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# Transcriptome Sequencing Data Sets for Determining Gene Expression Changes Mediated by Phase-Variable DNA Methyltransferases in Nontypeable *Haemophilus influenzae* Strains Isolated from Patients with Chronic Obstructive Pulmonary Disease

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**ABSTRACT** Nontypeable *Haemophilus influenzae* (NTHi) is a major bacterial cause of exacerbations in chronic obstructive pulmonary disease (COPD). Here, we report high-depth coverage transcriptome sequencing (RNA-seq) data from two NTHi strains, each encoding a different phase-variable methyltransferase. *modA* phase variation results in gene expression differences. These data will serve as an important resource for future studies.

Nontypeable *Haemophilus influenzae* (NTHi) is a human-adapted pathogen that causes exacerbations in chronic obstructive pulmonary disease (COPD) (1). Globally, COPD affects ~65 million people (2). Previous work examining NTHi pathobiology showed that phase-variable *modA* genes, encoding DNA methyltransferases, are involved in virulence by epigenetically regulating multiple genes (3–6). These systems are called phase-variable regulons (phasevarions) (7, 8). *ModA* allelic variants are defined by variation (<25% identity between alleles) in their central target recognition domain (TRD). The TRD determines methyltransferase specificity (9). Therefore, alleles containing different TRDs methylate different DNA sequences (7) and would control different phasevarions. Examination of an extensive panel of NTHi isolates from COPD patients (10) revealed the presence of two novel *modA* alleles, namely, *modA15* and *modA18*. We previously demonstrated that these methyltransferases are phase-variably expressed in prototype strains (11). Using matched *modA* on-off pairs of each of these prototype strains, we prepared triplicate biological replicates of total RNA using TRIzol (Thermo Fisher) according to the manufacturer's instructions. NTHi strains were grown to mid-log phase in brain heart infusion (BHI) broth (Oxoid, UK) at 37°C aerobically with 150 rpm shaking.

All methods used for transcriptome sequencing (RNA-seq) are identical to those described previously (12), except we used 100-bp paired-end sequence reads. Briefly, the Illumina Ribo-Zero Gold kit was used to prepare libraries, and libraries were assessed using an Agilent Bioanalyzer DNA 1000 chip. Quantitative PCR (qPCR) quantification assessed each library prior to normalizing (2 nM) and pooling using the Illumina cBot system and TruSeq paired-end (PE) cluster kit v3 reagents. The Illumina NovaSeq system with TruSeq sequencing by synthesis (SBS) kit v3 reagents was used to perform sequencing. Cleaned sequence reads were aligned against the respective reference genomes (GenBank accession numbers [CP029620](https://doi.org/10.1128/MRA.00526-19) [10P129H1; *modA15*] and [CP029621](https://doi.org/10.1128/MRA.00526-19) [84P36H1; *modA18*]). Reads were mapped to the respective genome using Bowtie2 aligner (v2.3.3.1), and transcripts were assembled with the read alignment and

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reference annotation-based assembly option in Stringtie v1.3.3 (<http://ccb.jhu.edu/software/stringtie/>). The Gencode annotation containing both coding and noncoding sequences for each genome was used to map reads (<http://www.gencodegenes.org/>). Raw gene count values were analyzed with edgeR (<https://bioconductor.org/packages/release/bioc/html/edgeR.html>), and the subread package (<http://subread.sourceforge.net/>) was used to quantitate counts using the featureCounts v1.5.3 utility. Gene expression differences were expressed as log<sub>2</sub> fold change of expression (logFC). Analysis also generated the following: average log count/million for each gene from all samples (logCPM values), quasi-likelihood F statistic for each gene from all samples (F values), *P* values to determine statistically different expression, and the false-discovery rate/adjusted *P* value for multiple-hypothesis testing.

Multiple genes are differentially regulated commensurate with *modA15* and *modA18* phase variation (44 genes differentially regulated between *modA15* on versus off; 10 genes differentially regulated between *modA18* on versus off). For example, the universal stress protein UspA (DLJ98\_08875) and the cold shock protein CspD (DLJ98\_02355) are both upregulated when *modA15* is off. In the *ModA18* phasevarion, two hypothetical genes (DLK00\_05505 and DLK00\_05510), encoding proteins of unknown function, are upregulated when *modA18* is on. These RNA-seq data will serve as a primary resource for studying the regulatory events resulting from genome-wide methylation differences by phase-variable methyltransferases during COPD and for determining the exact molecular mechanisms of *modA*-mediated gene expression differences.

**Data availability.** These data have been deposited in the NCBI Gene Expression Omnibus under the accession numbers [GSE129761](https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE129761) (*modA15* on versus off) and [GSE129764](https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE129764) (*modA18* on versus off).

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