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The Influence of Genetics on Cystic Fibrosis Phenotypes

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Technological advances in genetics have made feasible and affordable large studies to identify genetic variants that cause or modify a trait. Genetic studies have been carried out to assess variants in candidate genes, as well as polymorphisms throughout the genome, for their associations with heritable clinical outcomes of cystic fibrosis (CF), such as lung disease, meconium ileus, and CF-related diabetes. The candidate gene approach has identified some predicted relationships, while genome-wide surveys have identified several genes that would not have been obvious disease-modifying candidates, such as a methionine sulfoxide transferase gene that influences intestinal obstruction, or a region on chromosome 11 proximate to genes encoding a transcription factor and an apoptosis controller that associates with lung function. These unforeseen associations thus provide novel insight into disease pathophysiology, as well as suggesting new therapeutic strategies for CF.

Cystic fibrosis (CF) is a Mendelian “monogenic” recessive genetic disorder caused by mutations in the cystic fibrosis transmembrane conductance regulator (CFTR) gene (Welsh et al. 2001). There is a broad range of age-of-onset and disease activity for different organ systems in CF, including lung disease, meconium ileus, diabetes, and liver disease, even for CF patients who are homozygous for the most common mutation, F508del (Cutting 2010). Therefore, non-CFTR genetic variation and/or environmental influences must contribute to the variability of clinical phenotypes.

To enhance understanding of the pathogenesis of organ-system disease in CF and identify novel therapies, it is key to determine the magnitude of environmental and genetic effects of non-CFTR gene modifiers on clinical phenotypes. If variability in disease largely reflects environmental influences, there should be intensified focus on these factors, which might include the intensity of medical care, treatment of different types of lung infections, socioeconomic status, and effect of geographical temperature (Schechter 2004; Collaco et al. 2010, 2011). If the variability in clinical phenotypes largely reflects genetic variation in the genome, it is key to determine which genes are involved, and the mechanism of biological effects. Recent evidence indicates that there is a strong
influence of non-CFTR genetic variants on clinical phenotype in CF, but most of the genetic variation has not yet been defined (Mekus et al. 2000; Vanscoy et al. 2007).

Recent advances in genetic technology and the formation of The International CF Gene Modifier Consortium put CF at the cutting edge of gene-modifier research in “monogenic” disorders. This combination of events provides a critical opportunity for CF, because identification of the most important gene modifiers will provide insight into the mechanisms of disease pathogenesis, and offer the potential for novel prognostic approaches and therapies. This article provides an overview and update on the status of gene modifier research in CF.

HETEROGENEITY OF CF CLINICAL PHENOTYPES RELATED TO CFTR MUTATIONS

Overview

CF is the most common autosomal recessive disorder among Caucasians, and the median age of survival is only ~39 years, despite improved treatments. The cystic fibrosis transmembrane regulator (CFTR) gene encodes a 1480-amino acid protein that functions as a cAMP-mediated Cl⁻ channel, which plays a key role in hydrating airway secretions and regulating other cellular functions, including Na⁺ transport in airway epithelia (Welsh et al. 2001; Cutting 2010). Because clinical disease in CF results primarily from mutations in CFTR, CF is termed a “monogenic” disorder. There are >1500 mutations in CFTR (see Cystic Fibrosis Gene Analysis Consortium, www.genet.sickkids.on.ca/cftr/) and the most prevalent mutation results in deletion of phenylalanine at position 508 (F508del), which occurs on 70% of CF chromosomes in the U.S.

Heterogeneity of Disease Related to Mutations in CFTR

There is strong correlation between the general type of CFTR mutation and disease phenotype. Specifically, those mutations without residual function, such as F508del, are associated with pancreatic exocrine insufficiency, whereas ~10% of CFTR mutations retain some residual function and are associated with pancreatic exocrine sufficiency. Indeed, patients with at least one copy of a mutant CFTR allele with residual function have better nutritional status and milder lung disease, even though there is broad heterogeneity of disease severity (Mickle and Cutting 1993, 2000; The Cystic Fibrosis Genotype-Phenotype Consortium 1993). The development of meconium ileus, CF related diabetes (CFRD), and severe CF liver disease with portal hypertension (CFLD) is largely confined to patients with CFTR mutations with no residual function (Blackman et al. 2006, 2009b; Sontag et al. 2006; Bartlett et al. 2009; Moran et al. 2009). However, among patients carrying two mutations with no residual function, there is also a very broad range of lung disease severity, and there are strikingly different prevalences of meconium ileus, diabetes, and liver disease.

HERITABILITY OF CLINICAL PHENOTYPES

Overview

Before extensive genetic data was available, the prevailing concept was that the heterogeneity of clinical phenotypes reflected differing magnitudes of genetic versus environmental influences (Fig. 1) (Castaldo et al. 2001; Drumm 2001; Davies et al. 2005). For example, obstructive azoospermia (congenital bilateral absence of the vas deferens, CBAVD), occurs in nearly all CF males, regardless of whether the CFTR mutation has some residual function, or not; therefore, CBAVD is determined by genetics underlying mutations in CFTR (Cutting 2010). CF-related diabetes (CFRD) and severe CF liver disease with portal hypertension (CFLD) is limited to patients with two CFTR mutations without residual function (Bartlett et al. 2009; Blackman et al. 2009b; Moran et al. 2009). CFLD occurs before adulthood in the vast majority of CF patients who develop CFLD. The prevalence of CFLD is age-dependent, and occurs in ~25%–35% of CF adults. Lung disease occurs in patients regardless of whether CFTR mutations have residual function, or not, but the
range of disease severity is very great; thus, there are likely major contributions from both non-
\textit{CFTR} genetic modifiers and environmental influences (Mekus et al. 2000; Vanscoy et al. 2007).

Formal studies of genetic heritability of different clinical phenotypes have largely confirmed previous speculations (see above), and clearly show major roles for non-\textit{CFTR} genetic variation in the different clinical phenotypes (Table 1). Genes that cause many "monogenic" disorders have been described, but there is little knowledge of the other specific factors that contribute to variability of the clinical phenotype. Investigations of inbred mice provide evidence that overall genetic background has a major effect on clinical phenotype in monogenic

\begin{table}
\centering
\begin{tabular}{|l|c|c|c|c|c|l|}
\hline
Phenotype & Lung function & Persistent infection & Diabetes & Meconium ileus & BMI for height & References \\
\hline
Heritability, all subjects & 0.54–1.0 & 0.76–0.85 & \leq 1 & \leq 1 & \sim 0.6 & \sim 0.8 Blackman et al. 2006, 2008, 2009b; Vanscoy et al. 2007; Stanke et al. 2011; Green et al. 2012 \\
Heritability, F508del only & 0.56–0.86 & No estimate & No estimate & No estimate & No estimate & \sim 0.6 Vanscoy et al. 2007; Stanke et al. 2011 \\
\hline
\end{tabular}
\caption{Heritability estimates for CF-related phenotypes}
\end{table}

\textbf{Figure 1.} Schematic display of the magnitude of the effect of mutant \textit{CFTR} versus environmental influences and modifier genes. (Image courtesy of Peter Durie, Toronto.)
disorders, which is a concept that relates to non-
CFTR genetic influences on disease phenotype
in CF (Nadeau 2001). There are also CFTR-re-
lated “mono-organ” disorders, such as male in-
fertility, pancreatitis and sinusitis, which implicate complex genetic variation as important in
mono-organ disease. Therefore, the study of
modifier genes in CF is well-grounded from a
conceptual perspective.

Lung
There is a broad range of lung disease severity,
even for patients who are F508del homoygotes.
This is illustrated in a representative population
of Canadian CF patients by great variability in
forced expired volume in 1 sec (FEV1) (Fig. 2),
which is a measure of airflow obstruction that correlates with lung disease severity and survival in CF. Two twin/sibling studies have concluded that there is a strong non-CFTR genetic contribution (0.54–1.0) to the CF lung phenotype (Table 1—“Heritability”). A key feature for studying non-CFTR genotype-phenotype relationships is the ability to define a meaningful clinical phenotype in quantitative, or clear-cut binary, terms. Successful studies involve substantial effort to define the pertinent phenotype (see section below, “Challenges: Phenotypes”).

Meconium Ileus
About 15%–20% of CF neonates are afflicted by an intestinal blockage that develops in utero, a concretion composed of meconium and mucus, and characteristically forms in the distal ileum. Evidence of this condition, known as meconium ileus (MI), is often detected before birth by ultrasound as a dense region on the sonogram. Histologically, there is an increase in the number and size of goblet cells, and elongated villi and distended crypts are seen as well. MI appears to be under nearly complete control of genetic factors, as heritability of this trait approaches 100% (Table 1).

CF-Related Diabetes
CF-related diabetes (CFRD) is a common co-
morbidity in CF, with about a fifth of adoles-
cents affected and nearly half of adults (Moran et al. 2009). As a clinical entity, CFRD is distinct, although sharing properties of type 1 and type 2 diabetes (T1DM and T2DM, respectively). Type 1 features include reduced \( \beta \)-cell mass and diminished insulin secretion, but the etiology is not one of autoimmunity. Similar to type 2 diabetes, the pancreases of CFRD patients display amyloid deposition (Couce et al. 1996). Also, insulin resistance is common and ketosis

Figure 2. Lung disease severity assessed by FEV1 versus age in 1357 Canadian CF patients with biallelic loss-of-function (pancreatic insufficient) mutations. (Image courtesy of Peter Durie, Toronto.)
is rare, consistent with insulin secretion being reduced but not absent. Unlike type 2 diabetes, obesity is not a risk factor, nor are the dyslipidemia profiles characteristic of type 2 diabetes.

Whereas the morphometric and biochemical risk factors for T2DM do not appear to be risk factors for CFRD, the genetics point to a relationship between T2DM and CFRD. For example, having a relative with T2DM increases the risk of a CF patient for developing CFRD (Blackman et al. 2009a). In fact, the heritability of CFRD is high, approaching 1 (Table 1), substantially higher than for T2DM, suggesting that CFRD is much less influenced by environmental factors, such as diet, than T2DM with which it appears to share genetic etiology.

CF Liver Disease
CF liver disease begins with loss of CFTR (Cl– channel) function on the apical membrane of cholangiocytes. This dysfunction leads to defective (sluggish) bile flow and an inflammatory response with activation and proliferation of hepatic stellate cells, which results in cholangitis and fibrosis in focal portal tracts (Rowland and Bourke 2011). However, only a small subset (~5%) of patients with CF develops severe liver disease with portal hypertension (CFLD). Moreover, CFLD occurs at a median age of only 10 years, and >90% of CFLD occurs by age 20 years; thus, non-CFTR genetic variations likely contribute to risk for CFLD (Bartlett et al. 2009), but there are no formal estimates of heritability. Finally, there is also an increased prevalence of CFLD in males (2:1) over females, suggesting a role for gender and/or male/female hormones in this disorder.

MODIFIER GENETICS AND RECESSIVE MENDELIAN DISORDERS
Approaches to Study Genetic Modifiers in CF
The concept of gene modifiers in a Mendelian disorder such as cystic fibrosis shares some features, but is fundamentally different from the concept of genetic variants in non-Mendelian, complex genetic disorders, such as asthma. In complex genetic disorders, genetic variants interact with each other and the environment to cause the disorder (Van Heyningen and Yeyati 2004). In contrast, cystic fibrosis results from mutations in CFTR, and genetic variants that affect pathogenesis of mutant CFTR modify the severity of the phenotype (Drumm 2001; Cutting 2010). Genetic variation that has little effect in normal subjects may modify disease in cystic fibrosis.

Two different types of genetic studies have been used to discover gene modifiers in cystic fibrosis, i.e., case-control association studies and family-based (linkage) studies (Cutting 2010). The case-control (association) study is usually more robust for detecting common genetic variants, in part because of the larger number of subjects that can be enrolled in association studies, as compared with family-based studies. Enrollment into family-based twin/sibling linkage studies is more challenging, but it allows better definition of environmental versus genetic effects on the phenotype. Thus, complementary studies employing both the association and family-based methodologies would be optimal, and this strategy has been undertaken in cystic fibrosis by the International CF Gene Modifier Consortium.

Challenges: Phenotypes
Studies of genetic effects on phenotype require a meaningful and accurate definition of phenotype. For lung disease, FEV1 relates to clinical disease severity and outcome; therefore, the development of a quantitative lung phenotype has been developed for CF, using multiple measures of FEV1 and adjusting for mortality attrition (Taylor et al. 2011). This rigorous, quantitative lung phenotype has been used in a GWAS study of ~3500 CF patients (Wright et al. 2011). For meconium ileus, a binary (yes/no) approach is used to define phenotype, but requires careful review of source documents to be accurate (Wood et al. 2007; Sun et al. 2012). A recent publication has also determined that the heritability of chronic respiratory infection with Pseudomonas aeruginosa is very strong (0.76–0.85), which provides another lung phenotype
that can be studied for genetic modifiers (Green et al. 2012). For CF-related diabetes there is an increasing prevalence with age; therefore, genetic studies use age-of-onset ("survival") type analyses (Blackman et al. 2009b). For liver disease, a binary (yes/no) approach is used, but this requires extensive review of source documents to accurately determine which patients have severe liver disease with portal hypertension (CFLD) (Bartlett et al. 2009). Many genetic studies have been flawed by nonmeaningful or imprecise definition of phenotype, but the CF community has focused on establishing meaningful and accurate phenotypes.

IDENTIFIED MODIFIERS OF DISEASE PHENOTYPES BY CANDIDATE GENE APPROACH

Concept

Initial searches for non-CFTR genetic variants that influence pulmonary phenotype used the candidate gene approach, which is based on biological plausibility. This approach is attractive because testing a small number of carefully selected candidates avoids severe penalties from multiple comparisons during statistical analysis. However, the candidate gene approach has substantial bias, because of incomplete understanding of the pathophysiology of CF lung disease. Because an association between a genetic variant and disease phenotype can reflect relatively small differences in allele frequency, it is critical to replicate any association in a separate population, preferably several times. In CF, many candidate genes have been tested, but only a few candidates have been tested in >1000 CF patients, and only a few have been replicated (Table 2). A comprehensive review of all gene modifier studies in CF is beyond the scope of this article, so we report a summary of candidate genes that have been replicated in at least one organ in ≥2 populations entailing at least 500 individuals total (Table 2; see text below). Several recent publications review all candidate genes that have been tested in CF (Cutting 2005; Davies et al. 2005; Knowles 2006; Collaco and Cutting 2008).

Lung Disease

Lung disease in CF is linked to defective ion transport, which leads to inadequate hydration of airway mucus and results in defective mucociliary and cough clearance. Acute and chronic airway infection occurs secondary to defective mucociliary clearance, which contributes to an inflammatory response and adversely affects the phenotype. These defects of ion transport, mucus clearance, and infection and inflammation leads to the development of bronchiectasis. Therefore, genetic variation in many genes in these pathways could modify the clinical phenotype, and be associated with more severe (or milder) lung disease, and affect survival.

Multiple studies have tested whether functional variants in mannose binding lectin (MBL2) associate with lung disease severity in CF. MBL2 is biologically plausible, because of its role in innate immunity and the predilection of MBL2 deficient patients to bacterial and viral infection. Six studies involving >2000 CF patients have shown that MBL-deficient genotypes (OO or AO) are associated with worse lung function (Table 2) (Gabolde et al. 1999; Garred et al. 1999; Davies et al. 2004; Yarden et al. 2004; Trevisiol et al. 2005; Dorfman et al. 2008), and three studies reported earlier age of infection with Pseudomonas aeruginosa (Ps. a.) (Table 2) (Trevisiol et al. 2005; Dorfman et al. 2008; McDougal et al. 2010). Because severity of lung disease, Ps. a. infection, and age are correlated, a modeling framework addressed these confounding variables, and concluded that deficiency in MBL predisposes to earlier infection with Ps. a. and thereby leads to worse lung disease (McDougal et al. 2010). Several studies did not report an association of MBL deficient genotypes with CF lung disease, which might reflect large numbers of older patients and/or other factors related to bacterial infection (Cutting 2010).

Genetic variation in the endothelial receptor type A (EDNRA) has been associated with CF lung disease severity (Table 2), and replicated in three additional populations of CF patients (1478 total) (Darrah et al. 2010). The variant EDNRA (rs5335) is in the 3’ UTR, and the risk allele for worse lung disease is associated
with increased expression of EDNRA in cultured human tracheal smooth muscle cells, as well as increased proliferation of smooth muscle cells that are homozygous for the risk variant. Thus, the mechanism for adversely modifying CF lung disease likely reflects increased smooth muscle activity in airways and/or vasculature.

Transforming growth factor β1 (TGF-β1) modifies risk for airways disease in asthma and COPD, and has a key role in regulating airway inflammation and remodeling; hence, TGF-β1 has been repeatedly studied as a modifier of CF lung disease (Table 2). Alleles in the promoter (-509) and first exon (codon 10) of TGF-β1 are associated with worse lung function, and was replicated in the Genetic Modifier Study (GMS) of >1300 CF patients (Drumm et al. 2005). This discovery was independently replicated when haplotypes of alleles at -509 and codon 10 were analyzed in a separate population (Bremer et al. 2008). Further, other gene and environmental interactions are relevant to...
TGF-β1, because interactions with variants in MBL2 and second-hand smoke exposure modify TGF-β1 association with lung disease. A few small studies did not report the same association of TGF-β1 and CF lung disease (Cutting 2010).

Genetic variation in the interferon-related developmental regulator 1 (IFRD1) gene was identified as one of six candidate regions, based on study of 320 CF patients in the GMS study with “mild” ($n = 160$) versus “severe” ($n = 160$) lung disease (Table 2). Polymorphisms in IFRD1 replicated in the larger GMS sample, and then showed association in a family-based (twin and siblings) population (Gu et al. 2009). In vitro and studies in mice indicated that IFRD1 acts to modify neutrophil function during bacterial infection, via transcriptional mechanisms, but this gene has not been further replicated in additional CF populations.

IL-8 is a cytokine that is well-recognized to be associated with airways inflammation in CF, and known to mediate neutrophil chemotaxis. The association of IL-8 variants and CF lung disease was first noted in 737 CF patients and was most strongly shown in males (Table 2). The replication was verified for males in an additional 385 patients. The mechanism appears to reflect differential gene expression through IL-8 promoter variants.

**Meconium Ileus**

The heritability of MI (Table 1) clearly indicates that genes other than CFTR are involved in its etiology, but the candidate gene approach has not been applied to this CF trait. A novel candidate region approach was used however, in which a segment of mouse chromosome 7 that was found to influence intestinal obstruction in CF mice (Rozmahel et al. 1996) was examined for the corresponding region of the human genome for its effect on MI. Initial reports indicated that variants in this region did, in fact, associate with MI (Zielenksi et al. 1999), but a causative gene was never identified and subsequent studies failed to replicate this finding (Blackman et al. 2006). Although the genetic contribution to MI seems clear, the genes responsible remained elusive.

**CF-Related Diabetes**

The observation that family history of T2DM increases risk for CFRD, as described above, made obvious the need to test specific genes associated with T2DM for similar associations with CFRD. Thus, a candidate gene approach was initiated and showed that single nucleotide polymorphisms near the transcription factor 7-like 2 (TCF7L2) gene showed an association with type 2 diabetes with an estimated odds ratio of $\sim 1.5$ per allele (Blackman et al. 2009b).

In addition to showing a specific genetic relationship between T2DM and CFRD, this study showed an important concept regarding the confounding effects that nongenetic, or environmental factors may have on genetic studies, sometimes obscuring them. In this case, the nongenetic influences appear to be pharmacology, specifically steroidal anti-inflammatory drugs. Glucocorticoids are commonly used as an anti-inflammatory treatment for CF, and it is well established that a side effect of glucocorticoids is hyperglycemia, albeit transiently. Thus, this important pharmacotherapy for the lungs has the potential to mask differences in glycemic levels that would be caused by genetic variation. When controlling for the effect of this steroidal treatment by examining patients not recently using glucocorticoids, the risk associated with TCF7L2 alleles increased to 2.9 per allele (Blackman et al. 2009b).

Other evidence of similarities to T2DM is emerging as well. A screen of six genes associated with T2DM found that a polymorphism in Calpain-10 associated nominally with response to a glucose tolerance test (Derbel et al. 2006). Other T2DM genes are sure to be tested in the future, but as the effect of many of them is quite small it remains to be seen whether the cohort sizes available for CF studies will have the power to detect them.

**CF Liver Disease**

CF liver disease with portal hypertension (CFLD) occurs early in life in CF patients with biallelic loss-of-function (pancreatic insufficient) mutations in CFTR. The largest two-stage
study of CFLD focused on five candidate genes that had previously been studied in CF "liver
disease," but not all patients in previous studies
had portal hypertension (Bartlett et al. 2009).
The two-stage study showed strong association
(and replication) of the Z-allele of the α1-anti-
protease (SERPINA1) gene with CFLD in a total
of 260 patients with CFLD versus 1931 CF pa-
tients without CFLD (Table 2). The mechanism
is unclear, but probably does not reflect cumu-
lative efforts of the folding mutations in
CFTR (F508del) and SERPINA1 (Z-allele), because
these two genes are predominately expressed in
two different cell types (cholangiocytes and he-
atocytes, respectively) in the liver. Variants in
TGF-β1 were associated with CFLD in stage 1,
but did not replicate in stage 2 of the CFLD study
(Bartlett et al. 2009).

IDENTIFIED MODIFIERS OF DISEASE
PHENOTYPE BY GWAS ASSOCIATION
AND LINKAGE

The ability to perform unbiased study of rela-
tively common genetic polymorphisms across
the entire human genome in a cost-effective
fashion has recently come-of-age. To maximize
power to detect novel modifiers of CF pheno-
types using genome-wide approaches, three dif-
f erent study groups have joined forces and
created a consortium with > 3500 well-pheno-
typed CF patients to study modifiers in CF. This
consortium includes three different study de-
signs: (1) Gene Modifier Study (GMS), which
enrolled F508del homozygotes with extremes-
of-lung phenotype (Drumm et al. 2005); (2) Ca-
nadian Genetic Study (CGS), which enrolled
population-based CF patients; and (3) family-
based, twins and siblings (TSS), which enrolled
patients from families with more than one CF
child. The GMS and CGS are best-suited for as-
sociation analysis of unrelated patients, whereas
the TSS (family-based) study can be used for
both linkage and association analyses. Using
this complementary study design allows for dis-
covery by association (or by linkage) and then
testing for replication in the alternate popula-
tion(s).

Lung Disease

The combined genome-wide association and
linkage study (GWALS) genotyped 3467 CF pa-
tients (Illumina 610 Quad platform) to identi-
fy genomic regions associated with CF lung
disease severity. The strongest association sig-
nal ($p = 3 \times 10^{-8}$) was located at chromo-
some11p13, intergenic to EHF and APIP and
replicated in the TSS (family-based) patients
(Fig. 3) (Wright et al. 2011). EHF is an epithelia-
specific Ets transcription factor, and may be
an important regulator of differentiation under
stress conditions and inflammation. APIP is a
known inhibitor of apoptosis, and eQTLs for
APIP in lymphocytes suggests the risk SNP in
this region may be associated with increased ex-
pression of APIP, implying that inhibition of
apoptosis worsens CF lung disease. This is con-
gruent with the notion that inhibition of apo-
ptosis in airways delays clearance of neutrophils
and leads to a hyper-inflammatory state (Dib-
bert et al. 1999; Harris et al. 2005). Inhibition of
apoptosis may also contribute to goblet cell
metaplasia, which is a feature of CF airways dis-
ease (Dibbert et al. 1999; Harris et al. 2005).
Polymorphisms in the intergenic region of
chr11p13 may also affect regulatory domains,
which could alter expression of EHF and APIP
and other genes in the region, including ELF5,
PDHX, and CD44. Studies are ongoing to elu-
cidate the mechanism of disease pathogenesis
from genomic polymorphisms at chromo-
some11p13.

There was also a near-significant genome-
wide association at the chromosome 6p21.3
HLA class II DR locus. Genetic variation at
this locus has been widely recognized to play a
role in multiple infections and inflammatory
disorders, which would be relevant to the in-
fec tions and inflammatory nature of CF airway
disease (Handunnetthi et al. 2010). This locus is
very genetically heterogeneous, and studies are
underway to define the specific DR allele that
is associated with severity of CF lung disease.

There were five other regions of suggestive
association (Wright et al. 2011), which will be
further analyzed by an ongoing GWAS in an
additional ~3500 CF patients. Two previously
reported modifiers of CF lung disease, TGF-β1 and IFRD1, did not achieve genome-wide significance. However, TGF-β1 achieved p-values of $10^{-3}$ to $10^{-4}$ in GMS patients, depending on the covariates used in the analyses.

Linkage analysis of 486 sibling pairs from the family-based study identified a robust linkage signal at chr20q13.2 (LOD = 5.03). A region below the linkage peak was analyzed for association in the GMS and CGS patients, and replicated under a region-wide analytical approach ($p = 1/10^8$). The chr20p13.2 linkage region contains five genes (CBLN4; MC3R; CASS4; CSTF1; AURKA) that are expressed in fetal or adult lung, or bronchial epithelia. MC3R is of particular interest, because it has been implicated in regulation of energy balance and modulating neutrophil accumulation in lung inflammation, which are both relevant to CF lung disease (Savastano et al. 2009). Studies are underway to define the mechanism of lung disease in CF that reflects the linkage signal at chr20p13.2.

**Meconium Ileus**

The high heritability of MI certainly suggested that genes other than CFTR must be involved, and the availability of high density markers throughout the genome provided the opportunity to carry out linkage studies at high resolution. Using this approach, Blackman and colleagues found by genome-wide analyses several regions of suggestive linkage (logarithm of the odds of linkage >2.0) for contributing to, or protecting from development of, meconium ileus. These regions were found on chromosomes 4q35.1, 8p23.1, and 11q25 (MI contributing loci) and chromosomes 20p11.22 and 21q22.3 (protective loci) (Blackman et al. 2006).

The locus on chromosome 8 is a gene-dense region and was followed up by haplotype analysis and identified a three polymorphism haplotype that most strongly associated and corresponded to a gene, MSRA, encoding an enzyme, methionine sulfoxide reductase A, that maintains the amino acid methionine in a reduced state. This candidate was then examined functionally by assessing its effect on obstruction in a CF mouse model lacking functional MsrA, and found to dramatically decrease the incidence of obstruction compared to CF mice with functional MsrA (Henderson et al. 2012).

Another linkage study provided evidence for MI-influencing loci on chromosome 12p13.3. Using densely spaced markers in the 12p13.3 region in a cohort of pancreatic insufficient Canadian CF patients with and without MI, one gene, ADIPOR2 in MI ($p = 0.002$), was implicated. The frequency of alleles in this gene was
distorted from Hardy-Weinberg proportions, but only in the MI group. The deviation from expected proportions suggests a causative effect and the association was replicated in an independent sample of CF families. A second locus, but this one protective, was found on chromosome 4q13.3 and one candidate gene in the region, SLC4A4, a member of the solute transport family of proteins, showed evidence of association ($p = 0.002$) (Dorfman et al. 2009).

Association studies can also be used to identify genes involved in meconium ileus, and a novel modification of the association approach has identified several candidate genes. This approach, termed “informed GWAS” incorporates functional annotation into the analyses, giving higher import to those genes with a particular function attributed to them. Using a model in which genes influencing epithelial function were the focus, polymorphisms near SLC6A14 on the X chromosome and SLC26A9 on chromosome 1 accounted for $\approx 5\%$ of variance in MI and these associations were replicated in an independent sample of affected individuals (Sun et al. 2012).

Other

Genome-wide association data have not yet been reported for CFRD or CFLD, although the Consortium is analyzing data from GWAS1, and awaiting data from GWAS2.

CONCLUDING REMARKS

The ability to identify genes, or genomic regions, that influence disease processes hold enormous promise for better understanding those processes and for manipulating them for therapeutic gain. However, an important caveat of genetic screens is that they may identify genes involved in a process, but as such they are only markers; they do not provide information about the mechanisms by which the variants exert their phenotypic effects. Thus, for the results of any genetic study to have more than prognostic value, the genes must be followed up by functional studies to have clinical utility.

The potential for clinical application is not only great, it is near. There are already approved drugs that target the protein product encoded by several of the genes identified (Table 2), including antagonists to the endothelin and angiotensin receptors, drugs that alter TGF-B1 signaling and small molecule screening provides the potential to rapidly identify new drugs that will target these gene products or the pathways in which they act.

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