Genetic epidemiology of pulmonary function

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Respiratory diseases are major threats to human health.¹ It is believed that chronic obstructive pulmonary disease (COPD) is both environmental and genetic², however, specific genetic factors in the development of COPD have not been clearly identified, except for protease inhibitor types. Alpha₁-antitrypsin deficiency is rare in the general population⁴, and accounts for less than 2% of the cases of COPD.⁶, The molecular genetics of COPD has recently been reviewed by Barnes.⁸

Pulmonary function measures are the most important phenotypes of COPD. The respiratory muscles, the thorax, and the lungs are the components of the ventilatory apparatus which can be evaluated by appropriate function tests.9 The dynamic functional capacity of the ventilatory apparatus can be assessed by the volumetime or flow-volume manoeuvres. Airflow rates and volumes inhaled or exhaled over specific time intervals provide information on the flow resistive properties of the airways.¹⁰ These pulmonary function measures predict the development of lung diseases¹¹⁻¹³ and overall mortality.¹⁴⁻¹⁸ While the environmental determinants of pulmonary function have been extensively studied-for example, smoking and ambient air pollution-the genetic determinants have recently received increasing attention. Genetic epidemiological studies of pulmonary function are of potential importance in understanding normal pulmonary function and the aetiology and prevention of COPD and other respiratory diseases. This paper reviews

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Table 1 Family studies in pulmonary function

the familial aggregation and segregation of pulmonary function and presents evidence for different influences of heredity on airway function, lung volume, and airway-parenchymal dysanapsis (relative airway size to lung size). Some methodological issues related to segregation analysis are also discussed.

Family aggregation

FAMILY STUDIES

A number of family studies have provided evidence for familial resemblance of pulmonary function measures.¹⁹⁻³⁰ Studies have consistently shown significant parent-offspring and sibling-sibling correlations in lung volume and flow rate measures (table 1). Most studies found that spousal correlations in pulmonary function were trivial, although Higgins and Keller²⁷ reported a small but significant correlation in forced expiratory volume in one second (FEV₁) between spouses, and Kauffmann *et al*²¹ found significant spousal correlations in residual forced vital capacity (FVC), FEV₁, and forced expiratory flow between 25% and 75% of the vital capacity (FEF_{25-75%}).

One study found that familial correlations in pulmonary function were dependent on familial resemblance of body habitus and were no longer significant after the ponderal index (height/weight^{1/3}) was taken into consideration.²⁸ Other studies found that the familial correlations remained significant after adjustment for both height and weight²²⁻²⁶ or adjustment for height alone.²¹ Over-adjustment could

	.			Correlations				
Reference	Location and year	Study subjects	Lung function indices	ρ_{sp}	$ ho_{\scriptscriptstyle mo}$	$ ho_{fo}$	$ ho_{sib}$	Comments
Tager et al19	USA, 1976	148 households,	FEV ₁ , %	0.04	0.18*	0.25*	0.26*	FEV ₁ (%) less correlated than
		469 subjects	FEV ₁ score	0.05	0.11	0.16*	0.19*	FEV ₁ score
Schilling et al20	USA, 1977	376 families, 816 children and their	rFVC	0.07	0.19*	0.12*	0.27*	Separate parent-offspring
		parents	rFEV ₁	0.07	0.17*	0.11*	0.21*	correlations; smaller ones are
		-	rVmax ₅₀	0.07	0.15*	0.12*	0.16*	selected
Kauffmann et al ²¹	France, 1989	945 families, 1160 children and their	rFVC	0.18*	0.26*	0.19*	0.30*	Significant spousal correlations
		parents	rFEV ₁	0.20*	0.26*	0.15*	0.34*	
		-	rFEF _{25-75%}	0.23*	0.21*	0.16*	0.27*	
Coultas et al ²²	USA, 1991	733 households, 336 spouse pairs,	rFVC	0.11	0.26*	$=\rho_{mo}$	0.37*	Parent-offspring correlations: 6-17
		1059 parent-child pairs, 412 sib pairs	rFEV ₁	0.10	0.22*	$=\rho_{mo}$	0.24*	year group
Chen et al ²³⁻²⁵	Canada, 1996,	309 families, 1045 subjects	rFVC	0.08	0.17*	0.18*	0.35*	First degree relative correlations
	1997, 1998		rFEV ₁	0.04	0.11*	0.18*	0.11*	were not significantly different for
	-		rFEF _{25-75%}	0.11	0.15*	0.22*	0.19*	rFEV,, rFEF25 75% and rVmar50/FVC
			rVmax ₅₀ /FVC	0.11	0.16*	0.25*	0.27*	25 7570 max50
Givelber et al ²⁶	USA, 1998	1408 families, 5003 adult subjects	rFEV ₁	0.05	0.19*	0.12*	0.22*	Sib-sib correlation was greater than parent-offspring correlation

 FEV_1 = forced expiratory volume in one second; FVC = forced vital capacity; $FEF_{25-75\%}$ = forced expiratory flow between 25% and 75% of the vital capacity; $Vmax_{50}$ = maximal expiratory flow rate at 50% of vital capacity; ρ_{sp} = spousal correlation; ρ_{mo} = mother-offspring correlation; ρ_{fo} = father-offspring correlation; ρ_{sb} = sibling-sibling correlation; r = residual. *p<0.05.

Table 2 Twin studies in pulmonary function

	Location and year	Pairs of twins		I	Intrapair (differend	correlation ve)	
Reference		MZ	DZ	 Lung function indices 	MZ	DZ	Comments
Man and Zamel ³⁴	Canada, 1976	10	6	VC (l) Vmax ₆₀ (l/s) Vmax ₄₀ (l/s)	(0.30) (0.37) (0.30)	$(0.55)^{*}$ $(1.23)^{*}$ (0.63)	Difference seems larger for the geometry of larger airways than for smaller ones
Hubert et al ³²	USA, 1982	127	141	FVC FEV,	0.66† 0.62†	0.08 0.25†	Correlation in FVC between DZ twins was not significantly different from zero
Redline <i>et al</i> ³³	USA, 1987	256	158	FVC FEV ₁ FEF _{25-75%} FEV ₁ /FVC FEF _{25-75%} /FVC	0.76† 0.71† 0.52† 0.55† 0.53†	0.39† 0.16 0.21 0.18 0.21	Adjusted for body size and smoking
Ghio et al ³⁶	USA, 1989	47	27	$ \begin{array}{c} FVC (1) \\ FEV_1 (1) \\ FEF_{25-75\%} (1/s) \end{array} $	(0.28) (0.26) (0.58)	(0.36) (0.34) (0.77)	Differences between MZ and DZ were not significant after adjusting for height

 $MZ = monozygotic twins; DZ = dizygotic twins; VC = vital capacity; FEV_1 = forced expiratory volume in one second; FVC = forced vital capacity; FEF_{25-75\%} = forced expiratory flow between 25% and 75% of the vital capacity; Vmax₄₀, Vmax₆₀ = maximal expiratory flow rate at 40% and 60% of vital capacity. *p<0.05, difference between MZ and DZ.$

†p<0.05, difference from zero.

be one reason for the disappearance of familial aggregation of pulmonary function,²² and different analytical methodology could be another.

Chen et al²³⁻²⁵ compared the motheroffspring, father-offspring, and sibling-sibling correlations in various pulmonary function measures. No significant differences were found in the correlations of airway function measures including the FEV1, FEF25-75%, and maximal expiratory flow rate at 50% of vital capacity (Vmax₅₀). For FVC, however, the sibling-sibling correlation was greater than the parent-sibling correlation, which was consistent with the results from other studies.20 22 29 This additional resemblance between siblings may be due to shared sibling environment and environmental factors may have different impacts on the lung volume measure than on the flow rate measures.²⁹ Givelber et al²⁶ found that the sibling-sibling correlation in FEV₁ was greater than parent-offspring correlations. Since the FEV, was measured much earlier for the parents (1948-52) than for the offspring (1971-74), measurement error could be larger for data from the parents than those from the offspring due to outmoded spirometric techniques.²⁶ The results are not consistent in terms of differences between mother-offspring and father-offspring correlations for various pulmonary function measures. Givelber et al²⁶ suggested a greater mother-offspring correlation in FEV, compared with the fatheroffspring correlation. Coultas et al²² and Chen et al²³⁻²⁵ found that there was no significant difference between mother-offspring and fatheroffspring correlations.

TWIN STUDIES

Comparisons of monozygotic (MZ) and dizygotic (DZ) twins can be used to assess the relative importance of genetic and environmental effects, since MZ twins share 100% of genes while DZ twins share only 50%.³¹ A higher degree of similarities in MZ twins than in DZ twins suggests a genetic influence on pulmonary function phenotypes. Studies have shown that the intrapair correlation of pulmonary function measures was greater³² ³³ and the intrapair difference was smaller in MZ twins than in DZ twins.³⁴ ³⁵ Only one study³⁶ showed that the intrapair difference in pulmonary function between DZ twins was not significantly larger than that between MZ twins (table 2).

Two studies of MZ twins have suggested that genetic factors are important in determining susceptibility to airway dysfunction from cigarette smoke.^{37 38} Webster *et al*³⁷ studied 45 pairs of MZ twins and found that the intrapair difference in values of maximal expiratory flow rate at 60% of the vital capacity was small and was similar for smoking pairs and non-smoking pairs. There was a large difference between twins discordant for smoking. In a study of MZ twins raised apart Hankins et al38 examined the FEV₁, FEF_{25-75%}, and maximal expiratory flow rate at 70% of vital capacity and found similar results. However, the absence of DZ twins as a comparison group limits the ability to make strong inferences from these studies.

Genetic heritability

Both family and twin studies have clearly shown that pulmonary function including flow rate and lung volume measures are familial. The reason for the familial aggregation can be environmental, genetic, or both. Genetic heritability, which is the proportion of the genetic variance to the total phenotypic variance in a defined population, can be used to quantify the degree of genetic contributions. The genetic variance can be further divided into additive genetic variance, dominance variance, and epistatic variance (interlocus interaction). Heritability in the narrow sense, which is the proportion of additive genetic variance to the total phenotypic variance, is used to measure possible genetic effects not due to major gene segregation.

A number of family studies have examined the degree to which the observed familial aggregation of pulmonary function is attributable to genetic factors and have shown a moderate degree of heritability for various pulmonary function measures (table 3). Based on the data of 439 subjects from 108 families of patients without pulmonary disease, Astemborski *et al*³⁹ found that additive genetic variation accounted for 28% of the variation in

 Table 3
 Additive genetic heritability of lung function based on data from family studies

Reference	Location and year	Study subjects	Lung function indices	Additive genetic heritability	Comments
Lewitter and Tager ⁴¹	USA, 1984	404 families, 602 parents,	FEV ₁ score	42-47%	Path analysis; consistent over time
Devor and Crawford ²⁹	USA, 1984	756 children 96 families, 307 subjects	FEF _{25-75%} score rFVC rFEV	42–47% 20% 17%	Path analysis
Astemborski <i>et al</i> ³⁹	USA, 1985	108 families, 439 adults	rFEV ₁ rFEV ₁ /FVC	28% 24%	Variance components analysis; adult study population
Beaty <i>et al</i> ⁴⁰	USA, 1987	158 families, 781 subjects	rFEV ₁ rFEV ₁ /FVC	9% 25%	Variance components analysis; ascertained through a proband with obstructive lung disease
Cotch et al ⁴²	USA, 1990	384 families, 978 subjects	rFEV ₁ (cross sectional) rFEV ₁ (longitudinal)	36% 40%	Path analysis; no inter-generational differences
Coultas <i>et al</i> ²²	USA, 1991	733 households, 336 spouse pairs, 1059 parent-child group pairs, 412 sib pairs	rFVC (non-smoking parents) rFVC (smoking parents) rFEV _i (non-smoking parents) rFEV _i (smoking parents)	43% 42% 65% 44%	Path analysis; no substantial changes based on age and smoking status
Chen et al ^{23–25}	Canada 1996, 1997, 1998	309 families, 1045 subjects	rFEV ₁ rFEV ₅₀ rFEF _{25-75%} rVmax ₅₀ /FVC	26% 36% 34% 40%	Class D regressive model

 FEV_1 = forced expiratory volume in one second; FVC = forced vital capacity; $\text{FEF}_{25-75\%}$ = forced expiratory flow between 25% and 75% of the vital capacity; Vmax_{50} = maximal expiratory flow rate at 50% of vital capacity; r = residual.

residual FEV, and 24% of the variation in residual FEV₁/FVC. The estimators were smaller among families of patients with airway obstruction disease.40 Four studies used the path analysis approach to identify hereditary and environmental sources of familial aggregation for pulmonary function traits. Lewitter et al⁴¹ studied 404 nuclear families including 602 parents and 756 children and found that 42–47% of the variability in FEV_1 and $\text{FEF}_{25-75\%}$ could be explained by underlying genetic differences among the individuals. Another analysis by Cotch $et al^{42}$ showed a similar estimate of heritability of 36-40% for FEV₁ and no significant difference between white and black individuals. Coultas et al²² found an increased genetic variance in smokers. The genetic variances for FVC and FEV₁ were 10% and 25% greater for smokers than for non-smokers, respectively.22 The heritability estimates of pulmonary function measures were lower in another study. Based on the data from 305 men and 339 women Devor and Crawford²⁹ estimated that approximately 20% and 17% of the variation in FVC and FEV₁, respectively, was due to the transmission from parents to offspring. Smoking and age could alter the familial aggregation of pulmonary function, and different study designs may be the reason for the discrepancy.22 Shared heritable factors might not only influence the lung growth and development, but also the decline in pulmonary function in adults.²² In the recent Humboldt Family Study of 309 nuclear families Chen *et al*²³⁻²⁵ used a class D regressive model and estimated the additive genetic heritability as 26-40% for FEV₁, Vmax₅₀, FEF_{25-75%}, and Vmax₅₀/FVC.

The heritability estimates for different measures of pulmonary function in twin studies have been inconsistent. Hubert *et al*²⁹ studied 127 MZ and 141 DZ male twin pairs aged 42–56 years and estimated the heritability to be as high as 77% for FEV₁. In a study of 256 MZ and 158 DZ adult twins Redline *et al*³³ found that 40–75% of the measured variability in pulmonary function was accounted for by genetic influences. In another study, however, Ghio *et al*³⁶ found that the heritability was not significant after adjustment for height in 74 university student pairs of twins with an average age of 20 years. Twin studies usually provide an inflated estimate of heritability4 because twins share very similar, if not the same, exposures in utero, and share a more homogeneous living environment than other individuals. Since heritability is the ratio of the genetic variance to the total variance, heritability increases with decreasing variance due to environment. The heritability estimates can be more biased if the effects of environmental factors are more similar in MZ twins than in DZ twins. Some studies have shown that MZ twins have a greater concordance in smoking habits than DZ twins.^{32 44} Correlational analysis of the distribution of given traits within family members is one way to increase the robustness of the twin data analysis.45

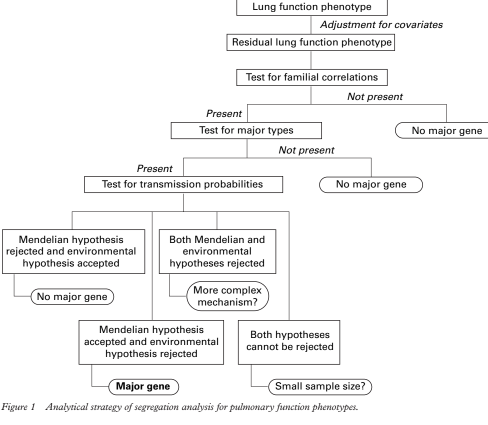
Heritability is a population-specific parameter and is affected by the environment in which the population developed. In addition, if there is an interaction between genotype and environment—for example, smoking may alter the genetic effects on pulmonary function—it is almost impossible to separate the genetic variance and environmental variance completely. Because of these limitations, heritability estimation should be explained with caution.

Major genetic effects on pulmonary function

SEGREGATION ANALYSIS

Genetic effects may be the consequence of a single gene (a major gene), a small number of genes (oligogenes), or a large number of genes each with a small effect (polygenes). In segregating families the relatively large effects of major genes should be detectable using the tools of segregation analysis, while the more general predictions of polygenes predict the overall patterns of correlation among relatives. The classical segregation analysis is used to identify Mendelian ratios when a phenotype is controlled by a major gene, which is traditionally assumed to result from segregation at a single locus having two alleles, A and B. The likelihood method is frequently used in





segregation analyses, and other approaches are also proposed including generalised estimating equations.⁴⁶

There are various strategies of model evaluation. Figure 1 gives an example of the analytical strategy for a segregation analysis. If a pulmonary function phenotype is familial, the next step is to determine mixtures of phenotypic distribution. Before there is evidence of Mendelian transmission, a more general term called "ousiotypes" or "types" has been suggested to describe the mixtures of distribution.⁴⁷ A key assumption is that individuals represent different ousiotypes or essential types that may reflect different genotypes for Mendelian models. The parameters of transmission probabilities can therefore be estimated, which are the probabilities of a parent transmitting the A allele to an offspring. Under Mendelian transmission, $\tau(AA) = 1$, $\tau(AB) = 0.5$, and $\tau(BB) =$ 0. A non-transmitted environmental effect was obtained with the three transmission probabilities being equal $(\tau(AA) = \tau(AB) = \tau(BB))$. The process of segregation analysis involves testing a series of models of inheritance, including Mendelian models such as dominant and recessive models plus non-genetic models, to identify the best fitting and most parsimonious model for a given set of family data.

AIRWAY FUNCTION

Based on data from 85 families with COPD and 56 families without pulmonary disease, Rybicki *et al*⁴⁸ used the class A regressive model and found that there were major genetic effects on FEV₁, and the major gene effects could explain all of the familial correlations for FEV₁ in families ascertained through a COPD proband. In families of patients without pulmonary disease, however, there were no familial correlations for FEV₁ and therefore no evidence of genetic control of FEV₁.⁴⁸ These results suggested substantial actiological heterogeneity in the control of FEV₁ between the families with COPD and those without the disease. The reasons for the lack of familial correlations in the families of those without COPD and its discrepancy with other studies were not discussed in the report.⁴⁸

The class A regressive model makes an assumption that siblings are correlated only through common parentage. This restriction, in the absence of a major gene, may lead to false inference of a major gene.⁴⁹ Chen et al^{23 25} studied 309 young families and used the class D regressive model in the segregation analysis, which allows additional correlation among siblings and is characterised by equal siblingsibling correlations. The data have suggested that FEV_1 , $\text{FEF}_{25-75\%}$, and Vmax_{50} are more likely to be controlled mainly by multiple loci-namely, many independent genes-each contributing in an additive fashion and/or common environmental factors are responsible for the familial resemblance of airway function. In a recent report of 5003 subjects from 1408 families in the Framingham study Givelber et al²⁶ provided consistent results. The most parsimonious model for FEV₁ included nontransmitted major types and residual familial correlations, and the Mendelian hypothesis was rejected.23 Based on the data from 309 families (1163 individuals) in the Tuscon children's respiratory study Holberg et al⁵⁰ also suggested polygenic control of FEV_1 or common environmental factors resulting in the familial aggregation of the trait; however, genetic heterogeneity might exist between families with and without

asthmatic members.⁵⁰ Silverman *et al*⁵¹ studied α_1 -antitrypsin deficient individuals in 44 nuclear families and found that there was an additional major gene other than the Pi locus influencing FEV₁. However, the major gene effect diminished after adjustment for pack-years of smoking.

LUNG VOLUME

Based on the data from 309 nuclear families Chen et al^{24} performed a segregation analysis for FVC, a measure of lung volume. Models with both major types and familial correlations gave the best fit for the data. However, neither Mendelian nor no parent-offspring transmission hypotheses were rejected. Heterogeneity may exist between families. The authors calculated the likelihood under the Mendelian model $(L_{\mbox{\tiny Mendelian}})$ and the environmental model $(\mathbf{L}_{\mbox{\tiny environmental}})$ and used the ln-likelihood values to sort families into groups that support one model of inheritance over another. In a subset of 196 families with a $ln(L_{Mendelian}/L_{environmental})$ value greater than zero, the families suggested a Mendelian gene leading to lower values of FVC, and the single locus explained all the familial aggregation of residual FVC. In the other 113 families in whom the ln(L_{Mendelian}/ L_{environmental}) value was less than zero the Mendelian hypothesis could not be rejected and the Mendelian model showed that a single locus accounted for all familial correlations except for the sibling-sibling correlation. However, the environmental hypothesis could not be rejected for this subgroup of families although the Mendelian model had a better fit than the environmental model based on the values of the Akaike's information criterion (AIC, see later).²⁴ The approach of dividing the families into two groups based on an individual likelihood ratio might remove certain confounding effects and increase the statistical power of detecting a major gene effect.²⁴

AIRWAY-PARENCHYMAL DYSANAPSIS

Green et al¹¹ found a low correlation between lung volume and maximal expiratory flow, and no obvious relationship between static lung recoil and Vmax₅₀, suggesting that there are substantial differences between individuals in airway size and function that are independent of lung size. Disproportionate but physiologically normal growth of airway and parenchymal components suggests a "dysanaptic" growth which may have an embryological basis.¹¹ The concept of airway-parenchymal dysanapsis was advanced by Mead⁵² who reasoned that subjects with large lungs do not necessarily have larger airways than those with small lungs. He used $Vmax_{50}/(VC \times Pst(L)_{50})$ as an index of airway-parenchymal dysanapsis in which Vmax₅₀ is the maximal expiratory flow rate at 50% of total volume and $Pst(L)_{50}$ is the maximal flow static recoil pressure characteristic at 50% of vital capacity (VC). Green *et al*¹¹ have shown that lung static recoil contributes

little to the variability between individuals and that the major variability in maximum flows is attributable to airway dimensions. The correlation between $Vmax_{50}/VC$ and $Vmax_{50}/(VC \times Pst(L)_{50})$ is high, ranging from 0.78 to 0.84.⁵³

The dysanapsis is a general phenomenon. Airway-parenchymal dysanapsis has been observed both in $adults^{12}$ ⁵⁴⁻⁵⁹ and children.^{60 61} Martin *et al*⁶⁰ found substantial interindividual variability of maximal expiratory flow rates relative to lung volumes during early childhood which remained constant during growth, suggesting that the dysanapsis originates in early childhood.

Chen *et al*²⁵ examined the major gene effects on the ratio of $Vmax_{50}$ to FVC. There was evidence for mixtures of distribution while the polygenic and sporadic model did not give a good fit to these data. The transmission of ousiotypes for $Vmax_{50}$ /FVC was not different from the Mendelian expectation, and the no parent-offspring transmission hypothesis was rejected, suggesting that there is a single locus gene or a cluster of genes working in unison to determine $Vmax_{50}$ /FVC.

It has been suggested that airwayparenchymal dysanapsis might have relevance for the pathogenesis of obstructive airway disease.^{11 12} A study by Litonjua *et al*¹³ indicated that airway-parenchymal dysanapsis, as measured by MMFR/FVC, was a significant predictor for the degree of bronchial hyperresponsiveness.

METHODOLOGICAL ISSUES

Adjustment for covariates

Pulmonary function phenotypes are most likely to be multifactorial, controlled by both genetic and environmental factors. Various factors including host characteristics, environmental factors, and history of respiratory symptoms and disease influence these pulmonary function measures. Adjustment for these variables is always a challenge. One approach is to include these variable covariates in regressive models; however, most of the variables tend to have inconsistent effects on pulmonary function measures in different age and sex groups. For example, body weight shows both "muscularity effect" (increase in pulmonary function with increasing weight) and "obesity effect" (decrease in pulmonary function with increasing weight). There is more "muscularity effect" than "obesity effect" in children and young adults but more "obesity effect" than "muscularity effect" in older adults, which is sex related.⁶² Cigarette smoking has a sex related effect on pulmonary function measures.63 64 Age itself positively predicts pulmonary function in children and young adults and negatively predicts pulmonary function in middle aged and older adults. Clearly, these variables cannot be appropriately adjusted by including them in the same regressive models in a segregation analysis. Preadjustment for the variables in different age and sex groups is therefore preferred. The adjusted values are used to fit models of inheritance. The relative importance of the covariates in relation to pulmonary function phenotype varies across age and sex groups, and this is reflected by the proportion of variation explained by these factors. Most studies of pulmonary function preadjusted these covariates including smoking, one of the most important determinants, in the segregation analyses.^{23-26 48 50}

It is debatable whether or not to adjust for history of respiratory symptoms and disease. A history of respiratory symptoms and disease may reduce pulmonary function but it can also be a surrogate measure for the effects of smoking and other environmental factors on the respiratory system. Adjustment for these variables may eliminate some confounding effects but may also reduce the variance of pulmonary function phenotypes unnecessarily. Comparisons of adjusted and unadjusted results are always helpful.

Selection of parsimonious models

The likelihood ratio test is usually used to select the most parsimonious model, which is minus twice the difference in the log, likelihood (In L) between models before and after reducing parameters. The test is based on a comparison of strictly hierarchical models. For several alternative non-hierarchical models the better fitting model is considered with a lower value of the Akaike's information criterion (AIC = $-2 \times \ln L + 2 \times \text{number of parameters}$ estimated).65 Although the AIC is not a statistical test and therefore provides no statistical inference, it is useful in identifying the most parsimonious model.

Statistical power

In segregation analysis a series of models of inheritance are fitted and the most parsimonious is chosen to explain the familial aggregation of a pulmonary function phenotype. Tests are typically based on "goodness of fit" measures and a type II error occurs when the genetic model is incorrect, but statistical testing fails to reject it because of small sample size.²⁸ Statistical power for segregation analysis is related to the size of gene effect and sample size.⁶⁶ Nuclear families with larger sibships are generally more informative³¹; however, the total number of subjects rather than sibship size per se may have more influence on the power.67 Large sample size increases the power in discriminating the completing model.

Gene-environmental interaction

A previous study has documented geneenvironment interactions in COPD.68 Another study has suggested that a gene-environment interaction may influence pulmonary function.⁴⁷ The pulmonary function phenotype expression of a gene may therefore depend on environmental variables such as smoking. Ignoring gene-environment interactions may result in underestimating the genetic effects on quantitative traits.69

Conclusions and future directions

Both family studies and twin studies have shown familial aggregation of various measures of pulmonary function. There is a moderate degree of genetic heritability for these pulmonary function measures. However, genetic factors may have different influences on phenotypes of airway function, lung volume, and airway-parenchymal dysanapsis. Airway function phenotypes are more likely to be controlled by many loci with no major gene effects and/or are due to common environmental fac-"normal" families. Aetiological tors in heterogeneity may exist in families with COPD or asthma and heredity may have different effects on normal airway function and airway dysfunction. There is evidence of major gene control of phenotypes of airway-parenchymal dysanapsis and lung volume. It would be interesting to investigate further the different effects of heredity on various pulmonary function phenotypes and their potential linkage to diseases of the lung. In particular, researchers should seek biological evidence for major gene controls of lung volume and airwayparenchymal dysanapsis of the lung.

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- 1 American Thoracic Society. Future directions for research on diseases of lung. Am J Respir Crit Care Med 1995;152:1713–35.
- Snider GL. Chronic obstructive pulmonary disease: risk factors, pathophysiology, and pathogenesis. Annu Rev Med 1989;40:411-29. 1989;40:411–29. 3 Higgins M. Risk factors associated with chronic obstructive
- Hung disease. Am NY Acad Sci 1989;624:7–17.
 4 Horne SL, Tennent R, Lovegrove A, et al. Pi type MZ and an increased risk of pneumonia. Clin Invest Med 1984;7:85-
- 5 Horne SL, Chen Y, Cockcroft DW, et al. Risk factors for reduced pulmonary function in women: a possible relation-ship between Pi phenotype, number of children, and pulmonary function. *Chest* 1992;**102**:158–63.
- Cohen BH. Chronic obstructive pulmonary disease: a challenge in genetic epidemiology. Am J Epidemiol 1980;112: 274 - 88
- Snider GL. Molecular epidemiology: a key to better understanding of chronic obstructive lung disease. Monaldi Arch Chest Dis 1995;50:3-6.
- 8 Barnes PJ. Molecular genetics of chronic obstructive pulmonary disease. *Thorax* 1999;54:245–52.
- 9 Wilson AF, ed. Pulmonary function testing: indications and interpretations. Orlando, Florida: Grune & Stratton, 1985.
- Altose MD. Practical aspects of pulmonary function testing. In: Baum GL, Wolinsky E, eds. Textbook of pulmonary diseases. 4th ed. Boston, Toronto: Little, Brown and Com-pany, 1989:101–13.
- 11 Green M, Mead J, Turner JM. Variability of maximum expiratory flow-volume curves. J Appl Physiol 1974;37:67– 73.
- 12 Brooks LJ, Byard PJ, Helms RC, et al. Relationship between lung volume and tracheal area as assess by acoustic reflec-tion. J Appl Physiol 1988;64:1050-4.
- 13 Litonjua AA, Sparrow D, Weiss ST. FEF₂₅₋₇₅/FVC as a measure of airway-parenchymal dysanapsis predicts bronchial hyperresponsiveness. Am J Respir Crit Care Med 1996;153(Suppl):A428.
- 14 Ashley F, Kannel WB, Sorlie PD, et al. Pulmonary function: relation to aging, cigarette habit, and mortality. Ann Intern Med 1975;82:739-45.
- 15 Beaty TH, Cohen BH, Newill CA, et al. Impaired pulmonary function as a risk factor for mortality. $Am \mathcal{J} Epi-$ demiol 1982; 116:102-13.
- 16 Menkes HA, Beaty TH, Cohen BH, et al. Nitrogen washout and mortality. Am Rev Respir Dis 1985;132:115-9.
- and mortality. Am Rev Respir Dis 1985;132:115-9.
 17 Hole DJ, Watt GC, Davey-Smith G, et al. Impaired lung function and mortality risk in men and women: findings from the Renfrew and Paisley Prospective Population Study. BMJ 1988;313:711-5.
 18 Sorlie P, Kannel W, O'Connor GT. Mortality associated with respiratory function and symptoms: the Framingham Study. Am Rev Respir Dis 1989;140:379-84.
 19 Tager IB, Rosner B, Tishler PV, et al. Household approximation of the production of
- aggregation of pulmonary function and chronic bronchitis. Am Rev Respir Dis 1976;114:485–92.
- O Schilling RSF, Letai AD, Hui SL, et al. Lung function, res-piratory disease, and smoking in families. Am J Epidemiol 1977:106:274-83.
- 21 Kauffmann F, Tager IB, Muñoz A, et al. Familial factors related to lung function in children aged 6–10 years. Am J Epidemiol 1989;**129**:1289–99.
- Coultas DB, Hanis CL, Howard CA, et al. Heritability of 22 ventilatory function in smoking and non-smoking New Mexico Hispanics. Am Rev Respir Dis 1991;144:770-5.

- 23 Chen Y, Horne SL, Rennie DC, *et al.* Segregation analysis of two lung function indices in a random sample of young families: the Humboldt Family Study. Genet Epidemiol 1996;13:35-47.
- 1990;13:30-47.
 Chen Y, Rennie DC, Lockinger LA, et al. Major genetic effect on forced vital capacity: the Humboldt Family Study. *Genet Epidemiol* 1997;14:63-76.
- 25 Chen Y, Dosman JA, Rennie DC, et al. Major genetic effects on airway-parenchymal dysanapsis of the lung: the Humboldt Family Study. Genet Epidemiol 1999;16:95-110.
- 26 Givelber RJ, Couropmitree NN, Gottlieb DJ, et al. Segrega-tion analysis of pulmonary function among families in the Framingham Study. Am J Respir Crit Care Med 1998;157: 1445-51
- 27 Higgins M, Keller J. Familial occurrence of chronic respiratory disease and familial resemblance in ventilatory capacity. 7 Chronic Dis 1975:28:239-51.
- Lebowitz MD, Knudson RJ, Burrows B. Family aggregation of pulmonary function measurements. Am Rev Respir Dis 1984;129:8-11.
- 29 Devor EJ, Crawford MH. Family resemblance for normal
- Devor EJ, Crawiord Mrl. Falling resembatic for normal pulmonary function. Ann Hum Biol 1984;11:439–48.
 Cotch MF, Beaty TH, Muñoz A, et al. Estimating familial aggregation while adjusting for covariates. Application to pulmonary function data from black and white sibships. Ann Epidemiol 1992;2:317–24. 31 Khoury MJ, Beaty TH, Cohen BH, eds. Fundamentals of
- genetic epidemiology. New York: Oxford University Press, 1993.
- 32 Hubert HB, Fabsitz RR, Feinleib M, et al. Genetic and
- Hubert HB, Fabsitz KK, Feinleib M, et al. Genetic and environmental influences on pulmonary function in adult twins. Am Rev Respir Dis 1982;125:409–15.
 Redline S, Tishler PV, Lewitter FI, et al. Assessment of genetic and nongenetic influences on pulmonary function. Am Rev Respir Dis 1987;135:217–22. 33
- 34 Man SFP, Zamel N. Genetic influence on normal variality of maximum expiratory flow-volume curves. J Appl Physiol 1976:41:874-7
- 35 Kawakami Y, Shida A, Yamamoto H, et al. Pattern of genetic influence on pulmonary function. *Chest* 1985;4:507–11. 36 Ghio AJ, Crapo RO, Elliott CG, et al. Heritability estimates
- Grano AJ, Ciapo RO, Finott GP, et al. Heritability estimates of pulmonary function. *Chest*, 1980;96:743-6.
 Webster PM, Lorimer EG, Man SFP, et al. Pulmonary function in identical twins: Comparison of nonsmokers and smokers. *Am Rev Respir Dis*, 1979;119-223-8.
 Hankins D, Drage C, Zamel N, et al. Pulmonary function in identical twins raised apart. *Am Rev Respir Dis* 1982;125: 119-21.
- 119 21
- 39 Astemborski JA, Beaty TH, Cohen BH. Variance components analysis of forced expiration in families. Am J Med Genet 1985;21:741-53.
- Genet 1953;21:741–35.
 Beaty TH, Liang KY, Seerey S, et al. Robust inference for variance components models in families ascertained through probands. II. Analysis of spirometric measures. *Genet Epidemiol* 1987;4:211–21.
 Lewitter FI, Tager IB, McGue M, et al. Genetic and
- environmental determinants of level of pulmonary function. *Am J Epidemiol* 1984;120:518–30.
 42 Cotch MF, Beaty TH, Cohen BH. Path analysis of familial
- resemblance of pulmonary function and cigarette smoking. Am Rev Respir Dis 1990;142:1337–43. Christian JC, Kang KW, Norton JJ Jr. Choice of an estimate
- 43 of genetic variance from twin data. Am J Hum Genet 1974; 26:154–61
- 24 Hannah MC, Hopper JL, Mathews JD. Twin concordance for a binary trait. II. Nested analysis if ever-smoking and ex-smoking traits and unnested analysis of "committed-committed analysis of "committed-committed analysis".
- smoking" trait. Am J Hum Genet 1985;37:153-65. 45 Redline S, Tishler PV, Rosner B, et al. Genotypic and phenotypic similarities in pulmonary function among family members of adult monozygotic and dizygotic twins. Am J Epidemiol 1989;**129**:827-36.
- 46 Beaty TH. Evolving methods in genetic epidemiology. I. Analysis of genetic and environmental factors in family studies. *Epidemiol Rev* 1997;1:14-23.

- 47 Cannings C, Tompson EA, Sholnick MH. Probability functions on complex pedigrees. Adv Appl Prob 1978;10:26–61. Rybicki BA, Beaty TH, Cohen BH. Major genetic
- mechanisms in pulmonary function. J Clin Epidemiol 1990; 43:667-75
- Demenais FM, Bonney GE. Equivalence of the mixed and regressive models for genetic analysis. I. Continuous traits. Genet Epidemiol 1989;6:597-617.
- Holberg CJ, Morgan WJ, Wright AL, *et al.* Differences in familial segregation of FEV₁ between asthmatic and non-asthmatic families: role of a maternal component. *Am* \tilde{J} Respir Crit Care Med 1998;158:162-9
- Silverman EK, Province MA, Campbell EJ, et al. Variability of pulmonary function in alpha-1-antitrypsin deficiency: 51 residual family resemblance beyond the effect of the Pi locus. Hum Hered 1990;40:340–55.
- 52 Mead J. Dysanapsis in normal lungs assessed by the relationship between maximal flow, static recoil, and vital capacity. Am Rev Respir Dis 1980;121:339-42.
- Tager IB, Weiss ST, Muñoz A, et al. Determinants of response to eucapneic hyerventilation with cold air in a population-based study. Am Rev Respir Dis 1986;134:502-
- 54 Castile RG, Hyatt RE, Rodarte JR. Determinants of maximal expiratory flow and density dependence in normal humans. *J Appl Physiol* 1980;**49**:897–904.
- Collins DV, Cutillo AG, Armstrong JD, et al. Large airway size, lung size, and maximal expiratory flow in healthy nonsmokers. Am Rev Respir Dis 1986;134:951-5.
- 56 Dolyniuk MV, Fahey PJ. Relationship of tracheal size to maximal expiratory airflow and density dependence. J Appl Physiol 1986;60:501-5.
- 57 Hoffstein V. Relationship between lung volume, maximal expiratory flow, forced expiratory volume in one second, and tracheal area in normal men and women. Am Rev Respir Dis 1986;134:956-61.
- 58 Knudson RJ, Schroter RC, Knudson DE, et al. Influence of airway geometry on expiratory flow limitation and density dependence. *Respir Physiol* 1983;52:113-23.
- Martin TR, Castile RG, Fredberg JJ, et al. Airway size is 59 related to sex but not lung size in normal adults. J Appl Physiol 1987;63:2042-7.
- Martin TR, Feldman HA, Fredberg JJ, et al. Relationship between maximal expiratory flows and lung volumes in 60 growing humans. J Appl Physiol 1988;65:822-8. Pagtakhan RD, Bjelland JC, Landau LI, et al.
- Sex differences in growth patterns of the airways and lung parenchyma in children. J Appl Physiol 1984;56:1204–10. Chen Y, Horne SL, Dosman JA. Body weight and body gain
- related to pulmonary function decline in adults: a six year follow up study. *Thorax* 1993;48:375–80.
- Chen Y, Horne SL, Dosman JA. Increased susceptibility to lung dysfunction in female smokers. Am Rev Respir Dis 1991;143:1224-30.
- Gold DR, Wang X, Wypij D, et al. Effects of cigarette smoking on lung function in adolescent boys and girls. N Engl J Med 1996;335:931-7.
- Akaike H. A new look at the statistical model identification. IEEE Trans Automat Conrol 1974;AC-19:719-23.
- MacLean CJ, Morton NE, Lew R. Analysis of family resem-66 blance. IV. Operational characteristics of segregation analy-sis. Am J Hum Genet 1975;27:365-84.
- Borecki IB, Province MA, Rao DC. Power of segregation analysis for detection of major gene effects on quantitative traits. *Genet Epidemiol* 1994;**11**:409–18.
- Khoury MJ, Beaty TH, Newill CA, et al. Genetic-68 environmental interactions in chronic airways obstruction. Int J Epidemiol 1986;15:65-72.
- Tiret L, Abel L, Takotovao R. Effect of ignoring genotype-environment interaction on segregation analysis of quantitative traits. *Genet Epidemiol* 1993;**10**:581–6. 69