Expression of a novel surfactant protein gene is associated with sites of extrapulmonary respiration in a lungless salamander

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SFTPC expression specificity

The expression pattern of SFTPC is highly conserved: all tetrapods express SFTPC exclusively in the lungs [1–9]. In anamniotes, SFTPC is expressed throughout the lung [5,6,10], whereas in mammals it is confined to alveolar type II cells. Four reports cite expression of SFTPC outside of the lungs in humans, but each report has methodological problems, including possible contamination, that may make such claims unreliable. Mo et al. (2006) report SFTPC in human fetal and adult skin [11]. This claim, however, relies on immunohistochemical data obtained with an antibody that may yield spurious labeling, and on RT-PCR data that was not followed up with sequencing. RT-PCR is subject to contamination and mispriming. Bräuer et al. (2009) report SFTPC expression in submandibular and parotid glands based on RT-PCR and western immunoblots [12]. The western blots reveal a protein of the expected size of the SFTPC pro-protein, and RT-PCR of focal cDNA is performed alongside a positive control, but there is no follow-up sequencing. Schicht et al. (2015) report SFTPC in saliva from human patients based on western blots and ELISA [13]. The antibody used to detect SFTPC is not identified, however, and the isolated band is at 16 kDa while SFTPC proprotein is 21 kDa and mature SFTPC 3.7 kDa [14]. Additionally, saliva may be subject to contamination by surfactant produced in the lungs. Finally, Schob et al. (2013) report SFTPC expression in central nervous system tissue and cerebrospinal fluid based on RT-PCR, but they fail to rule out the possibility of genomic DNA contamination [15]. Their western blots also fail to demonstrate a band at the expected size for SFTPC given the antibody they employed, and the authors express confusion about how mRNA for SFTPC could be present in cerebrospinal fluid. In sum, there are problems with all recent studies that cite extrapulmonary expression of SFTPC in humans. At the same time, numerous studies in mammals and frogs, including ISH and reporter knockins, have failed to demonstrate extrapulmonary expression of SFTPC. Nevertheless, it remains possible that humans and perhaps other animals endogenously express SFTPC outside of the lungs. Optimally, in situ hybridization and transcriptome sequencing should be used to validate the human results presented above.
Evidence for Duplication of SFTPC

SFTPC and SFTPC-like sequences diverge from one another within exonic regions, but not according to putative splice boundaries (Fig. S1a), which indicates that SFTPC-like is not an isoform of SFTPC. While SFTPC-like is divergent from SFTPC sequences, it is not an ortholog of a closely related BRICHOS domain-containing gene (Fig. S1b). We found SFTPC-like expressed in eleven species of lunged and lungless salamanders (Fig. S1a, b; Supplemental Data File 1); most also express SFTPC.

In Bayesian and maximum likelihood gene trees for SFTPC sequences from amphibians, amniotes, lungfish and coelacanth, tetrapod SFTPC and SFTPC-like fall into well-supported monophyletic lineages. However, support values for the split between SFTPC and SFTPC-like are low, leaving a polytomy including lungfish SFTPC, coelacanth SFTPC, salamander SFTPC-like and tetrapod SFTPC (Figs. S1, S2). Bayesian (Fig. S1b) and maximum likelihood (Fig. S2) gene trees both place coelacanth SFTPC sister to salamander SFTPC-like, but support values for a sister relationship to salamander SFTPC-like are low under the maximum likelihood approach. The presence of SFTPC in all amphibian and amniote groups and the absence of SFTPC-like in all surveyed tetrapods except salamanders argues that the most parsimonious explanation for the corresponding gene tree is that the tetrapod ortholog of SFTPC was duplicated in the salamander lineage following its divergence from frogs, followed by substantial sequence divergence between SFTPC and SFTPC-like (Fig. S1b). Low statistical support at the salamander SFTPC-like + coelacanth SFTPC node under maximum likelihood approaches should be interpreted as a polytomy, and long-branch attraction may cause an artifactual affiliation between coelacanth and lungfish SFTPC orthologs and salamander SFTPC-like. Therefore, origin of SFTPC-like from a salamander-specific gene duplication event is the most parsimonious explanation for the observed gene trees. However, alternative scenarios such as independent losses of SFTPC-like across lungfish, frogs and amniotes cannot be ruled out based purely on gene tree topology from Bayesian and maximum likelihood approaches alone.

Sequence alignments alone may not provide sufficient information to reconstruct gene evolution and patterns of gene gain and loss [16]. We applied PHYLDOG [16] to explicitly test for gene duplications of SFTPC. Given a guide tree with known phylogenetic relationships (Fig. S3), PHYLDOG predicts that SFTPC-like arose by gene duplication after the divergence of frogs and salamanders (Fig. S4). SFTPC-like has been meiotically mapped to linkage group 6 in Ambystoma.
mexicanum, a lunged salamander, and is located within a region syntenic to human chromosome 15 [17]. Finally, SFTPC-like and SFTPC have been assembled to two separate genomic scaffolds from A. mexicanum [18], supporting SFTPC-like’s origin via gene duplication.

Previous work identified a SFTPC gene duplication event in the lungfish Protopterus annectens [19]. Our analysis confirms that this duplication event is lineage specific and confined to P. annectens (Fig. S1b).
Supporting Information Tables:

**Table S1. Primers used to clone SFTPC and SFTPC-like from Ambystoma mexicanum, Desmognathus fuscus and Plethodon cinereus.**

| Gene       | Species                  | Forward                                       | Reverse                                      |
|------------|--------------------------|-----------------------------------------------|----------------------------------------------|
| SFTPC      | *A. mexicanum*           | 5'-CAC ACA GAR AMG ATT TTC CAG ATG-3'         | 5'-CGT CTT GTC CAT TTT TGT KAB GTA GCA-3'    |
| SFTPC-like | *A. mexicanum*           | 5'-AAG ATG GAA ACC GGC AGC AAG C-3'           | 5'-CGT CTT GTC CAT TTT TGT KAB GTA GCA-3'    |
| SFTPC-like | *D. fuscus*              | 5'-AAG ATG GAA ACC GGC AGC AAG C-3'           | 5'-AGT ATT GGA AGC GGT CTG GGT G-3'          |
| SFTPC-like | *P. cinereus*            | 5'-AAG ATG GAA ACC GGC AGC AAG C-3'           | 5'-GGT GTA GTC ATA GAC CAC-3'                |

**Table S2. Transcriptomes used to identify SFTPC and SFTPC-like.**

Accession numbers are listed in Supplemental Data File 1.

| Species                  | Transcriptome Source                                      |
|--------------------------|----------------------------------------------------------|
| Salamanders              |                                                          |
| *Ambystoma andersoni*    | Ryan Woodcock and Randal Voss                            |
| *Ambystoma mexicanum*    | Present study                                             |
| *Ambystoma tigrinum*     | Ryan Woodcock and Randal Voss                            |
| *Ambystoma tigrinum*     | [20]                                                      |
| *Andrias davidianus*     | [21]                                                      |
| *Cryptobranchus allemaniensis bishopi* | David Weisrock and Paul Hime                              |
| *Cynops cyanurus*        | David Weisrock and Paul Hime                              |
| *Desmognathus fuscus*    | David Weisrock and Justin Kratovil                        |
| *Ensatina eschscholtzii* | Rachel Mueller [22]                                       |
| *Hynobius chinensis*     | [23], reassembled by Paul Hime                            |
| *Hynobius retardatus*    | [24]                                                      |
| *Notophthalmus viridescens* | [25]                                                  |
| *Plethodon cinereus*     | Present study                                             |
| Frogs                    |                                                          |
| *Pelophylax nigromaculatus* | [26]                                                   |
| *Rana (Lithobates) pipiens* | [27]                                               |
| Dipnoi                   |                                                          |
| *Lepidosiren paradoxa*   | Igor Schneider [28]                                       |
| *Protopterus annectens*  | Chris Amemiya [28]; [19]                                 |
Figure S1. A novel form of Surfactant-associated protein C (SFTPC) is expressed in several species of salamanders. 

**a**, Amino acid alignment of SFTPC (yellow) and SFTPC-like (cyan) sequences reveals conservation of hydrophobic residues within the mature peptide domain. Full species names and accession numbers are listed in Supplemental Data File 1. Lungless (plethodontid) species are in red font. 

**b**, Bayesian 95% maximum clade credibility tree for SFTPC reveals SFTPC-like transcripts in 11 species of salamanders. SFTPC-like is not a related ortholog because it is nested within the SFTPC phylogeny. Node values are posterior probabilities; scale bar denotes expected changes per site. Salamander SFTPC is marked with a yellow box; salamander SFTPC-like is marked in cyan.

**c**, Predicted secondary structure of SFTPC-like from *Desmognathus fuscus*. SFTPC-like structure predictions (cyan) utilizing SWISS-MODEL [29], QUARK Ab initio predictions [30] and I-TASSER [31] are aligned with the resolved SFTPC mature peptide (yellow) [32].
Figure S2. Maximum likelihood gene tree for SFTPC and SFTPC-like. RAxML (v8.2.10) tree with 1000 bootstrap inferences and the JTT substitution matrix. Bootstrap support values from the maximum likelihood bootstraps are labeled on each node. Monophyly of coelacanth (\textit{Latimeria chalumnae}) SFTPC + salamander SFTPC-like (cyan box) is weakly supported. Salamander SFTPC is marked with a yellow box. Full species names and accession numbers are listed in Supplemental Data File 1. Lungless (plethodontid) species are in red font.
Figure S3. Guide tree for PHYLDOG. The NCBI taxonomic database was used to generate the tree topology, combined with Pyron and Wiens (2011) [33] for amphibian phylogenetic relationships.
**Figure S4. Gene duplications predicted by PHYLDOG.** The nodes where a gene duplication event is predicted are colored red. Blue nodes indicate divergence due to speciation events. PHYLDOG predicts that SFTPC-like (cyan box) originated due to gene duplication. Two other duplication events of SFTPC-like are predicted in salamanders. While some species of salamanders appear to express only one form of SFTPC-like, several species express SFTPC-like transcripts with slight sequence differences (Supplemental Data File 1). Further work is needed to determine if these sequence differences indeed represent further duplications of SFTPC-like or instead are the result of assembly error or alternative splicing. Only one SFTPC-like sequence per species was selected for phylogenetic analysis. Salamander SFTPC is marked with a yellow box. Lungless (plethodontid) species are in red font.
Figure S5. Histology of the integument in *Desmognathus fuscus* before and after metamorphosis. **a,** Tangential section through the gular region of a larva shows the layers of the integument (from left to right): flattened, cuticle-like keratinized layer (KL); an inner cell layer (IC); large cuboidal Leydig cells (LC) intermingled with capillaries and other supporting cells; basal lamina (BL). **b,** Sagittal section from the abdominal region of a larva. **c,** Transverse section from a recently metamorphosed specimen showing the acinous glands (AG) and a greatly thickened epidermis (EP). Mallory trichrome stain. Scale bars: **a,** 20 µm; **b,c,** 50 µm.
Figure S6. Additional images of SFTPC and SFTPC-like expression patterns. a, Wholemount embryos of *Ambystoma mexicanum* display SFTPC expression specific to the lungs (L). Lateral view; anterior is to the right. b, SFTPC sense control at the same stage shows no lung expression. c, Midsagittal section of *A. mexicanum* embryo stained for SFTPC shows expression in the trachea (T), but no expression in the integument or buccopharynx. Dorsal is up, anterior is to the right. d,e, SFTPC expression in *A. mexicanum* lung is confined to squamous epithelial cells lining the lumen. Sagittal section with anterior to the right. e is an enlargement of the boxed region in d. f, SFTPC-like expression in *Desmognathus fuscus* integument is confined to the apical cellular layer. Sagittal section with anterior to the left. g, SFTPC-like expression in adult *D. fuscus* buccal epithelium. Transverse section. Scale bars: d,e,f, 50 µm; a,b,c,g, 100 µm.
Figure S7. Ultrastructure of alveolar epithelial cells in adult *Ambystoma mexicanum*. **a**, Low magnification view of the pulmonary epithelium. The lumen of the lung is to the right. **b**, Enlargement of boxed area in **a**. **c,d**, Enlargement of boxed regions in **b** show lamellar bodies (LB) and secretory vesicles (SV). Scale bars: **a**, 2 µm; **b**, 1 µm; **c,d**, 200 nm.
Figure S8. Lamellar bodies in Desmognathus fuscus integument. a, b, Additional examples of lamellar bodies (LB) found in skin from a 24-mm larva of *D. fuscus*, a lungless salamander. The lamellar bodies are large (> 0.5 µm) and localized to the apical layer of epidermis. Each inset magnifies its corresponding boxed region (dashed lines). Apical is up in a and towards the upper right in b. Abbreviations: BL, basal lamina; LC, Leydig cell. Scale bars: a,b, 2 µm (insets: 500 µm).

Captions for Supplemental Data Files:

Supplemental Data File 1: Excel spreadsheet with all sequence data used for the study.
Supplemental Data File 2: A FASTA amino acid alignment used to generate the gene tree.
Supporting Information References:

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