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Inducible expression quantitative trait locus analysis of the *MUC5AC* gene in asthma in urban populations of children

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Abstract

Background: Mucus plugging can worsen asthma control, lead to reduced lung function and fatal exacerbations. MUC5AC is the secretory mucin implicated in mucus plugging, and *MUC5AC* gene expression has been associated with development of airway obstruction and asthma exacerbations in urban children with asthma. However, the genetic determinants of *MUC5AC* expression are not established.

Objective: To assess single-nucleotide polymorphisms (SNPs) that influence *MUC5AC* expression and relate to pulmonary functions in childhood asthma.

Methods: We used RNA-sequencing data from upper airway samples and performed cis-expression quantitative trait loci (eQTL) and allele specific expression (ASE) analyses in two cohorts of predominantly Black and Hispanic urban children, a high asthma-risk birth cohort and an exacerbation-prone asthma cohort. We further investigated inducible *MUC5AC* eQTLs during incipient asthma exacerbations. We tested significant eQTLs SNPs for associations with lung function measurements and investigated their functional consequences in DNA regulatory databases.

Results: We identified two independent groups of SNPs in the *MUC5AC* gene that were significantly associated with *MUC5AC* expression. Moreover, these SNPs showed stronger eQTL associations with *MUC5AC* expression during asthma exacerbations, consistent with inducible expression. SNPs in one group also showed significant association with decreased pulmonary functions. These SNPs included multiple EGR1 transcription factor binding sites suggesting a mechanism of effect.

Conclusions: These findings demonstrate the applicability of organ specific RNAsequencing data to determine genetic factors contributing to a key disease pathway. Specifically, they suggest important genetic variations that may underlie propensity to mucus plugging in asthma and could be important in targeted asthma phenotyping and disease management strategies.

Key Messages:

- Multiple *MUC5AC* SNPs associated with its expression and two independent groups of SNPs showed additive eQTL effects.
- By examining *MUC5AC* expression during respiratory illnesses, we showed an inducible increase in expression during asthma exacerbations according to genotype.
- By investigating DNA regulatory elements, we identified possible mechanisms for the observed associations of these SNPs.

Capsule summary: Our results add new information regarding the role of *MUC5AC* genotypes in childhood asthma pathogenesis in populations with high disease burden and suggest genetic predictors for targeted phenotyping and management strategies focused on mucin dysregulation.

Key words: Asthma, Expression quantitative trait, MUC5AC

Abbreviations:

ASE:	Allele Specific Expression
BH:	Benjamini-Hochberg
BMI:	Body Mass Index
CF:	Cystic Fibrosis
eQTL:	expression Quantitative Trait Locus
EGR1:	Early growth response protein 1
FDR:	False Discovery Rate
FEV1:	Forced Expiratory Volume in one second
FVC:	Forced Vital Capacity
GEO:	Gene Expression Omnibus
GWAS:	Genome Wide Association Study
ICS:	Inhaled Corticosteroid
LD:	Linkage Disequilibrium
SCS:	Systemic Corticosteroid
SNP:	Single Nucleotide Polymorphism

Introduction

Estimates of asthma heritability range from 35-90%, but genome-wide association studies (GWAS) only account for a minor fraction of the observed heritability¹⁻⁴. This is likely due in part to the racial, phenotypic, and molecular heterogeneity of asthma, but also to the limitations of GWAS approaches⁴. Moreover, the functional consequences of many asthma risk variants found by GWAS are incompletely understood⁵⁻⁹. Investigation of molecular and phenotypic consequences of genetic variation in or near genes implicated in asthma pathogenesis, and in targeted high-risk populations, can provide important new insights into the genetics of asthma with potential implications for population and personalized disease management.

Urban populations have increased asthma prevalence, suboptimal response to asthma therapies, and are more likely to have reduced lung function as they age¹⁰⁻¹³. Our previous studies highlighted *MUC5AC* overexpression as a specific characteristic in children who develop recurrent wheezing, high levels of allergic sensitization, and progressive loss of lung function during childhood¹⁴, and as a gene upregulated in the airways during incipient asthma exacerbations in children with exacerbation-prone disease¹⁵. Understanding genetic factors contributing to *MUC5AC* induction is critical given that overproduction can impair mucociliary transport and lead to mucus plugging^{16, 17}, which is the most common pathologic feature in patients with fatal asthma¹⁸⁻²⁰ and is a feature of severe disease in adult asthma²¹. Furthermore, genotypes in the *MUC5AC* region of chromosome 11 have recently been associated with asthma in adult European populations^{22, 23}.

We therefore hypothesized that *MUC5AC* genotypes could differentially regulate its expression in the airway, in particular during acute exacerbations of asthma. To test this hypothesis, we leveraged RNA-sequencing data from airway samples from two urban, predominantly Black and Hispanic cohorts of children to both genotype individuals and then perform combined expression quantitative trait loci (eQTL) and confirmatory allele specific expression (ASE) analyses from samples collected during disease control and during asthma exacerbations.

Methods

<u>Study design and sample composition:</u> The URECA birth cohort study started enrollment in 2005 in urban neighborhoods with high rates (>20%) of poverty in Baltimore, Boston, New York City, and St. Louis as previously described.²⁴ Pregnant women, whose children were at high risk for developing asthma due to a history of asthma, allergic rhinitis, or eczema in the mother or father, were recruited prenatally. Written informed consent was obtained from the families, and assent was obtained for children at ages 7 years and older. Asthma diagnosis was determined at age 10 and nasal cells were obtained at age 11 by brushing the anterior turbinate with a cytology brush. RNA was extracted and sequenced to analyze gene expression as previously described¹⁴ in 318 participants used for this analysis (Table 1); in brief total RNA was isolated from nasal epithelial brushes in RLT Plus lysis buffer (Qiagen) by using a Qiashredder column (Qiagen) and RNeasy mini kits (Qiagen). RNA-sequencing libraries were constructed from total RNA using SMART-Seq v4 Ultra Low Input RNA Kit (Takara) and single-read sequencing was carried out on a HiSeq2500 sequencer (Illumina), using a HiSeq SBS v4 Kit to generate 58-base reads. This data is publicly available at Gene Expression Omnibus (GEO) accession number GSE145505.

The MUPPITS study recruited children with established exacerbation-prone asthma between 2015 and 2017 across urban sites in 9 U.S. cities: Boston, New York, Detroit, Denver, Washington D.C., Chicago, Dallas, Cincinnati, and St. Louis, as previously described¹⁵. In brief, a participant was eligible for enrollment if he or she: was 6 to 17 years of age; was diagnosed with asthma by a clinician greater than 1 year prior to recruitment; had at least 2 asthma exacerbations in the prior year (defined as a requirement for systemic corticosteroids (SCS) and/or hospitalization); was treated with at least fluticasone 250 mcg 1 puff twice daily or its equivalent for those aged 6 to 11 years, or treated with at least fluticasone-salmeterol 250/50 mcg 1 puff twice daily or its equivalent for those aged 12 years and older; had peripheral blood eosinophils \geq 150 per mm³; was a non-smoker; lived in a census tract with a density of \geq 1000 families per square mile and at least 10% of families with income below the poverty level. Participants were followed prospectively for respiratory illnesses, during which they returned to the study site twice in the 6-day period after the start of symptoms for collection of nasal lavage samples and pulmonary function testing. An exacerbation illness was defined based on whether the participant was treated with SCS within 10 days following the onset of respiratory symptoms or not. RNA was extracted from nasal lavage samples and sequenced to analyze gene expression as previously described¹⁵ in 106 participants used for this analysis (Table 1); in brief, total RNA was isolated from nasal cell pellets in RNAprotect Saliva Reagent (Qiagen) by using a Qiashredder column (Qiagen) and RNeasy mini kits (Qiagen). Sequencing libraries were constructed from total RNA using TruSeq RNA Sample Preparation Kits v2 (Illumina) and singleread sequencing was carried out on a HiSeq2500 sequencer (Illumina), using a HiSeq SBS v4 Kit to generate 58-base reads. These data are publicly available at GEO accession number GSE115824.

<u>SNP determination and validation:</u> RNA-sequencing reads were first mapped to the GRCh38.p10 reference genome using the STAR aligner (v2.4.2a). Base calls were made at all SNP positions listed in NCBI dbSNP Build 151 using samtools (v1.7) with the arguments `samtools mpileup -f \$grch38FastaFile -B -C 50 -d 250 -l \$dbSnp151Bedfile -q 30 -Q 13 -g -e 20 -h 100 -L 250 -o 40 \$alignmentFile -o \$bcfCallFile`. The call files were converted to VCF format and filtered to exclude sites where fewer than 10 reads aligned using bcftools (v1.9). SNPs were further filtered to select those with at least 2 different genotypes represented among cohort subjects. MUPPITS SNPs and libraries were filtered to require at least 75% concordance in calls among libraries from the same donor and any non-concordant libraries were removed from the analysis. For 149 participants in the URECA cohort who previously had DNA-based microarray genotyping using an Illumina Custom 3000 platform (550,224

variants), 8,568 overlapping SNPs were compared between RNA-based and DNA-based calls and found to be 99.2% concordant.

Statistical analyses: SNPs spanning the MUC5AC gene, spanning from the transcriptional start site to the transcriptional end site (chr11:1157953-1201138), were used for eQTL analysis. SNPs were filtered to those genotyped in >50% of the URECA cohort and checked that they satisfied Hardy-Weinberg equilibrium by the chi square goodness-of-fit test with 2 degrees of freedom (Benjaminin-Hochberg [BH] false discovery rate adjusted p-value [FDR]>0.05) using the HardyWeinberg R package²⁵. MUC5AC expression was associated to SNP genotypes using a weighted linear model using the limma R package and empirical Bayes method^{26, 27} and considering each allele under an additive effect (0, 1, or 2 minor alleles). For multiallelic sites, any minor allele was counted as 1. For the URECA cohort, the model was adjusted for cell percentages in the sample and study site, which contributed to gene expression variability, as previously described¹⁴. For the MUPPITS cohort, the model was adjusted for cell percentages, study site, viral positivity, age, sequencing depth and a random effect for individual, which contributed to gene expression variability as previously described¹⁵. In both cohorts further adjusting for self-reported race gave very similar results, so race was not included in the final models. Multiple testing corrections were performed using the BH method and significant eQTL SNPs were defined as those with a FDR<0.05. Pairwise r^2 measures of linkage disequilibrium (LD) were calculated among significant SNPs for each cohort using the LDheatmap R package²⁸. A Euclidean distance matrix was constructed from the r^2 values in URECA and used for hierarchical clustering using the average method (hclust function in R), which divided SNPs into 3 cluster groups (called A, B, or other). Group A and B representative SNPs were selected from the eQTL significant SNPs as those with highest percentages of calls to avoid potential bias by smaller sample sizes. These SNP groups were assessed for independent eQTL effects by performing conditional analyses in which we included the representative SNP or SNPs (0, 1, or 2 minor alleles) as covariates in the eQTL linear model and investigated the effect on the remaining SNPs. In the MUPPITS cohort, the interaction effect between a representative SNP and exacerbation status was determined by adding an interaction term to the linear model. ASE was assessed by calculating the number of counts of the minor allele divided by the total counts (minor and major allele) within each individual heterozygous at a given SNP. Significance was determined using a binomial test and the log2 allelic fold change (log2-aFC) was calculated as the log2-ratio between the expression of the minor allele compared to the major allele in heterozygotes and reported on a non-log scale (aFC)²⁹. Associations between a group-representative SNP and FEV₁/FVC and FEV₁ % predicted values were tested using a linear model adjusted for sex (URECA) or sex and age (MUPPITS); all URECA participants were the same age (10 years) at the time of pulmonary function assessment.

<u>SNP functional assessment</u>: Functional consequences of the identified significant SNPs were assessed by investigating amino acid changes using dbSNP³⁰, transcriptional

regulatory consequences in ENCODE and ORegAnno using the UCSC genome browser interface³¹⁻³⁵, and predicted protein functional effects in Meta-SNP³⁶ and Polyphen-2³⁷.

<u>Comparison to DNA genotyping data</u>: LD between the group A or group B representative SNP to previously published non-coding SNPs was performed by examining DNA-sequencing based genotypes of those published SNPs in a subset of 243 (76.4%) individuals in the URECA cohort. LD was computed between these SNPs as described above using the LDheatmap R package²⁸.

Results

MUC5AC SNPs associated with its expression in a high asthma risk birth cohort

The URECA cohort consists of urban children who are predominantly Black and Hispanic with high rates of poverty. By age 10 years, 29.5% of these children had a diagnosis of asthma (Table 1). 86 SNPs within the MUC5AC gene were identified from nasal brush RNA-sequencing data for eQTL analysis. 43 of these showed a significant eQTL association with MUC5AC expression (FDRs<0.05) (Figure 1a, Table 2). Clustering of SNPs by LD (r^2), as well as conditional eQTL analyses demonstrated that there were at least two groups of SNPs each independently associated with MUC5AC expression (Figure 1b, Supplementary Figure 1, Supplementary Table 1). Group A contained 10 SNPs; at each group A SNP, individuals homozygous for the major allele showed the lowest MUC5AC expression levels, with an increase according to number of minor alleles (eQTL β -coefficients=0.12 to 0.21). Representative of group A, rs1132436 (eQTL β-coefficient=0.20; FDR=-1.63E-4), had a minor allele frequency of 35.6% and explained 5.4% of variance of MUC5AC expression in this population (Figure 1c). ASE, assessed by RNA copy number in heterozygotes, confirmed significantly higher expression of the minor allele with an allelic fold change (aFC)=2.25 at rs1132436 (p<2E-16) (Figure 1d). Group B contained 29 SNPs and at each SNP the minor allele was associated with decreased MUC5AC expression (eQTL β -coefficients = -0.13 to -0.31). Representative of group B, rs1292198170 (β-coefficient=-0.28, FDR=1.9E-3), had a minor allele frequency of 13.0% and accounted for 3.1% of variance of MUC5AC expression (Figure 1e). ASE confirmed lower expression of the minor allele with an aFC=0.42 (p<2E-16) (Figure 1f). rs1132436 in group A and rs1292198170 in group B showed an additive effect and together accounted for 7.9% of variance of MUC5AC expression (Figure 1g). The remaining 4 significant SNPs had low LD with the group A or B SNPs and with one another, and all had relatively low minor allele frequencies (<2.8%) so were not investigated further.

<u>MUC5AC SNPs showed an inducible effect on its expression during asthma</u> <u>exacerbations</u>

To validate these results and assess their relevance to asthma exacerbations, we conducted eQTL and ASE analyses in the MUPPITS cohort, a demographically similar population to URECA, but consisting entirely of children with established, exacerbation-prone asthma (Table 1)¹⁵. Participants had nasal lavage samples collected during respiratory illnesses, which were used for RNA-sequencing. 24 *MUC5AC* SNPs were detected in this dataset, predominantly near the 3' end of the gene. Among the group A SNPs, 7 of 10 were detected, 5 of which were significantly associated with *MUC5AC* expression (FDR<0.05); the other two showed similar trends (FDRs=0.07 and 0.12) (Figure 2a). eQTL effect sizes were larger in this dataset (rs1132436, β -coefficient=0.57; FDR=8.9E-3) compared to those estimated in URECA (Table 2). Conditional eQTL analysis and LD assessment demonstrated similar co-dependence among group A SNPs in this population (Supplemental Figure 2, Figure 2b, Supplementary Table 2).

Because the MUPPITS samples were collected during respiratory illnesses, we could test the effects of asthma exacerbations on the observed eQTLs. The *MUC5AC* eQTL association at group A SNP rs1132436 was significantly greater during respiratory illnesses that led to asthma exacerbations relative to uncomplicated upper respiratory illnesses (Δ FC=2.42, p=5.7E-6) (Figure 2c), which accounted for the overall larger effect sizes observed in this population. During exacerbations, rs1132436 explained 20.3% of variance in gene expression as compared to 1.0% of variance during limited upper respiratory illnesses. ASE confirmed significantly higher expression of the minor allele in heterozygotes (p=4.0E-5) (Figure 2d), and the aFC was higher during exacerbations (aFC=2.4) compared to limited upper respiratory illnesses (aFC=1.4).

Identification of group B SNPs, which are closer to the 5' end of the gene, was limited in the MUPPITS cohort because of library preparation method. We were able to identify genotypes at the group B SNP rs1292198170 in only 31.4% of the population. Likely due in part to the small sample size, neither this SNP nor other group B SNPs reached an eQTL FDR cutoff <0.05, but similar to URECA rs1292198170 showed a trend towards lower expression in heterozygotes (β -coefficient=-0.90, FDR=0.24). Despite the small sample size, we did observe a statistically significant inducible eQTL effect at this SNP during respiratory illnesses that led to asthma exacerbations relative to uncomplicated upper respiratory illnesses (Δ FC=0.27, p=3.9E-3) (Figure 2e). During exacerbations, rs1292198170 explained 5.2% of variance as compared to <1% of variance during limited upper respiratory illnesses. Similarly, ASE was significant in heterozygotes (p=8.0E-3) and the overall aFC was 0.32 (Figure 2f).

Together these two SNPs showed an additive inducible eQTL effect (Figure 2g) and explained 42.4% of variance during exacerbations as compared to 9.7% of variance during limited upper respiratory illnesses in the subset of the population with identifiable SNPs at both positions (30.4% of the population).

MUC5AC SNPs associated with pulmonary functions

We investigated the relationship of these SNPs with pulmonary function measures in each cohort. In the URECA cohort, the minor allele at rs1132436 (group A) was significantly associated with lower FEV₁/FVC values (β -coefficient=-1.33, p=1.8E-2) (Figure 3a) and showed a non-significant trend towards lower FEV₁ % predicted (β -coefficient=-1.74, p=0.13). The minor allele at rs1292198170 (group B) showed a non-significant trend towards higher values of FEV₁/FVC (β -coefficient=1.45, p=0.10) and FEV₁ % predicted (β -coefficient=2.58, p=0.16). In the MUPPITS cohort, rs1132436 (group A) was also associated with lower lung function as measured by both FEV₁/FVC (β -coefficient=-1.64, p=4.0E-2) and FEV₁ % predicted (β -coefficient=-5.71, p=2.2E-2) (Figure 3b). rs1292198170 (group B) did not show a relationship with pulmonary functions in the MUPPITS cohort (p-values>0.75), perhaps due in part to the small sample size genotyped at this SNP.

Functional inference of MUC5AC eQTL SNPs

To investigate potential functional consequences of the identified SNPs, we examined nucleotide and amino acid changes in dbSNP, DNA regulatory elements in ENCODE and ORegAnno through the UCSC genome browser, and predicted mutational consequences in Meta-SNP and Polyphen-2 (Table 2)³⁰⁻³⁷. DNA binding and regulatory elements spanning these SNPs revealed 4 Early Growth Response 1 (EGR1) transcription factor binding sites, 2 in group A and 2 in group B, as well as 3 CpG islands in group A. 16 of the significant SNPs lead to missense mutations, 4 in group A and 12 in group B. Predictions of functional consequences varied across tools with one missense mutation in block A and 3 in block B predicted as deleterious by at least one of the tools (Table 2).

LD of MUC5AC eQTL SNPs with prior disease associated SNPs

Using DNA genotype data from a subset of the URECA population (243 individuals, 76.4% of the population), we examined the LD between the SNPs identified in our study and non-coding *MUC5AC* SNPs previously associated with asthma or cystic fibrosis (CF) severity. rs11602802, which was previously associated with moderate-severe asthma in European ancestry²², was not in LD with either group of SNPs (rs1132436 r^2 = 0.08, rs1292198170 r^2 = 0.004) in our sample. rs12788104, which was significantly associated with adult asthma in British white individuals²³, was in weak LD with group A but not group B SNPs (rs1132436 r^2 = 0.37, rs1292198170 r^2 = 0.008). rs28514396, a *MUC5AC* intronic variant associated with CF severity³⁸ was in LD with group A but not group B SNPs (rs1132436 r^2 = 1, rs1292198170 r^2 = 0.09).

Discussion

MUC5AC hypersecretion plays a pathogenic role in asthma through its relation to airway hyperresponsiveness and mucus plugging. In this study, we used RNA-sequencing data from airway samples to derive genotyping information and perform

combined eQTL and ASE analyses focused on the *MUC5AC* region in two cohorts of predominantly African-American and Hispanic children using samples when children were well, and during incipient asthma exacerbations. We report two distinct groups of genetic polymorphisms in *MUC5AC* that independently relate to its expression, at least one of which is also significantly associated with decreased lung function. Moreover, we show that the effects of these SNPs on *MUC5AC* expression are increased during respiratory illnesses that result in asthma exacerbation, suggesting that inducible overexpression during exacerbations could contribute to exacerbation pathophysiology in genetically susceptible individuals. Functional assessment of these SNPs suggests potential causal effects, most notably they may alter EGR1 transcription factor binding.

Interestingly, despite long standing recognition of the role of mucus hypersecretion in asthma, GWAS studies have only recently linked specific MUC5AC polymorphisms, rs11602802²² and rs12788104²³, to asthma. These studies were restricted to individuals of European ancestry and the observed associations were with moderate-to-severe and adult-onset asthma. This relatively recent discovery of MUC5AC as an asthma risk locus may be due in part to the specificity of this finding to moderate-severe asthma, but may also be due in part to incomplete mapping of the MUC5AC gene until 2014³⁹. There was previously a large gap in the *MUC5AC* sequence including its large central exon, which spanned nearly all of the SNPs identified in the current study. These SNPs were missing from the data used in each of these prior GWAS studies. Since rs11602802 is not in LD with either group of our identified eQTL SNPs, it presumably has an independent effect in relation to asthma risk or may be specific to European populations. rs12788104 was in weak LD with group A SNPs in our URECA population, so may represent an overlapping genetic risk to that identified in our study. However, since the URECA cohort includes children with and without asthma, our results suggest that the identified group A and B SNPs are MUC5AC eQTLs independent of a diagnosis of asthma. Hence these SNPs may be more relevant to the pathophysiologic endotype of asthma than the risk of developing asthma. Consistent with our results, two prior targeted gene association studies have linked SNPs in LD with group A with pulmonary diseases: the non-synonymous coding SNP, rs1132440 in group A was modestly associated with bronchitis, wheeze, and asthma in a British cohort⁴⁰, and the intronic SNP, rs28514396 (r^2 =1 with rs1132436 in URECA) was associated with severity of lung disease in CF³⁸.

Prior work has shown EGR1 is an essential transcription factor for the expression of MUC5AC in airway epithelium^{41, 42}, and it is upregulated in airway epithelium in a model of respiratory virus infection⁴³. These lead us to hypothesize that the *MUC5AC* eQTLs in known EGR1 binding sites disrupt its binding as the most likely mechanism for the eQTL effects. Alternatively, given the number of missense mutations among the identified *MUC5AC* SNPs, it is also plausible that these altered codons could affect gene expression, protein function or both⁴⁴ and may thereby contribute to the observed eQTLs and/or associations with pulmonary function. Interestingly, the group A SNPs have been shown to be in high LD with an insertion in the central exon encoding

tandem repeats of an eight-amino-acid repeat rich in proline, threonine, and serine (PTS-TR domains)^{38, 39}. These domains are characteristic of mucin proteins and are heavily glycosylated. The group A SNPs are thus indicative of a higher molecular weight and highly glycosylated protein that may play an important role in mucus plug formation^{38, 39, 45}, which may functionally contribute to the associations observed with pulmonary function.

An important limitation of our study is that it assessed links between genotype, mRNA expression in the upper airway, and pulmonary physiology. Linking these genetic polymorphisms to lower airway MUC5AC protein expression and/or lung imaging consistent with mucus plugging^{16, 21} will help validate the pathogenic importance of these SNPs. Additionally, *in vitro* studies will be necessary to further investigate the functional effects of these SNPs to prove their mechanisms of effect. Our results suggest these as important next studies that can fully define the relevance of *MUC5AC* genotypes in the pathology of asthma, as well as potentially in other "muco-obstructive" lung diseases⁴⁶.

An important technical strength of our work is demonstration of the synergistic value of utilizing RNA-sequencing derived genotypes, organ specific gene expression, and sampling during disease exacerbation to gain novel insights into genetic contributions to a disease. These allowed for combined eQTL, ASE, and induction analyses to determine genetic factors contributing to a key pathway in asthma. We anticipate that the insights described in this study can lead to a more personalized understanding of asthma pathogenesis. Ideally, this will lead to strategies targeting mucus hypersecretion in genetically at-risk individuals as a way to reduce risk of disease inception and progression as well as risk of exacerbation⁴⁷.

In conclusion, our results add important new and detailed information regarding the role of *MUC5AC* genotypes in childhood asthma pathogenesis, demonstrating their direct effect on gene expression, relationships to pulmonary physiology, and identifying etiologies of their effects.

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Cohort population characteristics												
	URECA	MUPPITS										
Sex												
Female	151 (47.5%)	52 (49%)										
Male	167 (52.5%)	54 (51%)										
Race												
African-American, non-	233 (73.3%)	57 (53.8%)										
Hispanic												
Hispanic	59 (18.5%)	36 (33.9%)										
Other, non-Hispanic	26 (8.2%)	13 (12.3%)										
<u>Age in years, mean</u>	10*	10.8 [6-17]										
<u>Diagnosis</u>												
Asthma	94 (29.5%)	106 (100%)										
No asthma	224 (70.5%)	-										
ICS use	64 (20.1%)	106 (100%)										
Pulmonary function												
FEV ₁ /FVC, mean	82.7% [59.7% - 98.2%]	76.0% [57.2-96.0%]										
FEV ₁ percent predicted	99.2% [62.2% - 139.1%]	91.7% [42.3-136.9%										
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Gestational age in weeks	38.8 [34 - 42]	Not collected										
BMI percentage	69.2% [0.8% - 99.8%]	78.2% [0.9% - 99.9%										
Number of allergen	3.2 [0 - 13]	4.7 [0 - 15]										
sensitizations												
City of residence												
Baltimore	92 (28.9%)	-										
Boston	74 (23.2%)	14 (13.2%)										
Chicago	-	7 (6.6%)										
Cincinnati	-	7 (6.6%)										
Dallas	-	8 (7.5%)										
Denver	-	13 (12.3%)										
Detroit	-	14 (13.2%)										
New York	57 (17.9%)	23 (21.7%)										
St. Louis	95 (32.6%)	13 (12.3%)										
Washington DC	-	6 (5.7%)										

 Table 1. Demographics of URECA and MUPPITS cohorts.

Shown are the population characteristics of the URECA and MUPPITS cohorts. Shown are counts of mean values. () indicate percentage of the population. [] indicate ranges of values in the population. *URECA children were all age 10 at the time of pulmonary function assessment and asthma diagnosis, and were age 11 at time of nasal brush collection. BMI is body mass index; ICS is inhaled corticosteroid.

Table 2. Two independent blocks of SNPs show an eQTL effect on MUC5AC

The 43 significant eQTL SNPs from the URECA cohort are listed, divided into 2 independent groups, A and B, based on LD and conditional analyses. For each SNP we list the chromosomal position, percent of subjects called at that SNP, and minor allele frequency in the population. The β -coefficient and FDR values are shown, as well as the FDR after each conditional analysis and the r^2 with the SNP used to represent the group. The analogous results are presented with these SNPs in the MUPPITS cohort with blanks indicating SNPs that could not be called in the MUPPITS cohort. The base and amino acid changes are shown, as well as the identified DNA regulatory elements at that position, and functional predictions of amino acid changes.

Figure 1. MUC5AC SNPs showed an eQTL effect in the URECA cohort

a, The locus zoom plot for the URECA cohort showing eQTLs. Among the 86 cis SNPs detected, 43 showed significant association with *MUC5AC* expression (FDR< 0.05). Points are colored according to the minor allele frequency of the SNP and the point size reflects the percent of study subjects with a call at that SNP. **b**, An LD matrix shows the pairwise *r*² values among these 86 SNPs. **c**, For the block A representative SNP, rs1132436, *MUC5AC* gene expression was increased according to the number of minor allele reads. **d**, ASE of heterozygotes at rs1132436 showed an increased proportion of minor allele reads. **e**, For the block B representative SNP, rs1292198170, *MUC5AC* gene expression was decreased in heterozygotes. **f**, ASE of heterozygotes at rs1132436 and rs1292198170 showed an additive effect on *MUC5AC* expression. Boxplots show mean values, interquartile ranges, and 1.5 interquartile ranges.

Figure 2. *MUC5AC* SNPs showed an inducible eQTL effect during asthma exacerbations in the MUPPITS cohort

a, The locus zoom plot for the MUPPITS cohort showing eQTLs. Among the 24 cis SNPs detected, 5 showed significant association with MUC5AC expression (FDR< 0.05). Points are colored according to the minor allele frequency of the SNP and the point size reflects the percent of study subjects with a call at that SNP. b, An LD matrix shows the pairwise r^2 values among these 24 SNPs. **c**, For the block A representative SNP rs1132436, MUC5AC gene expression was increased according to the number of minor alleles and showed an inducible eQTL effect during exacerbations compared to non-exacerbation samples. d, ASE of heterozygotes at rs1132436 showed an increased proportion of minor allele reads. e, For the block B representative SNP, rs1292198170, MUC5AC gene expression showed a non-significant (FDR=0.24) decrease in heterozygotes relative to homozygotes and showed an inducible eQTL effect during exacerbations compared to non-exacerbation samples (p=3.9E-3). f, ASE of heterozygotes at rs1292198170 showed a decreased proportion of minor allele reads. g, Genotypes at rs1132436 and rs1292198170 showed an additive effect on MUC5AC expression. Boxplots show mean values, interguartile ranges, and 1.5 interguartile ranges.

Figure 3. MUC5AC SNP rs1132436 showed a significant association with pulmonary function in both URECA and MUPPITS.

a, The FEV₁/FVC ratio was significantly associated with the genotype at the block A representative SNP rs1132426 in the URECA cohort. **b**, Both the FEV₁/FVC ratio and FEV₁ % predicted were significantly associated with the genotype at the block A

representative SNP rs1132426 in the MUPPITS cohort. Boxplots show mean values, interquartile ranges, and 1.5 interquartile ranges.

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SNP		URECA co	ohort result	s					MUPPIT	S cohort res	ults				SNP details								
Rsid	Chromos omal position	% Subjs with call	Minor Allele Freq	FDR	Effect size	FDR, rs129219 8170 condition ed	LD R2 vs rs12921 98170	FDR, rs1132436 conditioned	LD R2 vs rs11324 36	FDR, dual conditione d, rs1132436 + rs1292198 170	% Subjs with call	Minor Allele Freq	FDR	Effect size	FDR, rs1132 436 conditi oned	LD R2 vs rs11324 36	Base chng	Codon change	Mutation al conseque nce	Amin o acid chang e	Meta- SNP predicti on	Polyphe n-2	DNA elements in region
Group A																							
rs1132436	1199741	95.9%	35.6%	1.48E-	0.199	9.87E-03	0.086	NA	NA	NA	71.7%	36.8%	8.89E-	0.574	NA	NA	C>T	[CCG]>[CTG]	Missense	P5521	Neutral	Neutral	none
rs1132440	1200591	98.4%	35.3%	2.80E- 04	0.207	9.87E-03	0.082	2.26E-01	0.986	8.10E-01	70.8%	38.2%	1.42E- 02	0.549	NA	1.000	G>C	[CGG]>[CGC]	Synonym ous	-			CpG island, CpG database
rs28633254	1199373	98.1%	35.4%	1.75E- 04	0.212	9.87E-03	0.089	5.66E-01	0.972	8.99E-01	69.8%	36.5%	3.04E- 03	0.665	4.95E- 01	0.977	C>T	[CCC]>[CCT]	Synonym ous				EGR1 TF binding site, ORegAnn
rs1132434	1195265	96.2%	35.8%	4.37E- 04	0.183	2.79E-02	0.098	9.08E-01	0.972	8.99E-01	50.0%	34.9%	1.42E- 02	0.636	1.00E+0 0	1.000	G>C	[CAG]>[CAC]	Missense	Q514 8H	Neutral	Neutral	none
rs1379997765	1188914	97.2%	35.9%	3.48E- 04	0.200	2.79E-02	0.096	8.28E-01	0.964	9.65E-01	-	-		-	-	-	G>A	[CGC]>[CAC]	Missense	R3590 H	Neutral	Deleteri ous	none
rs28415193	1195214	96.2%	35.6%	2.86E- 04	0.194	1.07E-02	0.081	3.81E-01	0.950	8.10E-01	43.4%	41.3%	6.77E- 02	0.509	1.00E+0 0	1.000	G>A	[ACG]>[ACA]	Synonym ous				CpG island, CpG database
rs1132433	1194354	96.2%	36.3%	1.22E- 03	0.160	6.29E-02	0.083	2.64E-01	0.938	8.10E-01	31.1%	48.5%	2.21E- 02	0.680	4.95E- 01	0.958	G>A	[ACG]>[ACA]	Synonym ous				CpG island, EGR1 TF binding site
rs878989392	1182583	58.2%	43.2%	1.07E- 02	0.113	6.29E-02	0.082	7.45E-01	0.894	8.99E-01	-	-	-	-	-	-	C>G, T	[CCC]>[GCC], [CCC]>[TCC]	Missense	P1480 A, P1480 S	Neutral, Neutral	No predicti on	none
rs28503875	1196902	96.2%	44.4%	1.84E- 02	0.121	4.05E-01	0.112	7.96E-01	0.690	9.65E-01	51.9%	46.4%	1.20E- 01	0.351	5.50E- 01	0.623	A>G	[CCA]>[CCG]	Synonym ous				none
rs996372487	1172511	66.0%	28.8%	1.48E- 04	0.193	3.28E-03	0.058	1.39E-01	0.494	8.10E-01	-	-	-	-	-	-	C>A	[GGC]>[GGA]	Synonym ous				none
Group B																							
rs1292198170	1191333	99.4%	13.0%	1.68E- 03	-0.281	NA	NA	1.30E-02	0.086	NA	31.1%	9.1%	2.45E- 01	-0.905	4.95E- 01	0.186	C>A	[GCC]>[GCA]	Synonym ous				none
rs1466012270	1185033	94.7%	14.6%	1.22E- 03	-0.214	9.02E-01	0.888	6.46E-02	0.109	9.55E-01	-	-	-	-	-	-			ous				none
rs1469470921	1185767	95.9%	13.8%	03 6.02F-	-0.260	5.19E-01	0.843	8.44F-03	0.095	8.10E-01	-	-	-	-	-	-	C>A	[ACC]>[AAC]	ous	T2541	Neutral	No	none
101100 170321	1105707	551576	151070	04	0.200	51152 01	0.010	0.112.00	0.035	0.102 01							0.11	[//////	Missense	N	neutiti	predicti on	lione
rs1248958955	1190666	98.7%	15.4%	2.60E- 04	-0.278	2.32E-01	0.830	8.96E-03	0.111	8.25E-01	18.9%	7.5%	7.16E- 01	-0.319	4.95E- 01	0.242	C>T	[ACA]>[ATA]	Missense	T4174 I	Neutral	No predicti on	none
rs1222080419	1187320	95.9%	15.1%	6.09E- 04	-0.229	9.02E-01	0.827	8.96E-03	0.114	8.99E-01	-	-	-	-	-	-	A>G	[AGC]>[GGC]	Missense	S3059 G	Neutral	No predicti	none
rs1330735919	1190684	98.7%	15.3%	7.37E- 04	-0.249	2.28E-01	0.823	1.24E-02	0.110	8.25E-01	22.6%	16.7%	1.20E- 01	-0.845	4.95E- 01	0.369	C>T	[GCC]>[GTC]	Missense	A4180 V	Neutral	No predicti	none
rs1269243287	1188446	97.2%	14.6%	8.65E- 04	-0.229	6.29E-01	0.823	8.96E-03	0.116	8.99E-01	-	-	-	-	-	-	C>T	[CCT]>[CTT]	Missense	P3434 L	Neutral	No predicti	none
rs1365388725	1184488	96.5%	14.8%	1.24E-	-0.224	8.22E-01	0.820	8.96E-03	0.118	8.25E-01	-	-	-	-	-	-	A>G	[AAC]>[GAC]	Missense	N2115	Neutral	Neutral	none
rs1462651135	1188935	95.0%	15.4%	1.48E- 04	-0.270	8.22E-02	0.819	7.03E-03	0.116	8.10E-01	-	-	-	-	-	-	G>A	[CGG]>[CAG]	Missense	R3597 Q	Neutral	Deleteri ous	none
rs1403452853	1185584	96.2%	15.2%	2.60E- 04	-0.250	5.18E-01	0.816	7.03E-03	0.126	8.25E-01	-	-	-	-	-	-	C>T	[GCT]>[GTT]	Synonym ous				none

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rs1391780761	1188422	97.8%	14.3%	9.20E- 03	-0.187	9.39E-01	0.809	6.16E-02	0.115	8.44E-01 Journa	al Pre-p	proof	- 	-	-	- 	G>C	[CGT]>[CCT]	Missense	R3426 P	Neutral	No predicti on	none
rs35779873	1197482	93.1%	13.9%	6.28E- 03	-0.184	5.55E-01	0.750	9.57E-02	0.096	8.10E-01	45.3%	13.5%	4.41E- 01	-0.351	4.95E- 01	0.221	C>T	[GTC]>[GTT]	Synonym ous				none
rs1444693715	1188653	78.6%	13.0%	1.48E- 04	-0.311	8.47E-01	0.744	7.03E-03	0.144	8.99E-01	-	-	-	-	-	-	G>A	[AGC]>[AAC]	Missense	S3503 N	Neutral	No predicti on	none
rs1368704331	1190025	98.4%	11.2%	1.90E- 03	-0.275	6.29E-01	0.732	1.30E-02	0.105	8.99E-01	-	-	-	-	-	-	C>A	[ACC]>[ACA]	Synonym ous				none
rs35915689	1197925	87.4%	16.0%	7.04E- 04	-0.205	8.17E-01	0.728	1.80E-02	0.133	9.92E-01	52.8%	16.1%	1.49E- 01	-0.517	4.95E- 01	0.247	C>G	[CCC]>[CCG]	Synonym ous				none
rs34831688	1198945	97.5%	16.6%	7.32E- 04	-0.226	5.40E-01	0.702	4.14E-02	0.119	8.31E-01	63.2%	14.2%	7.05E- 01	-0.113	4.95E- 01	0.170	C>T	[GAC]>[GAT]	Synonym ous				EGR1 TF binding site, ORegAnn o
rs35783651	1163029	80.5%	15.2%	2.79E- 03	-0.182	6.29E-01	0.691	6.16E-02	0.094	8.99E-01	-	-	-	-	-	-	C>G	[AGC]>[AGG]	Missense	S221R	Disease	No predicti on	EGR1 TF binding site, ORegAnn o
rs28691231	1164456	76.7%	16.2%	2.27E- 03	-0.180	8.47E-01	0.688	8.91E-02	0.119	8.99E-01	-	-	-	-	-	-	C>T	[TGC]>[TGT]	Synonym ous				none
rs13380	1201105	98.7%	13.9%	8.65E- 04	-0.251	3.39E-01	0.672	1.20E-02	0.108	8.44E-01	73.6%	11.5%	5.26E- 01	-0.189	4.95E- 01	0.177	A>C		3 Prime UTR	Variant			none
rs879185391	1176633	81.4%	16.0%	1.13E- 03	-0.191	7.70E-01	0.628	4.09E-02	0.151	9.87E-01	-	-		-	-	-	C>T	[GCC]>[GCT]	Synonym ous				none
rs879122882	1176540	82.1%	13.6%	3.57E- 02	-0.134	9.02E-01	0.627	2.26E-01	0.109	8.99E-01	-	-	-	-	-	-	C>T	[TGC]>[TGT]	Synonym ous				none
rs36132281	1177269	78.3%	15.3%	1.12E- 02	-0.150	6.60E-01	0.627	1.39E-01	0.126	8.99E-01	-		K	-	-	-	T>C	[TTT]>[TTT]]	Synonym ous				none
rs36142159	1176982	82.1%	16.1%	2.15E- 02	-0.137	8.22E-01	0.618	2.21E-01	0.161	8.25E-01		0	-	-	-	-	C>T	[TGC]>[TGT]	Synonym ous				none
rs13010	1201051	99.4%	20.7%	1.24E- 03	-0.216	4.20E-01	0.536	1.39E-01	0.150	8.99E-01	84.9%	17.8%	4.67E- 01	-0.191	6.08E- 01	0.224	A>C		3 Prime UTR	Variant			none
rs3087562	1200759	98.4%	20.4%	7.55E- 03	-0.171	6.70E-01	0.518	9.57E-02	0.153	8.99E-01	70.8%	16.7%	4.83E- 01	-0.200	4.95E- 01	0.236	C>T		3 Prime UTR	t Variant			none
rs11347	1201099	99.1%	20.2%	1.33E- 03	-0.211	4.90E-01	0.487	1.43E-01	0.155	8.99E-01	77.4%	15.2%	4.69E- 01	-0.206	4.95E- 01	0.228	C>T		3 Prime UTR	t Variant			none
rs55861305	1161529	67.3%	9.1%	1.24E- 03	-0.266	4.56E-01	0.405	3.05E-02	0.064	8.10E-01	-	-	-	-	-	-	G>T	[GTC]>[TTC]	Missense	V52F	Neutral	Deleteri ous	none
rs1222886276	1190064	94.0%	6.0%	3.60E- 02	-0.229	7.70E-01	0.317	2.47E-01	0.088	9.65E-01	-	-	-	-	-	-	C>A, T	[CAC]>[CAA], [CAC]>[CAT]	Missense, Synonym ous	H3973 Q	Neutral	No predicti on	none
Other																							
rs1204279823	1191305	96.9%	2.8%	7.71E- 04	-0.529	8.22E-02	0.141	3.00E-02	0.019	8.10E-01	19.8%	9.5%	2.45E- 01	-1.147	5.50E- 01	0.203	C>T	[TCT]>[TTT]	Missense	S4387 F	Neutral	Deleteri ous	none
rs34815853	1197927	87.1%	1.3%	3.13E- 02	0.439	2.19E-01	0.009	1.48E-01	0.027	8.10E-01	-	-	-	-	-	-	C>A	[GCG]>[GAG]	Missense	A5353 E	Neutral	Deleteri ous	none
rs34474233	1197926	86.8%	1.3%	3.16E- 02	0.437	2.19E-01	0.009	1.48E-01	0.027	8.10E-01	-	-	-	-	-	-	G>A	[GCG]>[ACG]	Missense	A5353 T	Neutral	Neutral	none
rs28403537	1167980	83.0%	1.3%	4.25E- 02	0.405	2.32E-01	0.009	1.86E-01	0.027	8.10E-01	-	-	-	-	-	-	C>T	[GCG]>[GTG]	Missense	A497V	Neutral	Neutral	none



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