

DATA REPORT

Novel variation at chr11p13 associated with cystic fibrosis lung disease severity

Hong Dang¹, Paul J Gallins², Rhonda G Pace¹, Xue-liang Guo¹, Jaclyn R Stonebraker¹, Harriet Corvol^{3,4}, Garry R Cutting^{5,6}, Mitchell L Drumm⁷, Lisa J Strug^{8,9}, Michael R Knowles¹ and Wanda K O'Neal¹

Published genome-wide association studies (GWASs) identified an intergenic region with regulatory features on chr11p13 associated with cystic fibrosis (CF) lung disease severity. Targeted resequencing in $n=377$, followed by imputation to $n=6,365$ CF subjects, was used to identify unrecognized genetic variants (including indels and microsatellite repeats) associated with phenotype. Highly significant associations were in strong linkage disequilibrium and were seen only in Phe508del homozygous CF subjects, indicating a *CFTR* genotype-specific mechanism.

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Lung disease in cystic fibrosis (CF) varies even among patients homozygous for the same genetic mutation (Phe508del), accounting for ~70% of CF alleles in Caucasian populations.¹ Using genome-wide association studies (GWASs) of 6,365 subjects, the International Cystic Fibrosis Gene Modifier Consortium (abbreviated Consortium) has identified genetic loci associated with CF lung disease severity, including an intergenic region on chr11p13.^{2,3} The region is flanked by genes of high potential relevance to CF lung disease, including: *EHF*, a transcription factor involved in epithelial differentiation;^{4,5} and *APIP*, a dual-function protein with roles in apoptosis and methionine salvage.^{6,7} The gene *PDHX* is also in the region and shares a promoter with *APIP*. Interestingly, the association signal in this region was stronger in 4,139 subjects homozygous for the most common CF mutation (Phe508del) compared with an analysis that included CF individuals with any severe *CFTR* genotype.²

Targeted resequencing is a viable approach to obtain a detailed genetic variant map to facilitate post-GWAS mechanistic studies. Toward this end, targeted resequencing (termed "ReSeqChr11") between the 5' end of *EHF* and the 3' end of *PDHX* (human reference genome assembly (hg19), chr11:34,641,749–35,017,674) was conducted using National Heart, Lung, and Blood Institute (NHLBI) Resequencing and Genotyping Services (rsng.nhlbi.nih.gov) (Figures 1a–c and Supplementary Figure 1). Resequencing was conducted in 377 homozygous Phe508del subjects selected from the larger Genetic Modifier Study cohort (University of North Carolina at Chapel Hill/Case Western Reserve University) that was recruited based on extremes of lung disease severity.⁸ Samples selected were balanced for gender, lung disease severity (KNORMA; equal numbers >0.4 and <0.4),⁹ and by the genotypes of the first reported³ single-nucleotide polymorphism (SNP) with lowest *P* value, rs12793173, that was found to be in strong linkage disequilibrium (LD) with the GWAS1+2² top SNP,

rs10742326 (Supplementary Table 1 and Supplementary Figures 1 and 2).^{2,3} All library preparation, enrichment, and sequencing were performed at University of Washington Genome Center at Seattle using a custom NimbleGen SeqCap probe library. Resequencing provided coverage of ~81% of the entire region with an average of 259× coverage in the sequenced regions. Gaps in coverage were because of highly repetitive features not compatible with the NimbleGen capture platform (Figure 1c).

The paired-end 49 bp sequence reads were mapped to reference genome hg19 by BWA v0.5.9-r16.¹⁰ SNP and insertion/deletion (indel) calls were made at the University of Washington using an automated software pipeline based on GATK toolkits v1.0-6125.^{11,12} The initial resequencing variant calls contained 4,800 variants, including SNPs/indels, with National Center for Biotechnology Information (NCBI) dbSNP134 annotation. Results for 94 previously genotyped SNPs were 99.98% concordant. SNP/indel calls were manually reviewed using both quality information from the variant call format (VCF) files and selected spot checks of sequencing read alignment from the binary alignment map (BAM) files, using the Integrative Genomics Viewer (IGV) genome browser.¹³ When reviewed by manual inspection in Integrative Genomics Viewer, only 40% of indels called by the GATK toolkit were verified as expected.¹⁴ The final, manually reviewed, SNP/indel calls from the 377 patient samples contained 2,991 variant calls over the resequenced region. In addition, 101 polymorphic microsatellites were called using GenoTan v0.1.5.¹⁵ We also identified a 113-bp deletion, corresponding to rs78669256 (115 bp deletion on hg19; 0.34 minor allele frequency (MAF) in our resequenced samples), that is part of a LINE element whose allele frequency and validation was unknown from single-nucleotide polymorphism database (dbSNP; Supplementary Figure 3).

The 2,991 variants were updated with NCBI dbSNP141 annotation through chromosomal location on hg19 (using

¹Marsico Lung Institute, CF/Pulmonary Research and Treatment Center, School of Medicine, University of North Carolina at Chapel Hill, Chapel Hill, NC, USA; ²Bioinformatics Research Center, North Carolina State University, Raleigh, NC, USA; ³Assistance Publique-Hôpitaux de Paris (AP-HP), Hôpital Trousseau, Pediatric Pulmonary Department, Institut National de la Santé et la Recherche Médicale (INSERM) U938, Paris, France; ⁴Sorbonne Universités, Université Pierre et Marie Curie (UPMC) Paris 06, Paris, France; ⁵McKusick-Nathans Institute of Genetic Medicine, Johns Hopkins University School of Medicine, Baltimore, MD, USA; ⁶Department of Pediatrics, Johns Hopkins University School of Medicine, Baltimore, MD, USA; ⁷Department of Pediatrics, School of Medicine, Case Western Reserve University, Cleveland, OH, USA; ⁸Program in Genetics and Genome Biology, The Hospital for Sick Children, Toronto, ON, Canada and ⁹Division of Biostatistics, Dalla Lana School of Public Health, University of Toronto, Toronto, ON, Canada.

Correspondence: H Dang (dangh@email.unc.edu)

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Table 1. Variants associated with CF lung disease severity with nominally more significant Phe508del homozygous *P* values than GWAS1+2² top SNP (rs10742326)

hg19 bp	dbSNP ID	Imputation quality (R^2)	Ref	Alt	MAF	ReSeqChr11 Phe508del/Phe508del, n = 4,139		ReSeqChr11 non-homozygous Phe508del, n = 2,226	
						β	<i>P</i> value	β	<i>P</i> value
34810590	rs374869483 ^a	0.992	TG	T	0.415	-0.117	1.81E-10	-0.045	6.79E-02
34810443	rs10836312	0.997	T	C	0.416	-0.116	2.42E-10	-0.045	7.10E-02
34808690	rs35532516 ^a	0.996	GA	G	0.416	-0.116	2.42E-10	-0.045	6.94E-02
34808486	rs1588354	0.996	T	A	0.416	-0.116	2.43E-10	-0.045	6.93E-02
34809626	rs10742325	0.996	T	G	0.416	-0.115	2.44E-10	-0.045	7.08E-02
34808920	rs10836310	0.983	T	C	0.416	-0.115	2.92E-10	-0.044	7.63E-02
34809166	rs10836311	0.983	G	A	0.416	-0.115	2.92E-10	-0.044	7.64E-02
34803694	rs11032868	0.974	G	T	0.415	-0.116	3.22E-10	-0.046	6.69E-02
34810010	rs10742326	0.981	G	A	0.416	-0.115	3.47E-10	-0.044	7.58E-02

Bold type highlights SNP of greatest significance from GWAS1+2.

Abbreviations: Alt, alternate allele; β , beta coefficient; CF, cystic fibrosis; dbSNP, database of single nucleotide polymorphisms (SNPs) and multiple small-scale variations that include insertions/deletions, microsatellites, and non-polymorphic variants; GWAS, genome-wide association study; MAF, minor allele frequency; SNP single-nucleotide polymorphism; Ref, reference allele.

^aVariants not in GWAS1+2 analysis.

ANNOVAR¹⁶ and University of California Santa Cruz (UCSC) genome databases), and 946 of them represent novel variants that tended to be rare (Supplementary Figure 4, black dots), with only 1 deletion (rs535729750 in dbSNP142) with MAF > 0.1 and only 23 deletions with MAF > 0.01.

For association with CF lung disease severity, imputation to the remainder ($n=6,365$) of the Consortium patient cohort² was performed using MACH and Minimac2, as previously described,¹⁷ except that reviewed variants from ReSeqChr11 were used as the reference set instead of 1000 Genomes variants. For microsatellite repeat polymorphisms, imputation was performed using Beagle v4.0¹⁸ because of its ability to impute multiple alleles at a single locus. Of the total 3,160 imputed variants over the resequenced region, 1,485 were of sufficient MAF (>0.01) and imputation quality ($R^2 > 0.3$, Supplementary Figure 4, vertical purple dashed line) to provide reliable tests of association with CF lung disease severity. Association testing was performed as previously described² with genotype PCs and sex as covariates. Briefly, the Consortium's quantitative lung phenotype⁹ was regressed on imputed allele dosages using linear regression for each Consortium cohort,² followed by a meta-analysis to obtain the final reported random effect *P* value. Microsatellite repeat variants were coded using the most common allele as the reference, and all other alleles as the alternative alleles before association testing.

Overall, the association *P* values are highly significant and reflect published GWAS results^{2,3} (Table 1, Figure 1, and Supplementary Figure 5). The top associated SNPs ($P < 10^{-7}$) for Phe508del homozygous subjects ($n=4,139$) were located between chr11:34,776,532 and 34,819,022, with the top SNP identified as rs374869483 (Figure 1d and blue asterisk Figure 1h), an indel located 580 bp downstream from the published top SNP (rs10742326)² (red asterisk, Figure 1h). Importantly, no significant association findings ($P < 10^{-5}$) were identified when the CF cohort was limited to non-Phe508del homozygous subjects ($n=2,226$) (Table 1 and Figure 1e), and interaction analysis between the top SNP (rs374869483) and Phe508del homozygous status was significant at $P=0.046$. This is consistent with previous results that reported a reduction in significance of the associations for this region in the entire CF cohort compared with Phe508del homozygous subjects² (Figure 1d). To evaluate association signals independent of the SNP with lowest *P* value, the imputed dosage of the conditioning SNP (rs10742326; top SNP in GWAS1+2 that is in LD with rs374869483 ($R^2=0.987$, $D'=0.995$)) was included in

the statistical model as a covariate (Supplementary Figure 6). No independent evidence of association was identified. In addition, SNP-set Kernel Association Test (SKAT) and Burden tests¹⁹ applied to the rare variants (MAF < 0.01) in *EHF*, *APIP*, *PDHX*, and several regulatory features from the 377 selected samples did not identify any significant associations (Supplementary Table 2). This analysis was limited to these subjects because rare variants cannot be imputed to the entire population.

Together, the analysis suggests that the causal mechanism is driven by one or more common variants in strong LD (Figure 2). Indeed Haploview software (default settings)²⁰ identified strong LD among all highly associated variants (Figures 1g-i and 2a-d). The top 5 haplotypes containing the 111 highly associated SNPs are shown in Figure 2e, with 3 most common haplotypes representing 38, 24, and 22% of all haplotypes. The fact that variants associated with disease are often in high LD and occur in context of local haplotype structure raised the possibility that the mechanism of regulation acts through multiple sites.

The intergenic nature of this region suggests it acts through complex gene regulatory functions. Regulatory features downloaded from SeattleSeq Annotation 138, UCSC genome browser, ENCODE (<http://www.genome.gov/10005107>),²¹⁻²⁵ and Roadmap Epigenomics servers, and annotation information over the resequenced region on chr11, were collated by chromosomal locations on hg19 (Supplementary Table 3).²⁶ Careful examination reveals complex, multiple, cell-type-specific regulatory features; for example, DNase I hypersensitive sites have been documented from human tracheal epithelium²⁷ that are distinct from those identified in K562 cells (Figure 1j vs. k). The two DNase I hypersensitive sites in human tracheal epithelium (11.2524 and 11.2525)²⁷ contain a large number of transcription factor binding sites (including four FOXA1 binding sites), as identified in chromatin immunoprecipitation sequencing assays (Figure 1l and Supplementary Table 4). In addition, epigenetic markers found in Calu3 cell line (goblet cell model from human lung adenocarcinoma) point to potential transcription enhancer activity overlapping the two FOXA1 binding segments (Supplementary Table 3). These features are potentially relevant given the known roles for FOXA1 in mucin production.^{28,29} Furthermore, the highly significant SNP rs10742325 (Table 1) has been found to show an allele-specific DNase I footprint (false discovery rate < 0.05).²⁶

HD, PJG, RGP, XG, JRS, MRK, and WKO; manuscript preparation: HD, RGP, JRS, MRK, and WKO.

COMPETING INTERESTS

The authors declare no conflict of interest.

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