Exposure to extracellular vesicles from *Pseudomonas aeruginosa* result in loss of DNA methylation at enhancer and DNase hypersensitive site regions in lung macrophages

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**ABSTRACT**

Various pathogens use differing strategies to evade host immune response including modulating the host’s epigenome. Here, we investigate if EVs secreted from *P. aeruginosa* alter DNA methylation in human lung macrophages, thereby potentially contributing to a dysfunctional innate immune response. Using a genome-wide DNA methylation approach, we demonstrate that *P. aeruginosa* EVs alter certain host cell DNA methylation patterns. We identified 1,185 differentially methylated CpGs (FDR < 0.05), which were significantly enriched for distal DNA regulatory elements including enhancer regions and DNase hypersensitive sites. Notably, all but one of the 1,185 differentially methylated CpGs were hypomethylated in association with EV exposure. Similarly, hypomethylated CpGs tracked to genes including *AXL, CFB* and *CCL23*. Gene expression analysis identified 310 genes exhibiting significantly altered expression 48 hours post *P. aeruginosa* EV treatment, with 75 different genes upregulated and 235 genes downregulated. Some CpGs associated with cytokines such as *CSF3* displayed strong negative correlations between DNA methylation and gene expression. Our infection model illustrates how secreted products (EVs) from bacteria can alter DNA methylation of the host epigenome. Changes in DNA methylation in distal DNA regulatory regions in turn can modulate cellular gene expression and potential downstream cellular processes.

**Background**

Extracellular vesicles (EVs) are circular lipid bilayer particles, 20–1000 nm in diameter, secreted by nearly all known eukaryotic and prokaryotic cell types [1]. EVs include eukaryotic exosomes (20–150 nm) and microvesicles (100–1000 nm), which contain a variety of macromolecules including RNA, DNA and proteins, as well as outer membrane vesicles (OMVs) (20–500 nm) that contain virulence and other factors [1]. EVs transport these virulence factors to facilitate infection and can trigger inflammatory responses in host cells [2].

Multiple studies have characterized the contents of *P. aeruginosa* extracellular vesicles. Some examples of proteins include OprF, OprG, IcmP and LasA which have roles in antibiotic resistance, proteolysis and bacteria-host interactions [3]. *P. aeruginosa* EVs also carry immune modulatory components like lipopolysaccharide (LPS) that may affect cells that express pattern recognition receptors [1,4]. After being released from the bacteria, EVs from other Gram-negative bacteria can exist in natural environments like water drains, soil, and house dust [1]. Furthermore, Gram-negative EVs are pathogenic on their own outside of the live bacteria [5,6]. To better understand the mechanism behind pathogenic effects of bacteria in the lung, it is important to study the effects of isolated EVs on host immune cells. Such studies will contribute to understanding the bacterial effects of
diseases such as cystic fibrosis (CF), in which pathogenic species like *P. aeruginosa* deliver virulence factors by secreting EVs into the lung and accelerates decline the pulmonary function in patients [7–11].

Although some work has been done to understand the impact of bacterial EVs in gene expression of host cells, not as much has been done to discern epigenetic responses to bacterial EVs. Epigenetics is the study of heritable changes in gene function caused by mechanisms other than changes in the underlying DNA sequence [12]. One of the most widely studied epigenetic modifications is DNA methylation of cytosine at CpG dinucleotides [13]. CpG islands, regions of dense CpG contents, are predominantly located in promoter regions of genes [14,15]. Methylation of CpG regions regulates gene expression by recruiting proteins involved in gene repression or by inhibiting the binding of transcription factor(s) to DNA [16]. DNA methylation controls methylation of histone H3K4 at regulatory regions, where hypomethylated regulatory regions such as promoters and putative enhancers have enriched with H3K4me3 marks thereby increasing active transcription [17]. Additionally, DNA methylation at enhancer regions can present bivalent chromatin states as it can co-exist with active H3K27ac [18]. When DNA methylation marks decrease, the H3K27ac marks are also reduced, decreasing active transcription [18]. The role of DNA methylation and its loss at DNase hypersensitive sites (DHS) is less studied. However, it is evident that DHS regions are depleted in areas of dense CpGs [19]. Historically, alterations in DNA methylation have been attributed to chronic or longer-term environmental exposures, however, recent work has begun to investigate whether shorter-term exposures also result in DNA methylation changes.

Recent studies have investigated how pathogens can modulate host epigenome, specifically the DNA methylation landscape of cells of the innate immune system. Sinclair et al. have shown genome-wide DNA methylation changes in *Anaplasma phagocytophilum*-infected human neutrophils [20]. Cizmeci et al. demonstrated altered DNA methylation associated with *Burkholderia pseudomallei* infection in macrophages [21]. Pacis et al. in an infection model of *Mycobacterium tuberculosis* in human dendritic cells [22], provide evidence of demethylation of distal DNA regulatory elements in genes, some of which regulate the activation of key immune transcription factors. However, epigenetic effects of EVs from *P. aeruginosa*, a common type of bacteria that causes infections, has not yet been studied to our knowledge. Therefore, the goal of this study is to assess whether EVs isolated from *P. aeruginosa* alter DNA methylation in human lung macrophages thereby potentially modulating the ability of macrophages to mount a proper immune response to the infection.

**Results**

**Subject characteristics and EV characterization**

Bronchoalveolar lavage (BAL) fluid was obtained from the right upper lobe of healthy subjects (n = 8) and a pure population of lung macrophages was isolated as previously described [23,24]. A cohort of five females and three males was used for DNA methylation analysis, whereas four females and three males were used in the gene expression analysis, due to lack of RNA availability from one of the original female subjects. Mean age of the study population was 27.1 years old. Subject characteristics are shown in Table 1.

As the mucoid phenotype of the *P. aeruginosa* is the strain associated with increased morbidity and mortality, EVs from clinically-driven mucoid strains were used for exposure [25]. Isolation of EVs was done via affinity resin as described in the Supplemental Methods. Negative stain transmission electron microscopy was used for visualization of EVs (Supplementary Figure 1a). Additionally, nanoparticle tracking analysis was used for particle counting and validation of particle size. EV particle size ranged from 20 to 600 nm. A representative trace confirming particle size from the NanoSight NS300 instrument is shown in Supplemental Figure 1b. To characterize

| Table 1. Subject characteristics. | DNA methylation (n=8) | Nanostring (n=7) |
|----------------------------------|-----------------------|-----------------|
| Age                              | 27.1 (23-31)          | 27.3 (23-31)    |
| Sex                              |                       |                 |
| Female                           | 5 (62.5%)             | 4 (57.1%)       |
| Male                             | 3 (37.5%)             | 3 (42.9%)       |
the contents of *P. aeruginosa* EVs, we performed mass spectrometry on isolated EVs. They were found to contain virulence factors such as LasB and CbpD and protease associated proteins such as PrpL and LasA. Full list of proteins identified from mass spectrometry from *P. aeruginosa* EVs can be found in Supplementary Table 1.

**Differential methylation with *P. aeruginosa* EV treatment**

Unsupervised hierarchical clustering of top 10,000 most variable CpGs resulted in clustering by subject rather than by treatment (Supplementary Figure 2). Clustering by subjects is not unexpected to observe genetic effects even after dropping known SNP-associated array probes as treatment effects of *P. aeruginosa* EVs may not affect majority of the CpGs analysed.

To observe the effects of *P. aeruginosa* EV treatment on DNA methylation of lung macrophages, we compared methylation status of DNA from matched control lung macrophages and *P. aeruginosa* EV treated macrophages. We identified 1,185 CpGs with differential methylation in association with EV treatment (FDR < 0.05, Figure 1(a)). Some of the most differentially methylated CpGs include those associated with genes such as *BCAT1*, *CNIH3*, and *YPEL2*. Remarkably, all but one of the significantly differentially methylated CpGs were hypomethylated in treated lung macrophages compared to the controls. The only hypermethylated CpG was associated with gene *EVL*. Full list of differentially methylated CpGs and their annotations are provided in Supplementary Table 2.

As expected, supervised hierarchical clustering of the 1,185 CpG with differential methylation indicated a clear distinction between control and *P. aeruginosa* EV treated lung macrophages (Figure 1(b)). To better understand where the differentially methylated CpGs were located, we conducted Mantel-Haenszel odds ratio enrichment tests for genomic contexts of the 1,185 CpGs. Interestingly, differentially methylated CpGs were significantly enriched in DNase hypersensitive sites (DHS) (OR = 8.91, 95% CI = 7.04–11.41) and Fantom5 enhancer regions (OR = 7.91, 95% CI = 6.95–8.98) (Figure 2(a)). In addition, differentially methylated CpGs were significantly depleted for gene promoter regions (OR = 0.29, 95% CI = 0.24–0.36). Based on the observed depletion of CpGs at promoter regions, the differentially methylated CpGs were expectedly significantly enriched in the open sea region (OR = 5.61, 95% CI = 4.61–6.88) in relation to known CpG island locations (Figure 2(b)). Similar to the CpG probes used in analysis, the differentially methylated CpGs were most enriched in regions within 200 base pairs (bps) upstream of the transcription start site (TSS) (Supplementary Figure 3). However, the CpGs were enriched in regions 1–2 kbps downstream from the TSS compared to the analysed CpGs, which corresponds to the enrichment of differentially methylated CpGs at gene body sites (OR = 1.34, 95% CI = 1.19–1.51) and introns (OR = 1.94, CI = 1.72–2.18).

**Differential gene expression with *P. aeruginosa* EV treatment**

To further understand effects of *P. aeruginosa* EV treatment on the lung macrophages, we measured gene expression with the NanoString nCounter human v3 Cancer/Immune RNA expression assay. To expand on studies that have measured targeted gene expression of inflammatory response to *P. aeruginosa* EVs, our studies were designed to broaden the scope of the genes measured [1]. Of 541 genes used in differential gene expression analysis, a majority (310) of genes were significantly differentially expressed in association with *P. aeruginosa* EV treatment (Figure 3(a)). There were 3-fold more downregulated genes (235 genes) in the EV-treated lung macrophages compared to upregulated genes (75 genes). Hierarchical clustering of all 541 genes used in downstream analysis indicated a distinction between control lung macrophages and lung macrophages treated with EVs (Figure 3(b)).

Top downregulated genes included genes that encode cell membrane proteins such as *FN1*, *TREM2*, *CD22*, and *CR1*. The most highly upregulated genes included genes that encode for cytokines and chemokines such as *PPBP*, *IL19*, *IL6*, and *CXCL6*. We conducted gene ontology analysis for both downregulated and upregulated genes to specifically assess what biological processes were
associated with directional changes in gene expression. Strikingly, the downregulated genes were associated with biological processes related to regulation of viral processes and MHC class II presentation (Supplementary Figure 4a). Upregulated genes were found to be associated
with biological processes regulating white blood cell migration or chemotaxis (Supplementary Figure 4b). A full list of differentially expressed genes (FDR < 0.05) can be found in Supplementary Table 3.

To gain a deeper understanding of the biological function of these differentially methylated CpGs, we assessed gene ontologies of nearby genes of these CpG sites. Because not all CpGs were annotated to a specific gene, we used the Genomic Regions Enrichment Annotations Tool (GREAT) to predict the functions of the non-coding regions using their nearby genes [26]. Given the relative purity of cell type in our samples, significantly hypomethylated CpGs were expectedly associated with genes involved in active immune response biological processes such as wound healing, positive regulation of angiogenesis and haemopoiesis (Supplementary Figure 5). Wound healing involved in inflammatory response biological process had the highest fold enrichment relative to other biological processes. Further GREAT analysis indicated that these hypomethylated CpGs were associated with cellular components such as PI3K complex and focal adhesion as well as molecular functions such as MAP kinase activity and histone acetyltransferase binding (Supplementary Figure 6A, 6B). Interestingly, gene ontology analysis of 735 genes annotated to differentially methylated CpGs revealed that the CpGs were enriched in positive regulation of phosphorylation and signal transduction processes (Supplementary Figure 7).

**Correlation between changes in methylation and changes in gene expression**

We next sought to assess the potential relationship between CpG hypomethylation and downstream gene expression in lung macrophages treated with *P. aeruginosa* EVs. The 1,185 differentially methylated CpGs tracked to 565 genes. Of these
565 genes, 38 unique genes had matching NanoString gene expression data. As some CpGs tracked to the same genes on the NanoString gene expression panel, we were able to assess correlations between 57 CpGs and their respective gene expression levels. 31 of the 57 CpGs showed significant negative correlation and four CpGs positive correlation between methylation and gene expression, respectively, while 22 did not have significant correlation at the 0.05 significance.
level. Complete results for the correlation of gene expression versus DNA methylation is available in Supplementary Table 4.

One particular gene that is relevant to macrophages, CSF3, had four associated CpGs that were hypomethylated. CSF3 encodes a cytokine that is essential in controlling differentiation and function of granulocytes [27]. Spearman rank correlation indicated that hypomethylation was significantly associated with increased expression of CSF3 for all four CpGs which are located upstream of the transcription start site (all \( P < 0.01 \), Figure 4(a–d)). The CpG site (cg07472953) farthest upstream of the transcription start site was still within 1.5kb, and also tracked to a Fantom5 enhancer region (Figure 4(a–e)). The other three hypomethylated CpGs were the only CpGs measured on the array platform within 200 bps of the transcription start site. All four hypomethylated CpGs were located within the same DHS and were not in a CpG-dense CpG island. Full annotation of all CSF3 associated CpGs are included in Supplementary Table 5.

Among the remaining CpGs tested, we observed four CpGs with positive correlations between methylation and gene expression at three different
genes. Decreased methylation of CpGs related to *P. aeruginosa* EV treatment was significantly associated with decreased gene expression levels of *AXL*, *MME*, and *CD46* (Figure 5(a–b)). All four CpGs tracked to the gene body regions of these genes, and the observed association of gene body hypermethylation with increased gene expression is consistent with prior literature [28–30]. Three CpGs except cg15815551 (Figure 5D) were located in either shelf or shore region in relation to CpG island. Full annotation of these CpGs are highlighted in light green in Supplementary Table 2.

**Discussion**

The goal of this study was to determine if EVs from *P. aeruginosa* could alter the host epigenome. Although prior work has examined epigenomic alterations related to bacterial infection [20–22], our study addresses a gap in understanding the impact of *P. aeruginosa* EV exposure on host immune cell epigenetic gene regulation. Using human lung macrophages and *P. aeruginosa* extracellular vesicles as an *in vitro* model, we demonstrate that macrophages exposed to *P. aeruginosa* EVs exhibit decreased DNA methylation at known gene regulatory elements and that these changes are associated with altered gene expression.

We observed a broadly consistent pattern of DNA hypomethylation associated with *P. aeruginosa* EV exposure that tracked to *cis*-regulatory regions of biological processes important in active immune function such as wound healing and haematopoiesis. What was more interesting is that the ontology of only the genes annotated to CpGs revealed that genes with active phosphorylation processes were enriched. This suggests that the lung macrophages activate innate immune pathways in response to *P. aeruginosa* EVs. Additionally, the hypomethylated CpGs...
were located in distal regulatory elements such as enhancers and DNase hypersensitive sites and depleted in promoter regions. It is possible that hypomethylation of these distal regulatory elements may be changing the chromatin conformation to poise the lung macrophages to be prepared for active transcription, but not directly controlling active transcription. Few CpGs in HDAC9 and TET2, genes important in transcriptional regulation, were hypomethylated. It suggests EVs may potentially activate master epigenetic regulatory genes to induce a response in the lung macrophages. The significant correlations observed in a majority of the assessed relationships between methylation and gene expression indicate the possibility of epigenetic mechanisms contributing to modulation of immune responses to *P. aeruginosa* EVs in lung macrophages.

Our investigation is consistent with the study of Pacis et al. demonstrating DNA methylation changes in response to bacterial infection (i.e. *M. tuberculosis*) in both human dendritic cells and monocyte-derived macrophages in several key observations [31]. First, our results overwhelmingly demonstrate DNA hypomethylation in our EV infection model. Second, our study shows significant enrichment of differentially methylated CpGs in enhancer regions. And finally, differential methylation was noted to overlap in numerous genes in both studies, including: *NFKB1*, *CREB5*, and *BCL2*.

*NFKB1*, *CREB5*, and *BCL2* have important functions in innate immunity in response to infections. In macrophages, the NF-κB family has essential pro-inflammatory functions in response to infections. NF-κB transcription factors induce large number of cytokines and chemokines such as IL-1β and IL-6 [32]. We observed hypomethylation of CpG in the 5′ UTR region of *NFKB1* and substantial increase in expression of *IL1B* and *IL6*. NF-κB also regulates *BCL2* to promote cell survival. The CREB family is also involved in anti-apoptotic cell survival in macrophages through NF-κB-dependent pathway [33]. Hypomethylation of these critical genes may contribute to survival of macrophages in response to pathogens like *M. tuberculosis* and *P. aeruginosa*. Further investigation is needed to validate the involvement of these genes in an active *P. aeruginosa* infection.

However, not all studies focusing on host DNA methylation changes in the setting of infection demonstrate consistent results. This suggests that methylation events may be pathogen-specific. While Cizmeci et al. observed DNA hypomethylation in a human macrophage cell line upon exposure to *Burkholderia pseudomallei* consistent with our results, the work of Marr et al. demonstrates DNA hypermethylation events associated with *Leishmania donovani* infection of THP-1 macrophages [21,34,35–39]. Additional infection models and testing of alternate disease-related pathogens or pathogen-derived secreted products are required to better understand DNA alterations in host-pathogen interactions.

*P. aeruginosa* infection in cystic fibrosis patients is common and leads to poor patient outcomes. Understanding the molecular mechanisms underlying immune response to this opportunistic pathogen could reveal novel strategies to combat chronic infections. Our study reports epigenetic and transcriptional reprogramming of lung macrophages from exposure to *P. aeruginosa* extracellular vesicles. Future work examining epigenetic reprogramming of lung macrophages in CF patients with active *P. aeruginosa* infection as well as other CF pathogens, either singularly or co-colonized is needed to validate our lung infection model observations. With further studies in immune response to bacterial pathogens in CF, opportunities to develop better options for preventing and treating infections in CF and other immunocompromised patients may be gleaned.

**Conclusions**

In this study, we have demonstrated extensive epigenetic changes in DNA methylation, primarily hypomethylation in enhancer and DNase hypersensitive regions and gene body regions in host lung macrophages upon exposure to *P. aeruginosa* EVs. Consistent with changes in these distal DNA regulatory elements, we observed coinciding changes in gene expression. These results are consistent with an immune evasion strategy employed by a prokaryote in a host-pathogen interaction,
potentially leading to an altered innate immune response.

**Methods**

This study was approved by the Dartmouth Hitchcock Institutional Review Board (#22,781). All subjects were healthy non-smokers. Eight volunteer subjects, all Caucasian, with a mean age of 27.1 (range 23–31, 3 males and 5 females) were included in the study and specimens from 7 subjects were amenable for use in the gene expression portion of this study (3 males and 4 females).

**Bronchoalveolar lavage and macrophage isolation**

Subjects underwent flexible bronchoscopy following local anaesthesia with lidocaine to the posterior pharynx and intravenous sedation. A bronchoscope was inserted trans-orally and advanced through the vocal cords. Bronchoalveolar lavage (BAL) fluid was obtained from tertiary airways via rinses with 20 ml of sterile saline followed by 10 ml of air and repeated for a total of 5 times per airway. Lung macrophages were isolated as previously described [23,40].

**Experimental design**

Right upper lobe (RUL) lung macrophages were used in this study, as RUL is the predominant lung lobe infected in CF patients and our previous work has shown that regional differences occur in lung macrophages [24,41]. Cells were plated in 6-well tissue culture dishes seeded at 1 x 10^6 cells per well. Cells were allowed to set down for 60 minutes. Lung macrophages were allowed to set down for 60 minutes. Cells were then incubated in the presence or absence (control) of *P. aeruginosa* EVs at a ratio of 100:1 (EVs:cell) per Cecil et al., for up to 48 hours. We have used the 48-h time point to simulate chronic infection [42].

The *P. aeruginosa* clinical isolate strain used to generate EVs in this study is designated DH1137 and was a gift from Dr. Deborah Hogan, Department of Microbiology and Immunology, Geisel School of Medicine at Dartmouth. The isolate was originally described as strain NC-AMT0101-2 by Smith et al [43]. Additionally, *P. aeruginosa* strain NC-AMT0101-2 has been characterized as LasR sufficient and mucoid phenotype by the work of Hoffman et al [44]. For EV isolation and characterization details see Appendix 1: Supplemental methods.

**DNA methylation array**

Epigenome-wide DNA methylation profiling was performed via the Infinium Methylation EPIC Bead Chips (Illumina Inc., San Diego, CA) to measure methylation at more than 850,000 CpG sites as previously described [45]. Briefly, DNA was extracted from bronchoalveolar lavage-derived cells via Qiagen (Germantown, MD) DNeasy Blood and Tissue Kit. DNA was quantitated on a Qubit 3.0 Fluorometer (Life Technologies, Carlsbad, CA). Bisulphite conversion of DNA was carried out with the Zymo EZ DNA methylation kit (Zymo Research, Irvine, CA) and EPIC array hybridization and scanning were performed at the University of Southern California Molecular Genomics Core.

**DNA methylation array data processing**

Raw intensity data files (IDATs) from the MethylationEPIC BeadChips were processed by the minfi R/Bioconductor analysis pipeline (version 1.30) annotation file version ilm10b4.hg19 [46]. 78,379 cross reactive and SNP-associated probes annotated by Zhou, et al and 19,064 probes on sex chromosomes were excluded [47]. The quality control was performed using ENmix R package [48]. 9,432 probes those failing to meet a detection P-value of 0.00001 in ≥ 10% samples, were excluded. After all exclusions, 759,003 CpGs were remained for analysis. The filtered data was then normalized using preprocessFunnorm in minfi to remove any unwanted technical variation.

Annotations of CpGs such as genomic context or relation to CpG Island were provided in the Illumina EPIC B4 manifest file. The ‘enhancer’ context was defined by having a FANTOM5 enhancer record. ‘DNase hypersensitive site’ context was defined by having a record in the ‘DNase_Hypersensitive_NAME’ in the annotation.
‘Gene body’ transcriprional context was defined by having a ‘Body’ in the UCSC_RefGene_Group. Likewise, ‘3’ UTR and ‘5’ UTR were defined by having ‘UTR3’ and ‘UTR5’, respectively, in the UCSC_RefGene_Group. ‘Promoter’, ‘Intergenic’, ‘Intron’ and ‘Exon’ genomic contexts were defined by finding overlapping genomic regions of the CpGs and each context using the ilm10b4.hg19 annotation. Relation to CpG Island were defined by the ‘Relation_to_UCSC_CpG_Island’ in the Illumina EPIC annotation file. If no record of relation to the CpG island was indicated, the CpG was considered to be in the ‘Open Sea’ region.

**Differential methylation analysis**

Based on the prior analysis of distribution of the methylation beta value variance for the processed CpGs, we selected 115,556 CpGs with variance greater than 0.001 for downstream analyses. Differential methylation analysis of Illumina EPIC data was conducted with M-values (log2 ratio of beta value) using limma (v3.40.6) and q value (v2.16.0) R packages in R (v3.6.1) [49,50]. Differentially methylated CpGs were identified by fitting the M-values into a linear mixed-effects model adjusted for age, sex, repeat measures from single subject. The repeated measures were accounted for by calculating correlation coefficient by blocking by subject using the duplicateCorrelation function in limma. The correlation coefficient was found to be 0.857. The differentially methylated CpGs were deemed to be significant if the Q-value (p-values adjusted for false discovery rate) was under the threshold value of 0.05.

Differentially methylated CpGs were compared to the 115,556 CpGs for enrichment tests of genomic context using Mantel-Haenszel test, adjusting for Illumina EPIC array probe types. Functional significance of the CpGs were analysed with Genomic Regions Enrichment of Annotations Tool (GREAT) [26]. Distance of differentially methylated CpGs to the nearest transcriptional start sites (TSS) were annotated with TxDb.Hsapiens.UCSC.hg19.knownGene annotation file using the ChIPpeakAnno (v3.18.2) [51].

**Gene expression quantification**

RNA was isolated via RNaseasy mini kit (Qiagen, Germantown, MD) as outlined in manufacturer’s instructions, and quantitated on a Qubit 3.0 Fluorometer (Life Technologies, Carlsbad, CA). The digital multiplexed NanoString nCounter human v3 Cancer/Immune RNA expression assay (NanoString Technologies, Seattle, WA) was performed according to manufacturer’s instructions with total RNA. Briefly, 200ng total RNA was assayed by overnight hybridization (65°C) with nCounter gene-specific reporter and capture probes. Following a 20-hour hybridization, reporter and capture probes are washed away using the automated nCounter sample prep station and probe/target complexes are aligned and immobilized in the nCounter Cartridge. Cartridges are then placed in the nCounter digital analyser for data collection. nSolver Analysis software (NanoString) (v3.0) was used for data analysis including background correction by subtracting the mean of the six negative controls included on the NanoString platform and normalization using the average geometric mean of the thirty housekeeping genes included in the assay.

**Gene expression analysis**

Differential gene expression analysis of Nanostring data was conducted using edgeR (v3.26.8) and limma (v3.40.6) packages in R (v3.6.1) [49,52]. The normalized data from Nanostring were scaled to counts per million (cpm). 541 genes with greater than 10cpm in at least 2 samples were used for downstream analysis. Likelihood ratio tests were conducted after fitting the gene expression data to a negative binomial generalized linear model. 310 genes were determined as differentially expressed based on its likelihood ratio test FDR value threshold of 0.05. Gene ontology analysis was conducted with Over Representation Analysis of biological processes gene ontology through WebGestalt (Web-based Gene SeT AnaLysis Toolkit) 2017 [53].

**Abbreviations:**

EV: Extracellular vesicle; CF: Cystic fibrosis; NTA: Nanoparticle tracking analysis; BAL: Bronchoalveolar lavage
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Authors’ contributions
DAA and AA designed the study. JAD, DLM, and DSA provided clinical samples. DAA and HFH performed the experimental work. MKL performed data analysis with input from BCC. MKL and DAA wrote the manuscript. All authors approved the final version of the manuscript.

Ethics approval and consent to participate
This study was approved by the Dartmouth Hitchcock Institutional Review Board (#22781).

Data availability statement
DNA methylation array data has been deposited into Gene Expression Omnibus under accession number GSE142801. A token for access by reviewers is available by contacting the corresponding author. All other data used in this study is available upon request from the corresponding author.

Disclosure statement
The authors state that there are no conflicts of interest (financial nor personal) associated with this work.

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