Comparative transcriptomics in human COPD reveals dysregulated genes uniquely expressed in ferrets

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Abstract

Background: Chronic obstructive pulmonary disease (COPD) is a progressive lung disease with poor treatment options. However, most mouse models of COPD produce a primarily emphysematous disease not recapitulating clinically meaningful COPD features like chronic bronchitis.

Methods: Wild-type ferrets (Mustela putorius furo) were divided randomly into two groups: whole body cigarette smoke exposure and air controls. Ferrets were exposed to smoke from 1R6F research cigarettes, twice daily for six months. RNA-sequencing was performed on RNA isolated from lung tissue. Comparative transcriptomics analyses of COPD in ferrets, mice, and humans were done to find the uniquely expressed genes. Further, Real-time PCR was performed to confirmed RNA-Seq data on multiple selected genes.

Results: RNA-sequence analysis identified 420 differentially expressed genes (DEGs) that were associated with the development of COPD in ferrets. By comparative analysis, we identified 25 DEGs that are uniquely expressed in ferrets and humans, but not mice. Among DEGs, a number were related to mucociliary clearance (NEK-6, HAS1, and KL), while others have been correlated with abnormal lung function (IL-18), inflammation (TREM1, CTSB), or oxidative stress (SRX1, AHRR). Multiple cellular pathways were aberrantly altered in the COPD ferret model, including pathways associated with COPD pathogenesis in humans. Validation of these selected unique DEGs using real-time PCR demonstrated > absolute 2-fold changes in mRNA versus air controls, consistent with RNA-seq analysis.

Conclusion: Cigarette smoke-induced COPD in ferrets modulates gene expression consistent with human COPD and suggests that the ferret model may be uniquely well suited for the study of aspects of the disease.

Keywords: COPD, Chronic bronchitis, Ferret model, Gene expression, Cigarette smoke

Background

Chronic obstructive pulmonary disease (COPD) is a heterogeneous disease diagnosed typically by irreversible airflow obstruction and persistent respiratory symptoms [1, 2]. COPD remains among the top five leading causes of death worldwide [3] and still lacks pharmacologic treatments that can mitigate COPD progression and persistent symptoms of the chronic bronchitis phenotype that include chronic cough and sputum production. Genetic studies can reveal novel mechanisms and suggest potential therapeutic strategies for investigation.

Cigarette smoking and chronic exposure to other inhaled pollutants are principal environmental risk factors driving COPD pathogenesis known to exert...
dysregulation of gene expression in the lung as well as extrapulmonary tissues [4–6]. However, transcriptomics findings in human studies cannot control for phenotypic variations that can be achieved using experimental systems, and animal studies have largely been limited to studies in mice [5, 6]. Mice are among the most utilized models to study COPD [7, 8], yet produce primarily very mild emphysematous disease and do not develop features pathognomonic for chronic bronchitis or mucus hypersecretion [9]. To remedy this, we developed a cigarette smoke exposure model in ferrets that recapitulates features of emphysema as well as COPD-associated chronic bronchitis including airway remodeling, mucous metaplasia, muco-obstructive lung disease and periodic infectious exacerbations [10–13].

In the current report, we conducted transcriptome profiling of ferrets with COPD and compared findings to prior results from human COPD lungs [14] and a mouse model of COPD [6]. Overall, we sought to identify whether our ferret model of COPD captures unique genetic signatures in comparison to mouse models that can help improve understanding of the molecular pathogenesis of human COPD and promote the development of new and effective therapies.

Materials and methods

Cigarette smoke exposure in ferrets

Age and sex-matched ferrets (Mustela putorius furo) were procured from Marshall BioResources and exposed to room air or cigarette smoke generated by an automated cigarette smoke generator (TSE Systems, Chesterfield, MO). Smoke exposure consisted of 1 h of smoke from 1R6F research cigarettes, twice daily for at least 6 months. Animals were continuously monitored for particular matter (200 µg/L) and CO levels (~1% to ~3%). All animal experiments were approved by the Institutional Biosafety Committee of the University of Alabama at Birmingham (IACUC 20232).

RNA isolation

Following euthanasia, RNA was extracted from freshly frozen ferret lung samples using a RNeasy Mini Kit (Qiagen, Germantown, MD, USA) according to the manufacturer’s instructions. RNA integrity and concentration were examined prior to processing for RNA-seq.

Next-generation sequencing on Illumina platforms

mRNA sequencing was performed on an Illumina NextSeq 500 System as described by the manufacturer (Illumina Inc., San Diego, CA, USA). Briefly, the quality of the total RNA was assessed using the Agilent 2100 Bioanalyzer. RNA with an RNA Integrity Number (RIN) of 7.0 or above was used for sequencing library preparation. We used the Agilent SureSelect Strand Specific mRNA library kit as per the manufacturer’s instructions (Agilent, Santa Clara, CA, USA). Library construction began with two rounds of polyA selection using oligo dT containing magnetic beads. The resulting mRNA was randomly fragmented with cations and heat, which was followed by first strand synthesis using random primers with inclusion of Actinomycin D (2.4 ng/µL final concentration). Second strand cDNA production was done with standard techniques and the ends of the resulting cDNA were made blunt, A-tailed and adaptors ligated for indexing to allow for multiplexing during sequencing. The cDNA libraries were quantitated using qPCR in a Roche LightCycler 480 with the Kapa Biosystems kit for Illumina library quantitation (Kapa Biosystems, Woburn, MA, USA) prior to cluster generation. Cluster generation was performed according to the manufacturer’s recommendations for onboard clustering (Illumina). Paired end 75 bp sequencing runs were completed to allow for better alignment of the sequences to the reference genome.

RNA-seq analysis

RNA-seq data from 12 individual ferrets (six ferrets for air and smoke experimental conditions, respectively) were sequenced using an Illumina NextSeq 500 System (Additional file 1: Table S1). Paired end 75 bp reads were generated. Pre-alignment quality assessments of the raw fastq sequences were carried out using FastQC (version 0.11.7) (Andrews, 2010). The number of paired ends reads for the twelve samples range from 35 to 57 M (Additional file 2: Table S2, Additional file 3: Fig. S1). The raw fastq sequences were aligned to the Mustela putorius furo reference genome (GenBank assembly accession: GCA_000215625.1) [15]. The alignments were carried out using STAR (version 2.7.1a) [16]. --outFilterType BySJout --outFilterMultimapNmax 20 --alignSJoverhangMin 8 --alignSJdbOverhangMin 1 --outFilterMismatchNmax 999 --outFilterMismatchNoverReadLmax 0.04 --alignIntronMin 20 -alignIntronMax 1000000 -alignMatesGapMax 1000000; as per ENCODE STAR parameter options for RNA-seq). Post-alignment quality assessments were carried out with RSeQC (version 2.6.3) [17]. Samtools (version 0.0.19) [18] and IGV (version 2.6.2) were used for indexing and viewing the alignments respectively. Gene expression was quantified as gene level counts using the htseq-count function (version 0.12.3) [19]. The Ensembl gene annotations for Mustela putorius furo (genebuild-last-updated 2016-05) was used [20]. The htseq-count default parameters were used except for the strand parameter which was set to reverse to consider the strandedness of library. DESeq2 filtered the low counts; genes for which there are less than 3 samples with normalized counts greater than or equal to 4 were
filtered out. For each sample, the read counts were normalized using the Median Ratio Normalization method. Batch effects were evaluated using the ComBat function in sva (version 3.33.1) [21]; principal component analysis plots with the normalized gene expression data were used. Differentially expressed genes (DEGs) were identified using DESeq2 (version 1.28) [22]. DESeq2 was configured by adding batch and sex to the design to control for these factors whilst DESeq2 tests for association due to the condition [22]. Genes were differentially expressed if the absolute log2Fold change was > 1, and the adjusted p-value was < 0.1. The transformed normalized gene expression data with batch removal and adjustments for sex, using the limma function removeBatchEffect (version 3.44.0) [23], was used for downstream analyses (such as clustering analysis and heatmap generation (Fig. 1). The complex heatmap package (version 2.7.1.1015) [23] was used to generate heatmaps. The volcano plots were generated with an R Bioconductor package EnhancedVolcano (version 1.12.0) [24]. Circos plots were drawn with Circos (version 0.699) [25]. Interactivenn was used to generate Venn diagrams [26].

**Functional enrichment analysis**

To assess which biological functions were altered upon smoke exposure, over-representation enrichment analysis was performed using the WEB-based GEne Set AnaLysis Toolkit (Webgestalt) [27]. The functional database selected was gene ontology (biological process); the genome was selected as the reference set. A list of gene symbols was supplied (those genes that mapped between the ferret and human [20]; the genes supplied were DEGs. The organism of interest selected was Human. The hypergeometric test was used for over-representation evaluation for the lists of statistically altered DEGs. The Benjamini and Hochberg method was used to calculate the adjusted p-values (q). Other functional databases used were the gene ontology for molecular function and the biological pathway (KEGG). The detection of over-represented gene ontologies in the differentially expressed genes was also performed using the clusterProfiler package (version 4.0) [28, 29].

**Real-time PCR validation**

For qRT-PCR analysis, RNA was isolated using miRNeasy Mini Kit as per manufacturer protocol (Qiagen, Cat No.: 217004, USA). 50 nanogram of RNA was used
for further amplification by using TaqMan™ RNA-to-
CT™ 1-Step Kit (Thermo Scientific). Analyses of mRNA
levels were carried out using StepOnePlus Real-Time
PCR System (Life Technologies, Carlsbad, CA, USA),
TaqMan Universal PCR Mastermix (Applied Biosystems,
Waltham, MA, USA) and gene-specific Prime PCR Probe
Assays was designed and order from thermofisher (Assay
ID: APEPYE2, APT2FKD, APU696A, APFVTYY, APG-
ZM1W, APHF64U, APKCAPR, APMF4AN, APNKXXK,
APPRTFH and APRWKZF). GAPDH was used for refer-
ence housekeeping gene. Relative expressions were quan-
tified with the $2^{-\Delta\Delta CT}$ method [30].

**Data access**
The experiments performed in this study are available in
GEO under the accession number (GSE193749).

**Results**
**Evaluation of differentially expressed genes in smoke-exposed ferret lungs**
The transcriptome profile of COPD ferrets has not been
previously reported. To determine the gene expres-
sion signature of the ferret model, lung parenchyma was
derived from cigarette smoke-exposed or air control fer-
rets and assessed using RNA-Seq. In the ferret transcrip-
tomics data, the overall read mapping rate to the ferret
reference genome was between ranged between 80 and
92% and did not significantly vary by smoking status.
Principal component analysis of global transcriptomics
demonstrated clear discrimination of smoke and con-
trol exposed ferrets (Fig. 1A). Hierarchical clustering of
DEGs, shown by heat map after adjustment of transcript
levels by batch and sex (Fig. 1B), demonstrated distinct
expression patterns by exposure condition, with pro-
nounced alterations of the transcriptome in response to
smoke, in smoke exposed ferrets. Overall, a total of 420
DEGs were associated with smoke exposure in COPD
ferrets (Fig. 2, Additional file 1: Table S1) Cytochrome
P450 family 1 subfamily B member 1 (CYP1B1) was
highly upregulated in the model (5.1-fold increase, Ben-
jamini–Hochberg adjusted p-value = 2.41E−44), in line
with previous observations in humans with COPD [31]
and rodent exposure models [32], and thus served as an
important quality control.

**Gene ontology enrichment analysis**
To support these pathway analyses, we further acquired
an overview of Gene Ontology by evaluating the func-
tional annotations of ferret DEGs based on GO terms.
The top 15 associated functions in each component are
presented in Fig. 3 and the complete annotation is pro-
vided in Additional file 4: Table S3. Disease relevant path-
ways that were altered in biological processes (GO: BP)
primarily included neutrophil degranulation, neutro-
phil activation, and respiratory burst, whereas G protein-
coupled receptor signaling pathway and cell adhesion
were down regulated (Fig. 3A; Additional file 4: Table S3).
Molecular functions (GO: MF) notable included receptor
signaling pathways and oxidoreductase activity (Fig. 3B;
Additional file 4: Table S3), whereas cellular component
GO terms included extracellular matrix and NADPH oxi-
dase (Fig. 3C; Additional file 4: Table S3).

**Functional and gene ontology (GO) enrichment analysis of
differentially expressed genes in smoke-exposed ferret lungs**
DEGs associated with cigarette smoke exposure identi-
fied from the analysis of ferret lung were next subjected
to functional enrichment analysis. Top ten upregulated
and downregulated KEGG pathways are shown in Table 1
in rank order by p-value, noting many were not statisti-
cally significant by FDR.

**Comparison of differentially expressed genes in
cigarette smoke-exposed ferret lungs, mouse lungs, and human
COPD**
A unique property of the COPD ferret model is that it exhibits classical pathophysiological features of air-
way disease, including goblet cell hyperplasia, mucus hyperexpression, and delayed mucus clearance, as well
phenotypical features of chronic bronchitis including
Fig. 3 Gene ontology (GO) annotation plots for differentially expressed genes (DEGs) identified in smoke-exposed ferrets. Bubble plots showing GO terms affected for A biological Process, B molecular function, and C cellular component categories, by DEG count and FDR in smoke exposed ferrets as compared to controls.
cough [10–12]. We next sought to understand whether COPD ferrets can provide insights into the molecular basis of human COPD that otherwise cannot be examined in COPD mouse models. We next sought to evaluate the extent of overlap between the genetic signatures of DEGs in ferrets as compared to human and/or mice. To do that, overlap was assessed between the COPD ferret gene expression signature and DEGs in datasets from Obeidat et al. evaluating mice that were smoke-exposed for 24 weeks duration [6] and Bosse et al. that evaluated humans with severely affected COPD [14], a population most relevant to the duration and intensity of chronic smoke exposure experienced by ferrets [12]. Of the 420 DEGs identified in COPD ferrets, 266 DEGs also had analyzable gene expression data generated in human [14] and mouse [6] models enabling comparative transcriptomic analyses between ferret, human, and mouse lungs. As depicted in Fig. 4A Venn diagram and detailed in the Circos plot in Fig. 4B that illustrates commonalities between the ferret, human, and mouse genomes, 52 (18.5%) of these 266 DEGs in ferrets were common to all three species, whereas 77 (28.9%) were also differentially expressed in humans, and 90 (33.8%) were differentially expressed in mice (Top 25 listed in Table 2; Complete list in Additional file 5: Table S4). Interestingly, 25 DEGs were differentially expressed in human and ferret, but not mice (Table 3); among these genes uniquely modeled in ferrets were those previously determined to be relevant to COPD pathogenesis, with upregulation observed for IL-18, KL, SRXN1, TREM1 and downregulation observed for HAS1 and PNPLA3 [33–36]. Of note, the change in KL and PLPLA3 expression in ferret were discordant with changes in humans found by Bosse et al. Among the 25 most statistically significant DEGs common in ferret and human, three other genes also showed discordant regulation between the two species: GZMA, SNAP25, and EGFLAM (Table 3). These genes are important in airway goblet cells, although studies of its expression and localization in ferrets have not been performed.

### RT-PCR validation analysis of key genes in COPD ferret model

To validate RNA-Seq results, we conducted RT-PCR using ferret tissues for selected DEGs (Fig. 5) that were

### Table 1 KEGG pathway analysis showing up-regulated (A) and down-regulated (B) pathways

| (A) Up-regulated genes—KEGG | Gene set | Description | Size | Overlap | Expect | ER | p-value | FDR |
|----------------------------|----------|-------------|------|---------|--------|----|---------|-----|
| hsa04145                   | Phagosome |             | 152  | 10      | 1.91   | 5.23| 1.89E−05| 0.004|
| hsa04142                   | Lysosome  |             | 123  | 9       | 1.55   | 5.81| 2.18E−05| 0.004|
| hsa04610                   | Complement and coagulation cascades | | 79   | 5       | 0.99   | 5.03| 0.003070148| 0.250|
| hsa05144                   | Malaria   |             | 49   | 4       | 0.62   | 6.49| 0.003233764| 0.250|
| hsa04611                   | Platelet activation | | 123  | 6       | 1.55   | 3.88| 0.004416005| 0.250|
| hsa04380                   | Osteoclast differentiation | | 128  | 6       | 1.61   | 3.72| 0.005361306| 0.250|
| hsa05323                   | Rheumatoid arthritis | | 90   | 5       | 1.13   | 4.41| 0.005373572| 0.250|
| hsa04210                   | Apoptosis  |             | 136  | 6       | 1.71   | 3.51| 0.007171521| 0.266|
| hsa04640                   | Hematopoietic cell lineage | | 97   | 5       | 1.22   | 4.10| 0.007356998| 0.266|
| hsa04060                   | Cytokine-cytokine receptor interaction | | 294  | 9       | 3.70   | 2.43| 0.011381961| 0.347|

| (B) Down-regulated genes—KEGG | Gene set | Description | Size | Overlap | Expect | ER | p-value | FDR |
|----------------------------|----------|-------------|------|---------|--------|----|---------|-----|
| hsa04060                   | Cytokine-cytokine receptor interaction | 294  | 8       | 1.30   | 6.16 | 3.08E−05| 0.01|
| hsa04080                   | Neuroactive ligand-receptor interaction | 277  | 6       | 1.22   | 4.90 | 0.001172172| 0.19|
| hsa04930                   | Type II diabetes mellitus | 46   | 2       | 0.20   | 9.84 | 0.017356504| 1.00|
| hsa04672                   | Intestinal immune network for IgA production | 49   | 2       | 0.22   | 9.24 | 0.01955958| 1.00|
| hsa03320                   | PPAR signaling pathway | 74   | 2       | 0.33   | 6.12 | 0.041960528| 1.00|
| hsa04020                   | Calcium signaling pathway | 183  | 3       | 0.81   | 3.71 | 0.046110736| 1.00|
| hsa04052                   | Chemokine signaling pathway | 189  | 3       | 0.84   | 3.59 | 0.049930188| 1.00|
| hsa04350                   | TGF-beta signaling pathway | 84   | 2       | 0.37   | 5.39 | 0.052704308| 1.00|
| hsa04974                   | Protein digestion and absorption | 90   | 2       | 0.40   | 5.03 | 0.059574237| 1.00|
| hsa04750                   | Inflammatory mediator regulation of TRP channels | 99   | 2       | 0.44   | 4.57 | 0.07042317| 1.00|

**Top ten gene sets annotated are displayed**

**ER enrichment ratio; FDR False Discovery Rate**
uniquely modeled in ferrets, in addition to CYP1B1 as an exposure control, MMP12 as an important gene associated with COPD progression, and HAS1 as a disease of interest related to mucosal defense. Quantification of mRNA expression in smoke-exposed vs. air control ferrets by RT-PCR demonstrated expected changes with smoke exposure (Fig. 5), generally consistent with the RNA-seq analysis and providing validation for the utility of COPD ferrets in modeling a set of genes that can uniquely inform on the pathogenesis of COPD.

Discussion
Various studies using cigarette smoke induced animal models have demonstrated genes associated with smoking by transcriptomic analysis [4, 5]. However, comparative studies using smoke exposed mouse models of COPD have limitations, since mice do not exhibit all features of the complex human COPD sub-phenotypes. In this study, we assessed the transcriptome profile of lung parenchyma in a ferret model of cigarette smoke-induced COPD. Gene expression profiling has not been previously reported in COPD ferrets, which uniquely exhibit features of airway remodeling in comparison to rodent models of cigarette smoke exposure, while also exhibiting emphysema [10–13]. A total of 420 DEGs associated with smoke exposure were identified, many of which have been observed in humans, mice, or both. Furthermore, building upon a recent study by Obeidat et al. who compared the expression profiles of smoke exposed (both short term and long term) murine model with COPD human lungs [6], we identified multiple additional DEGs and associated pathways bearing pathophysiological relevance to COPD that were uniquely expressed in ferrets and humans, but not mice. To our knowledge, this is currently the only RNA-seq based transcriptomics analysis in the COPD ferret model, and due to the unique features of ferrets, provides opportunities for detection of novel pathways with human relevance.

A major goal of the analysis was to identify gene targets that might be uniquely modeled in ferrets, given the human-like phenotype it exhibits. Our analysis revealed 25 aberrantly regulated genes that were exclusive to humans and the ferret COPD model, but not mice. These included several DEGs that have previously been studied for their role in COPD and that may be implicated in mucus-related aspects of the disease. IL-18 has been correlated with pulmonary function in COPD patients and, in an IL-18 transgenic mouse model, was identified as a key driver regulating downstream responses including mucus metaplasia, airway fibrosis, and vascular remodeling [38]. KL-deficient mice have been observed to exhibit delayed mucociliary clearance [34], and downregulation of KL protein KLOTHO has been widely observed in COPD with cigarette exposure [39, 40]. That we observed increased expression of KL may indicate that smoke exposed ferrets are responding to delayed mucociliary clearance, as recently shown [34]. The discordant expression of KL in ferrets compared to humans could be due to the possibility that higher expression of KL acts as a compensatory response to oxidative stress due to smoking, a finding reported by other studies [41], including in human smokers without COPD [42]. Other DEGs common to humans and ferret of interest included the antioxidant enzyme sulfiredoxin-1 (SRX1), which was shown to be upregulated in the lungs of nuclear factor erythroid-2-related factor 2 (Nrf2+) mice upon acute exposure to cigarette smoke, conferring a protective effect against oxidative stress, although was decreased in lung tissue from patients with advanced COPD [43]. This may have been prominent since ferrets exhibit a mild form of disease, given their age and relative duration of cigarette smoke exposure. Upregulation of triggering receptor expressed on myeloid cells 1 (TREM1) has been observed in cigarette-smoke mice, exacerbating pyroptosis-mediated lung injury and inflammation [44]. These pathways could represent opportunities for further investigation in COPD pathogenesis using the ferret model.

Prior investigations evaluating gene expression profiles of smoked-exposed mouse [6] and human COPD [14] lungs enabled cross-species transcriptomics analysis for

![See figure on next page.](image-url)

**Fig. 4** A Venn diagram depicting overlap between differentially expressed genes (DEGs) in smoke-exposed ferret, human COPD, and smoke-exposed murine lungs. The number of DEGs in common between smoke-exposed ferrets, human COPD [14], and smoke-exposed mice [6] are shown. B Circos plot of smoking-related genes overlapping between smoke-exposed ferret, human COPD, and smoke-exposed murine lungs. The genochromosomal positions are based on the human genome and shown inside the circles. The first pair of circles (light blue and dark blue) represents the dysregulated genes from the long term (24 weeks) smoke-exposed mouse [6]. The upregulated genes are plotted in the dark blue circle pointing outward whilst the downregulated genes are plotted in the light blue circle pointing inward. The second pair of circles (light red and dark red) show the ferret genes dysregulated due to smoke exposure. The upregulated genes are in the dark red circle and the downregulated genes are in the light red circle. The third pair of circles (light green and dark green) show the “human smoking” gene signature [6, 14]. The upregulated genes are in the dark green circle, whilst the downregulated genes are in the light green circle. The length of the line is proportional to the $-\log_{10}$ FDR values for differential expression in human and for the $-\log_{10}$ FDR values in murine and for the $-\log_{10} p_{\text{adj}}$ for the ferret. Genes highlighted are statistically significant from the respective studies.
Fig. 4  (See legend on previous page.)
Table 2 The top 25 smoke related DEGs identified in the ferret, mouse and human lungs shown in order of statistically significance for the ferret study

| FERRET | log₂FC | p-value | P_adj | MOUSE | log₂FC | p-value | P_adj | HUMAN | Wilcox p-value | Fold change |
|---------|--------|---------|-------|--------|--------|---------|-------|-------|----------------|-------------|
| CD84    | 2.22   | 4.64E−56| 3.81E−52| CD84  | 1.40   | 2.11E−13| 1.01E−10| CD84  | 2.14E−09      | 1.40        |
| CYP1B1  | 2.33   | 5.85E−48| 2.41E−44| CYP1B1| 0.89   | 9.33E−11| 1.28E−08| CYP1B1| 1.14E−19      | 4.53        |
| ATP6V0D2| 3.99   | 1.3E−09 | 3.42E−26| ATP6V0D2| 1.04  | 1.11E−06| 3.18E−05| ATP6V0D2| 7.64E−16      | 5.89        |
| PLD3     | 1.30   | 2.55E−25| 2.62E−25| PLD3  | 1.23   | 8.27E−17| 4.37E−13| PLD3  | 1.17E−07      | 1.32        |
| CD180    | 1.63   | 1.4E−27 | 1.19E−24| CD180 | 0.39   | 4.42E−03| 2.42E−02| CD180 | 1.66E−08      | 1.59        |
| IGSF6    | 1.14   | 2.64E−25| 1.89E−22| IGSF6| 0.37   | 1.90E−03| 1.24E−02| IGSF6| 5.24E−14      | 1.96        |
| TYROBP   | 1.57   | 3.37E−25| 2.31E−22| TYROBP| 0.77   | 3.38E−08| 1.85E−06| TYROBP| 7.77E−12      | 1.37        |
| P2RX4    | 1.04   | 3.09E−24| 1.89E−21| P2RX4| 0.30   | 3.68E−05| 5.37E−04| P2RX4| 3.39E−10      | 1.36        |

The statistical metrics reported for the mouse and human are computed from the mouse 24 week smoke exposed study [6] and the human smoking study [14] respectively. The log₂FC is log fold change; the p_adj is adjusted p-value.

266 of the 420 total DEGs detected in the COPD ferret model. There was substantial overlap in DEGs detected across all three species, with involvement seen in genes with widely studied roles in COPD such as CYP1B1 [31], matrix metalloproteinase 12 (MMP12) [45], and aryl hydrocarbon receptor repressor (AHRR) [46], underscoring the validity of the smoke exposure model in ferrets for evaluation of the lung transcriptome. Common DEGs between ferret, human, and mouse also reinforced the contribution of additional genes to COPD pathogenesis. Among other overlapping DEGs, TREM2 has been observed to promote macrophage survival in the lungs [47], nimA-related protein kinase 6 (NEK6) has been implicated in COPD-associated ciliopathies [48], and cathepsin B (CTSB) induced emphysema and secretory cell hyperplasia upon intratracheal administration to hamster lungs [49]. Additional genes of interest that were substantially upregulated and have direct relevance to COPD and/or cigarette smoke exposure included SERPINA1 (2.17 log₂ fold change; log₂FC; p_adj = 5.13E−26), ATP6V0D2 (3.99 log₂FC; p_adj = 3.42E−26), MMP-12 (2.69 log₂FC; p_adj = 6.20E−12), AHRR (1.74 log₂FC; p_adj = 8.32E−05), and CTSB (1.83 log₂FC; p_adj = 4.64E−18), among others [4, 50].

Downregulation of hyaluronic acid synthase 1 (HAS1) has been associated with abnormally low production of the high molecular weight form of HA (HMW-HA) in COPD [51]. While RNA-Seq revealed a trend, RT-PCR, conducted because of prior interest in this area, indicated downregulation of HAS1. Reduced HAS1, and thus lower HMW-HA, could also lead to abnormal HMW/LMW HA ratios, which would alter mucus clearance and also induce inflammation [51].

Multiple cellular pathways were aberrantly altered in the COPD ferret model, including pathways that are also associated with COPD pathogenesis in humans. Of the pathways were significant changes were observed (Table 1 and Additional file 4: Table S3), those that have been reported to be involved with COPD pathogenesis included activation of mitogen-activated protein kinase (MAPK), phosphatidylinositol 3 kinase/serine threonine protein kinase (PI3K-AKT), phagosome,
Table 3  25 Smoke-related DEGs identified in the ferret and human lungs shown in order of statistically significance for the ferret study

| FERRET  | \( \log_{2}FC \) | p-value       | \( p_{adj} \) | Human  | Wilcox p-value | Fold change |
|----------|-----------------|---------------|---------------|--------|----------------|-------------|
| DHRS9    | 4.62            | 2.31E-66      | 3.80E-61      | DHRS9  | 3.38E-08       | 1.70        |
| LIPA     | 1.62            | 3.60E-35      | 7.41E-32      | LIPA   | 1.40E-12       | 2.16        |
| RAB7B    | 1.33            | 1.63E-30      | 2.43E-27      | RAB7B  | 8.24E-09       | 1.52        |
| TMEM92   | 1.30            | 1.38E-20      | 4.73E-18      | TMEM92 | 3.28E-10       | 1.48        |
| HSD17B14 | 1.37            | 1.26E-19      | 3.84E-17      | HSD17B14 | 1.51E-12     | 1.69        |
| CSAR1    | 1.17            | 1.42E-16      | 3.25E-14      | CSAR1  | 4.37E-12       | 1.42        |
| WDFY4    | 1.21            | 3.60E-16      | 7.90E-14      | WDFY4  | 1.44E-07       | 1.29        |
| IL18     | 1.03            | 5.78E-15      | 1.03E-12      | IL18   | 2.26E-10       | 1.42        |
| ALPL     | 1.33            | 2.34E-14      | 4.04E-12      | ALPL   | 2.71E-09       | 1.39        |
| CLEC7A   | 1.39            | 4.59E-14      | 7.62E-12      | CLEC7A | 2.62E-12       | 1.35        |
| TMEM144  | 1.06            | 1.63E-13      | 2.50E-11      | TMEM144 | 4.05E-11     | 1.38        |
| TREM1    | 1.12            | 1.38E-12      | 1.84E-10      | TREM1  | 2.24E-08       | 1.41        |
| KBTBD8   | 1.31            | 1.14E-09      | 9.94E-08      | KBTBD8 | 2.97E-09       | 1.61        |
| GLDN     | 1.46            | 2.83E-07      | 1.57E-05      | GLDN   | 2.28E-10       | 1.44        |
| SRXN1    | 1.02            | 6.73E-07      | 3.38E-05      | SRXN1  | 5.16E-08       | 1.36        |
| GLS2     | 1.09            | 1.04E-06      | 4.97E-05      | GLS2   | 2.46E-12       | 1.54        |
| GZMA     | 1.81            | 1.36E-06      | 6.24E-05      | GZMA   | 5.53E-08       | 0.65        |
| RPP25    | 2.07            | 3.87E-06      | 1.61E-04      | RPP25  | 4.64E-09       | 1.33        |
| SNAP25   | – 3.94          | 5.62E-05      | 1.63E-03      | SNAP25 | 2.56E-09       | 2.10        |
| KL       | 1.23            | 7.98E-04      | 1.43E-02      | KL     | 1.80E-08       | 0.62        |
| INSC     | – 1.36          | 2.44E-03      | 3.37E-02      | INSC   | 1.75E-07       | 0.72        |
| PNPLA3   | – 1.03          | 2.52E-03      | 3.46E-02      | PNPLA3 | 7.50E-13       | 2.34        |
| EGFLAM   | – 1.15          | 2.75E-03      | 3.69E-02      | EGFLAM | 1.01E-06      | 1.52        |
| MAL      | – 1.17          | 9.75E-03      | 9.24E-02      | MAL    | 2.72E-07       | 0.73        |
| FANCE    | 1.07            | 1.07E-02      | 9.85E-02      | FANCE  | 8.44E-11       | 1.30        |

The statistical metrics reported for the human are computed from the human smoking study [14]. The \( \log_{2}FC \) is \( \log_{2} \) fold change; the \( p_{adj} \) is adjusted p-value. Genes listed in bold are discordant between ferret and human.

cytokine-cytokine receptor interaction, extracellular matrix (ECM)-receptor interaction, chemokines, apoptotic signaling pathways, inhibition of transforming growth factor beta (TGF-ß), peroxisome proliferator-activated receptors (PPAR), and inflammatory mediator regulation of transient receptor potential (TRP) channel pathways [52].

Our study, while unique in the analysis of ferrets, is not without limitations. Ferrets are an outbred species, and thus genetic heterogeneity may have reduced the ability to detect some important DEGs, especially given the sample size encountered. We focused our analysis on parenchyma tissue, noting this was most similar to the specimens collected in humans and other species for which we compared, and this samples also include the small airways, where airway disease is prominent [12, 53]; nevertheless, the analysis of airway epithelia could be informative in the future, especially given pathological changes in the large airways of ferrets upon smoking [10–12]. Since ferrets are sexually dimorphic, it is not surprising that there were important differences by ferret sex, although this was controlled for in the differential gene analysis and included where possible in comparison to other data sets. Importantly, analysis of RNA has evolved throughout the study, providing a limitation in the inter-species comparisons. The RNA-Seq analysis and RT-PCR validation in this study was based on mRNA expression; validation by protein analysis will be useful in the future, noting some protein reagents in ferrets need further characterization. To assure specificity of our comparisons, we required at least one-fold change, in addition to statistical significance for the DEG to be included in the ferret study, which differed from those included in mice-human comparators. In a minority of cases for genes uniquely modeled in ferrets, the directional change in differential expression in ferrets and humans were discordant. Although this can be explained for KL due to the potential for evolving changes over time, further research will be needed for other genes where this was the case, such as GZMA, SNAP25, KL, PNPLA3 and EGFLAM.
Fig. 5  RT-PCR validation of differentially expressed genes (DEGs) unique to smoke-exposed ferret and human COPD lungs as compared to raw data from RNA-Seq. RT-PCR (right) and RNA-Seq (left) measurements for A CYP1B1; B MMP12; C IL-18; D KL; E TREM1; F TREM144; G SFRN1; H ATP6V0D2; I PNPLA3; and J HAS1 in smoke-exposed and air control ferret lungs. Mean ± SEM. RT PCR N = 6/condition, 3 males and 3 females per group, comparison by Mann–Whitney
Conclusion

In summary, our analysis provides a large RNA-seq based transcriptome analysis set in COPD and involves a ferret model that uniquely exhibits features of chronic bronchitis. Our data revealed 420 DEGs, including multiple genes established as abnormal among COPD patients, such as pathways involving neutrophil activation, immune regulation, and oxidative metabolism. We also identified 25 DEGs that are uniquely expressed in ferrets and humans, but not mice, some of which implicate abnormal mucus clearance, a finding uniquely modeled by ferrets. Selected genes were validated with real time PCR, and these were concordant with the RNA-Seq expression. Together, the transcriptomic profile and comparison with other available models of COPD identifies a unique set of genes that could be relevant to disease pathogenesis. This high-resolution view of the lung transcriptome associated with COPD may ultimately provide a valuable resource for the research community and warrants further mechanistic investigations.

Abbreviations

COPD: Chronic obstructive pulmonary disease; qPCR: Quantitative polymerase chain reaction; DEGs: Differentially expressed genes; KEGG: Kyoto Encyclopedia of Genes and Genomes; CYP1B1: Cytochrome P450 family 1 subfamily B member 1; GO: Gene Ontology; BP: Biological process; MF: Molecular functions; KL: KLOTHO; SRX1: Sulfiredoxin-1; TREM1: Triggering receptor expressed on myeloid cells 1; Nrf: Nuclear factor erythroid-2-related factor 2; MMP12: Matrix metalloproteinase 12; AHRR: Aryl hydrocarbon receptor repressor; NEK6: NIMA-related protein kinase 6; HAS1: Hyaluronic acid synthase 1; HMW-HA: High molecular weight form of HA; MAPK: Mitogen-activated protein kinase;
Supplementary Information
The online version contains supplementary material available at https://doi.org/10.1186/s12931-022-02198-0.

Additional file 1: Table S1. RNA-seq data for 12 samples (six biological replicates each for air and smoke experimental conditions) sequenced using an Illumina NextSeq 500 System. [GEO Accession Number GSE193749].

Additional file 2: Fig. S1. The proportion of paired ends reads for the ferret lung samples, by individual ferrets.

Additional file 3: Table S2. The number of paired ends reads for the ferret lung samples.

Additional file 4: Table S3. Complete annotation GO TERMS in smoke exposed ferrets compared to controls.

Additional file 5: Table S4. Complete list (52) of DEGs in common exposed ferrets compared to controls.

Acknowledgements
Not applicable.

Author contributions
SSH, EMS; SRR conceived of and designed the research; SSH, YJKE, DSM, DDS, and SMR analyzed data and interpreted results; SSH, YJKE, EFL, and SMR prepared figures; SSH, YJKE, EFL, and SMR wrote the manuscript; YJKE, SMR supervised the project; all authors had an opportunity to edit the manuscript and approved of its submission. All authors read and approved the final manuscript.

Funding
These studies were supported by NIH National Heart, Lung, and Blood Institute (NHLBI) Grant R3S HL135816 (to S.M.R.) and National Institute of Diabetes and Digestive and Kidney Diseases Grant P30 DK072482 (to S.M.R.). Funded in part by a grant from the Alpha-1 Foundation 919960 (To S.S.H).

Availability of data and materials
The datasets analyzed during the current study are available from the corresponding author on reasonable request.

Declarations
Ethics approval and consent to participate
All animal experiments were approved by the Institutional Biosafety Committee of the University of Alabama at Birmingham (IACUC 20232).

Consent for publication
Not applicable.

Competing interests
The author(s) declare(s) that they have no competing interests.

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Received: 14 June 2022 Accepted: 19 September 2022
Published online: 10 October 2022

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