene expression profiles of mid-foregut tissue, containing endoderm and mesoderm cells. The stages are: 16–20 somite embryos, 21–25 somite embryos and 25–30 somite embryos. The earliest samples (16–20 somites) show evidence of primitive thyroid and liver/pancreas budding which provide morphologic limits at the extremes of the mid-foregut field from which the lung is derived. All embryos with greater than 25 somites have initiated formation of bilateral lung buds. Although some variability is expected in the relationship between somite number and morphology, there is a high degree of uniformity within the range of number of somites used to group the earliest and latest samples for molecular analysis.

Pregnant CD-1 mice were purchased from Charles River with pregnancy timed by the presence of a vaginal plug at day E0.5 of gestation. Dams were sacrificed at 6–12 hour intervals between gestational days E8.5 and E10 to obtain embryos containing between 16 and 30 somites. After hysterotomy, embryos were placed in PBS at 4°C prior to dissection. Each embryo was examined by dissecting microscopy to determine its somite number. Embryos were divided into three groups according to somite number: 16–20, 21–25, and 26–30 somites. This grouping correlates well with morphological features described above: 16–20 somite embryos have thyroid and liver buds but lack lung buds; all 26–30 somite embryos have lung, thyroid, and liver buds. The 20–25 somite group includes a small number of embryos in which lung buds have just commenced to emerge but are difficult to visualize. To obtain mid-foregut tissue, the neural tube and heart were dissected away for the embryo to expose the foregut. The foregut tissue posterior to the pharyngeal arches and anterior to the liver was excised using tungsten needles. Collected tissues were placed immediately in RNeasy™ buffer (QIAGEN). Five to ten mid-foreguts were pooled for each of the three somite groups for RNA analysis.

For histological analysis embryos or isolated mid-foreguts were fixed in 4% freshly prepared paraformaldehyde in PBS, stored in fixative overnight at 4°C, and embedded in paraffin following standard processing with ethanol dehydration. For laser capture microdissection embryos were fixed in 70% alcohol overnight at 4°C, dehydrated and embedded in paraffin in RNAse-free conditions (Goldsworthy et al., 1999). Paraffin blocks were examined in a dissecting microscope with fiberoptic lighting to determine the orientation of the embryo. Excess paraffin was trimmed away, and the embryo was then reembedded oriented in a supine position (ventral side up). Six micron transverse sections were prepared for further study.

### **2.2 RNA purification and amplification and microarray quality controls**

Total RNA was isolated from dissected tissues using RNeasy™ micropurification kit (QIAGEN) according to the manufacturer's directions, followed by treatment on column with DNA-*free*™ DNase (Ambion). RNA concentrations were measured in 1 μl (1/10 of the sample) in a NanoDrop ND-1000 spectrophotometer (NanoDrop Technologies). RNA (100 ng) was amplified using the RiboAmp HS kit (Arcturus Engineering, Inc) as described by the manufacturer. RNA was labelled using biotinylated ribonucleotides during the second *in vitro* transcription step using ENZO kit (Affymetrix). After two rounds of amplification, 10– 15 μg of amplified RNA (aRNA) were obtained. The quality of amplified RNA obtained from microdissected foreguts was evaluated by examining the size distribution of the aRNA and aRNA from E18.5 fetal lung, used as control, in an agarose gel stained with Sybr Gold (Molecular Probes). A similar size distribution of aRNA in the three samples, ranging from  $\sim$ 200bp to  $>$  2kb is shown in (Figure S2A, supplementary material). Thus the handling and time required for isolation of the samples has not resulted in degraded RNA unsuitable for microarray hybridization. Acceptable correlation values  $(R^2)$  values indicating reproducibility of microarrays are shown in Figure S2B (technical replicates  $R^2 = 0.973$ ) and Figure S2C (biological replicates  $R^2 = 0.969$ ) in the supplemental material.

### **2.3 Microarray experiments**

Gene expression profiles of 3 independent mRNA samples for each of the three somite number groups were determined using MOE430 A2.0 microarrays (Affymetrix), containing 22,690 probe sets. Genes represented on this array are available at

[http://www.affymetrix.com/products/arrays/specific/mouse430\\_2.affx.](http://www.affymetrix.com/products/arrays/specific/mouse430_2.affx) Each scanned image was evaluated for significant artifacts. Bacterial genes spiked into the hybridization mixture (bioB and bioC) were used as positive quality controls for hybridization procedures. Arrays showing similar quality control parameters (Table S1, supplementary material) were used for data analysis. Background and noise measurements were below 100 in all arrays. The average ratio of signals from mid-sequence and 3′ probe sets for GAPDH and actin was consistent

within all arrays, indicating similar efficiency of the amplification step. This allows comparison of amplified genes between independent arrays. As expected these ratios were higher than in non-amplified samples due to the preferential amplification of the 3′ end of the mRNA using polydT. Minimal degradation of the isolated mRNA could have occurred, even though strict RNAse-free conditions were used in every step of the experiments to avoid this problem. Data from each array was scaled (target intensity of 100) to normalize the results for inter-array comparisons. Reproducibility of the amplification and hybridization experiments was determined by correlation analysis of the microarray data obtained in replicates of 26–30 somite foregut RNA, starting from the same RNA (technical replicates) or from RNA from different embryos (biological replicates).

### **2.4 Microarray data analyses**

A single weighted mean expression level for each gene was derived using Microarray Suite (MAS) 5.0 software (Affymetrix). Using a one-sided Wilcoxon signed-rank test, the MAS 5.0 software generated a detection p-value for each gene indicating whether or not the transcript was reliably detected. An initial mild filter was applied to select genes with detection p-value less than 0.05 in at least one of the nine arrays. A total of 13,371 probe sets passed the filter and were used in further statistical analyses. A parametric ANOVA was performed to find genes differentially expressed between two or more of the groups. 1076 genes passed the ANOVA test at  $p < 0.05$ , a significantly larger number than expected by chance (650 genes would be expected by chance). 80% of the 1076 genes were indeed detected in more than 7 arrays of the 9 arrays analyzed. While a significant proportion of genes identified on statistical analysis may represent false positives, we limited our biological analysis to those whose fold changes were greater than 1.8 fold. In addition, a large number of biologically relevant genes were validated by other methods. We performed a K-means clustering analysis with the 1076 genes that passed the ANOVA test ( $p \le 0.05$ ), to determine groups of genes that have similar time course profiles of expression levels over the three time points studied. To make the different genes comparable to each other in the cluster groups, the average of each sample group was calculated and z-score-normalized so the mean of the averages is zero and the standard deviation is one. K-means clustering was performed in Spotfire DecisionSite arbitrarily setting the number of clusters (k) equal to five. This method allows the identification of genes enriched in each of the somite groups. A Student's t-test was also performed ( $p \leq$ 0.05) to compare the earliest vs. the latest time point (16–20 somites to 26–30 somites). We used EASE (Expression Analysis Systematic Explorer,

[http://david.abcc.ncifcrf.gov/ease/ease.jsp,](http://david.abcc.ncifcrf.gov/ease/ease.jsp) and the GO functional classification database to discover enriched biological themes within the lists of genes which are increased or decreased  $(> 1.8$  fold) as the lung bud forms  $(16–20)$  somite samples vs.  $26–30$  somite samples). We have opted to use 1.8 fold as cut-off level due to the high number of genes showing a statistically significant change between 1.8 and 2-fold. The genes used in the EASE analysis pass ANOVA  $(p \le 0.05)$  and Student's t tests ( $p \le 0.05$ ). EASE analyzes a list of Affymetrix ID numbers of the genes under study (input list) and finds over-represented biological "themes" in the list of genes, compared to the total number of genes for each biological theme in the array. The significance of each category is determined by two statistical values that are used to sort the categories i.e. the standard Fisher exact probability, and a conservative EASE score that identifies robust categories. We considered categories with an EASE score  $\leq 0.05$  as overrepresented.

### **2.5 Real Time RT-PCR**

Selected genes identified as differentially expressed in the mid-foregut by microarray analysis were validated by real time RT-PCR (QRT-PCR) in non-amplified mid-foregut RNA samples. Total non-amplified RNA from embryonic mid-foreguts grouped as before was treated with DNA-*free*™ DNase (Ambion) and reversed transcribed (RT) (1 μg of RNA in 25 μl reaction

volume) using AMV reverse transcriptase (Promega). RT reactions, diluted appropriately to obtain a signal in fewer than 34 cycles, were analyzed by QRT-PCR in an ABI 7000 Sequence Detection System (Applied Biosystems). PrimerExpress software version 2.0 (Applied Biosystems) was used to design primers for SybrGreen reactions, while TaqMan primers and probes used were Assays-on-Demand (Applied Biosystems) (Table S2, supplementary material). Reactions were performed in 50 μl using SybrGreen PCR master mix or TaqMan PCR universal master mix (Applied Biosystems). Optimal reaction conditions were determined for all primers and probes and dissociation curves were generated for primers used in SybrGreen reactions to confirm a single PCR amplification product. A calibration curve, generated with serial dilutions of reverse transcribed E18.5 total lung or E8.5 total embryo RNA (n=3), was used to determine the relative message concentration for each gene tested. Data were normalized to relative concentration of GAPDH mRNA amplified from the same RT reaction. Using equal amounts of total RNA from 16–20, 21–25 and 26–30 somite groups we established that the relative expression level of GAPDH is similar at all three time points (data not shown). GAPDH was therefore used to normalize gene expression in QRT-PCR experiments.

### **2.6 Whole mount in situ hybridization**

Whole mount in situ hybridization was performed as described by Wilkinson (Xu, 1998) and modified by Desai et al (Desai et al., 2004). Anti-sense and sense probes were generated by RT-PCR using oligonucleotides containing adaptors for T3 or T7 promoters (see oligonucleotide sequences in Table S8, supplementary material). Purified PCR fragments were used as templates in the *in vitro* transcription reaction (Maxiscript kit, Ambion) to synthetize sense and antisense riboprobes labeled with digoxigenin (DIG). After hybridization and staining of the embryos with BMpurple (Boehringer Mannheim), some were dehydrated and embedded in paraffin. Transverse sections (10 μm) were obtained as described above, and counterstained with Fast Red using standard methods.

### **2.7 Laser Capture Microdissection (LCM) of foregut endoderm**

We isolated endoderm cells from the anterior (thyroid region), mid (lung region) and posterior (liver/pancreas region) foregut of 21–25 and 26–30 somite embryos by laser capture microdissection. Embryos were fixed, embedded, and oriented in a supine position as described above. Six micron transverse sections were placed on plain glass slides, dried for 30 min at  $37^{\circ}$ C and immediately deparaffinized and dehydrated in xylenes ( $2 \times 5$  min), absolute ethanol (30 sec) and xylenes  $(2 \times 5 \text{ min})$ . Slides were dried at room temperature for 10 min and placed in slide boxes containing Drierite to keep them dry while dissecting. Foregut endoderm was microdissected using a PixCell I laser capture microdisection system (Arcturus) with a laser spot size of 30 μ, pulse power 15 mW, pulse width 15.2 ms. Sections from two embryos were collected per CapSure LCM cap (Arcturus) and total RNA was purified as described above. RNA purified from 6–10 embryos was pooled for QRT-PCR and  $[32P]RT-PCR$  analyses.

### **Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

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**Figure 1. Mid-foregut tissue was isolated to study gene expression profiles at three developmental stages**

**A.** Graphic representation of the morphology of the foregut in embryos containing 11–25 somites (E8.0–E9.5), and in embryos containing more than 25 somites (E9.5–10). Endoderm derived organs (thyroid, pancreas, liver, and lung) are depicted. The relative position of those organs to the cardiac mesoderm is represented. **B.** Microscopic appearances of partially dissected mid-foreguts from 16–20 somite embryos (E8.5) (red arrowhead), and from 26–30 somite embryos (E9.5). The mid-foregut tissue between the black dashed lines was collected for further analyses. The earliest samples (16–20 somites) show evidence of primitive thyroid and liver/pancreas budding at the extremes of the mid-foregut field from which the lung is derived. Embryos with greater than 25 somites have initiated formation of bilateral lung buds

(outlined in yellow). Ba, branchial arches; Li, liver; Lu, lung buds, outlined by yellow dashed lines. **C.** TTF-1 (Nkx2.1, Titf1) and FGF10 mRNA levels, assessed by real time RT-PCR, in mid-foreguts dissected from 16–20, 21–25 and 26–30 somite mid-foreguts. Titf-1 and FGF10 mRNA levels are normalized to GAPDH. Data are expressed relative to 16–20 somite midforegut samples. n=3 Error bars represent standard error of the mean. \* indicates t-test  $\leq 0.05$ .



### **Figure 2.**

Cluster graphs of (**A**) genes up-regulated ≥ 1.8 fold, and (**B**) genes down regulated ≥ −1.8 fold from 16–20 somite to 26–30 somite embryos (detection  $p \le 0.05$ , ANOVA  $p \le 0.05$ , and Student's test  $p \le 0.05$ ). Green color indicates low expression, red indicates high expression. Three independent arrays were analyzed for each developmental stage (arrays 1–3, 16–20 somite mid-foregut samples; arrays 4–6, 21–25 somite mid-foregut samples; arrays 7–9, 26– 30 somite mid-foregut samples). **C.** Pie chart showing the GO classification of the functional groups of the 104 genes up-regulated shown in A. **D.** Pie chart showing the GO classification of the functional groups of the 119 genes down-regulated shown in B.



### **Figure 3.**

Patterns of gene expression identified by K-means clustering analysis of genes that pass the ANOVA test comparing the three developmental points studied 16–20, 21–25, and 26–30 somite embryos (ANOVA  $p \le 0.05$ ). **A.** genes that are down-regulated from 16–20 to 26–30 somites; **B.** genes that are down-regulated from 16–20 to 21–25 somites; **C.** genes that peak at 21–25 somites; **D.** genes that are up-regulated from 16–20 to 26–30 somites; **E.** genes that are up-regulated from 21–25 to 26–30 somites. The complete list of genes is in the supplemental material. Axes: x, developmental stage; y, z-score normalized values, 0=mean of the average for each gene.  $+/-1$  = standard deviation.



### **Figure 4.**

Real time RT-PCR validation of selected transcription related genes identified by microarrays. **A.** Expression levels obtained in the microarray analysis of amplified RNA from 16–20 somite mid-foreguts (relative value =1, black bars) compared to 26–30 somite mid-foreguts (fold change relative to 16–20 somite samples, hatched bars). **B.** Real time RT-PCR validation of the genes depicted in A. Non-amplified RNA from 16–20 somite mid-foreguts (relative value =1, black bars) compared to 26–30 somite mid-foreguts (fold change relative to 16–20 somite samples, hatched bars). Data are normalized to GAPDH expression level. n=3. Error bars represent standard error of the mean.

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**Before** 

After

Cap



### **Figure 5.**

**A.** Representative laser capture microdisection of foregut endoderm from (anterior) thyroid region, (mid) lung region, and (posterior) liver/pancreas region from embryos containing 21– 25 somites. (Before) picture of a transverse tissue section before laser capture, (after) tissue remaining on the slide after dissection, (cap) tissue collected on the cap membrane. Nt, notochord; Fg, foregut; Th, thyroid bud; Li, Liver bud; H, heart. **B.** Real time RT-PCR of three selected transcription factors that show distinctive patterns of expression along anterior (A), mid (M) and posterior (P) foregut endoderm from 21–25 somite embryos. **C.** Real time RT-PCR of three selected transcription genes that show distinctive patterns of expression along anterior (A), mid (M) and posterior (P) foregut endoderm from 26–30 somite embryos.

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### **Figure 6.**

Pattern of expression of selected genes in the primary lung buds and developing lung determined by whole mount in situ hybridization. **A.** Isl1 is mainly expressed in the mesenchyme of the primary lung buds (yellow arrowheads) and the stomach of embryonic foreguts at E9.5. **B.** Transverse section of paraffin embedded whole embryos after WMISH, counter stained with Fast Red, shows expression of Isl1 in the neural tube (red arrows), the tips of the forming lung buds and the sinus venosus at E9.5 (yellow arrowheads). **C.** Expression levels of Isl1 mRNA were determined by real time RT-PCR at different stages of lung development and in adult lung. Isl1 expression level is normalized to GAPDH. n=3. Error bars represent standard error of the mean. **D.** FoxG1 is expressed in the mesenchyme of the primary

lung buds (yellow arrowheads) and stomach at E9.5. **E.** FoxG1 is diffusely detected in the lung mesenchyme at E11.5 **F.** Carm1 is concentrated in the mesenchyme at the tips of the lung branches at E11.5 (yellow arrowheads). **G.** Zfp26 is expressed in the mesenchyme close to the epithelium in the forming primary buds at E9.5 (yellow arrowheads). **H.** Zfp26 is faintly detected at the tips of the branches at E10.5 (yellow arrowheads). **I.** Cbx4 is expressed in the epithelium of the lung branches at E10.5 (black arrowheads). **J.** Musculin is absent in the primary lung buds at E9.5 but is expressed in the mid foregut along the esophagus. It is also expressed in the branchial arches and stomach, likely in muscle progenitor cells. **K.** Musculin is not detected in the lung at E10.5 but expression continues in the branchial arches. **L.** Rfx5 is expressed in the subepithelial mesenchyme alongside the foregut, including the primary lung buds at E9.5 (yellow arrowheads). Fg, foregut; Nt, neural tube; H, heart, Sv, sinus venosus; Ba, branchial arches; Lb, primary lung bud; Lu, lung; Tr, trachea.



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**Table 1**



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 $\mathbf{I}$ 

 $\Big|$  =  $\Big|$ 

 $2342$ 

 $2.5$ <br> $1.6$ <br> $1.0$ 

*Suz12* 1449661\_at 1.0 2.5 2.9 *a Ing3* 1450760\_a\_at 1.0 1.6 2.4 *t Smcy* 1424903\_at 1.0 1.0 2.2 **a** 

 $\frac{1449661_a}{1450760_a}$ <br>1450760\_a\_at<br>1424903\_at

 $\frac{Suz12}{Ing3}$ 





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# **Table 2**<br>Selected families of transcription factors detected in the mid-foregut. Selected families of transcription factors detected in the mid-foregut.



 $a = ANOVA$   $p \le 0.05$ a= ANOVA p ≤ 0.05 The table includes the average signal for genes detected in the mid-foregut and lung regions (detection p  $\leq 0.05$ ) at 16–20s, 21–25s and 26–30s developmental stages. Only genes with a signal  $\geq 100$  at any stage are The table includes the average signal for genes detected in the mid-foregut and lung regions (detection p ≤ 0.05, 21–25s and 26–30s developmental stages. Only genes with a signal ≥ 100 at any stage are listed. Most genes do not change the level of expression as the lung buds. A few of them change the level of expression significantly indicated by Student's t and/or ANOVA tests.

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 NIH-PA Author ManuscriptNIH-PA Author Manuscript **Table 3**<br>Selected signaling pathways detected in the mid-foregut.

Selected signaling pathways detected in the mid-foregut.





t= Student's t-test  $p \le 0.05$ t= Student's t-test  $p \le 0.05$ 

a= ANOVA  $p \le 0.05$ a= ANOVA p ≤ 0.05 The table includes the average signal for genes detected in the mid-foregut and lung regions (detection  $p \le 0.05$ ) at 16-20s, 21-25s and 26-30s developmental stages. Only genes with a signal  $\ge 100$  at The table includes the average signal for genes detected in the mid-foregut and lung regions (detection p ≤ 0.05, 21–25s and 26–30s developmental stages. Only genes with a signal ≥ 100 at any stage are listed. Most genes do not change the level of expression as the lung buds. A few of them change the level of expression significantly indicated by Student's t and/or ANOVA tests. any stage are listed. Most genes do not change the level of expression as the lung buds. A few of them change the level of expression significantly indicated by Student's t and/or ANOVA tests.

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