Markers of early disease and prognosis in COPD

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Abstract: COPD is a complex disease with multiple pathological components, which we unfortunately tend to ignore when spirometry is used as the only method to evaluate the disorder. Additional measures are needed to allow a more complete and clinically relevant assessment of COPD. The earliest potential risk factors of disease in COPD are variations in the genetic background. Genetic variations are present from conception and can determine lifelong changes in enzyme activities and protein concentrations. In contrast, measurements in blood, sputum, exhaled breath, broncho-alveolar lavage, and lung biopsies may vary substantially over time. This review explores potential markers of early disease and prognosis in COPD by examining genetic markers in the $\alpha_1$-antitrypsin, cystic fibrosis transmembrane conductance regulator (CFTR), and MBL-2 genes, and by examining the biochemical markers fibrinogen and C-reactive protein (CRP), which correlate with degree of pulmonary inflammation during stable conditions of COPD. Chronic lung inflammation appears to contribute to the pathogenesis of COPD, and markers of this process have promising predictive value in COPD. To implement markers for COPD in clinical practice, besides those already established for the $\alpha_1$-antitrypsin gene, further research and validation studies are needed.

Keywords: chronic obstructive pulmonary disease, biomarker, pathogenesis, prognosis, genetics

Introduction

Chronic obstructive pulmonary disease is defined as a disease state characterized by airflow limitation that is not fully reversible; the airflow limitation is usually both progressive and associated with an abnormal inflammatory response of the lungs to noxious particles or gases. According to the WHO Global Burden of Disease Study, COPD is the fifth leading cause of death worldwide. Projections from the WHO predict that COPD will continue to increase in the years to come, challenging the health services in countries where cigarette smoking is prevalent. Consequently, we need to improve the diagnostic and therapeutical tools against COPD.

COPD is a complex disease with multiple pathological components, which we unfortunately tend to ignore when spirometry is used as the only method to evaluate the disorder. Additional measures are needed to allow a more complete and clinically relevant assessment of COPD. This may enable better phenotyping of different types of the disease and help improve the evaluation of disease activity and efficacy of therapy. The earliest potential risk factors of disease in COPD are variations in the genetic background. Genetic variations are present from conception and can determine lifelong changes in enzyme activities and protein concentrations. In contrast, measurements in blood, sputum, exhaled breath, broncho-alveolar lavage, and lung biopsies may vary substantially over time. The best known inherited risk factor for early-onset COPD is genetically deficient plasma levels of $\alpha_1$-antitrypsin.

In this review we have explored potential markers of early disease and prognosis in COPD using data from the prospective epidemiological study, the Copenhagen City Heart Study, and information from two Danish National Registers covering all hospital discharges and causes of death in the country. The investigations are focused
on genetic markers in the α1-antitrypsin and cystic fibrosis transmembrane conductance regulator (CFTR) genes, the two most important known genes for obstructive lung disease, and the MBL-2 gene in which functional polymorphisms have been associated with features of COPD in previous reports.\textsuperscript{13,14} Since COPD is defined by an abnormal inflammatory response to noxious particles or gases, elevated lung inflammation could also represent an early pathological event in COPD. Fibrinogen and C-reactive protein (CRP) correlate with degree of pulmonary inflammation during stable conditions of COPD,\textsuperscript{15,16} and could potentially be useful as markers of early disease and prognosis in COPD.

**α1-antitrypsin deficiency and COPD**

The Z allele substitutes lysine for glutamic acid at position 342 in the α1-antitrypsin gene, while the S allele substitutes valine for glutamic acid at position 264. The Z allele and to a lesser degree the S allele can cause α1-antitrypsin to polymerise in hepatocytes leading to reduced plasma levels of α1-antitrypsin and higher risk for COPD.\textsuperscript{17} Severe α1-antitrypsin ZZ deficiency is the most important known genetic risk factor for COPD. Individuals with less severe α1-antitrypsin deficiency (MS, MZ, SZ genotypes) could also have an increased susceptibility for COPD. We compared plasma α1-antitrypsin level, decline in lung function, and risk of hospitalization for COPD in α1-antitrypsin MS, MZ, SZ genotypes versus controls.\textsuperscript{18,19} We found a stepwise reduction in plasma α1-antitrypsin with increasing severity of the α1-antitrypsin genotype (Figure 1). The MZ, SZ and ZZ genotypes were associated with reduced plasma α1-antitrypsin, greater annual FEV\textsubscript{1} decline (ZZ only borderline), and with greater risks of spirometry-defined airway obstruction and COPD (MZ and SZ only borderline for airway obstruction) (Figure 1). Although MS was associated with lower plasma α1-antitrypsin level, this genotype did not confer greater FEV\textsubscript{1} decline or increased risk of airway obstruction and COPD.

To put these results into an international context, we identified previous studies on COPD risk in intermediate α1-antitrypsin deficiency, aggregated the results using meta-analyses, and calculated summary risk estimates for COPD in MS, MZ, and SZ individuals\textsuperscript{20,21} (Figure 2). Seventeen studies were included in meta-analyses on MS, 16 in meta-analyses on MZ, and 6 in meta-analyses on SZ. The summary odds ratio for MS was increased at 1.2 (1.0–1.4), the summary odds ratio for MZ was increased at 2.3 (1.6–3.4), and the summary odds ratio for SZ was increased at 3.3 (1.2–8.6). Previous publications indicate that the magnitude of the COPD risk in MZ differs by study design. Thus, we performed a meta-analysis for MZ stratified by study design. After stratification, the summary odds ratio for MZ was 1.5 (0.97–2.3) in cross-sectional studies and 3.0 (2.1–4.3) in case-control studies. The summary odds ratios for MS and SZ, and the summary odds ratio for MZ in cross-sectional studies, did not differ considerably from the results provided by the Copenhagen City Heart Study.\textsuperscript{18–21}
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**Figure 2** Cross-sectional and case-control studies of COPD risk in protease inhibitor MS, MZ, and SZ heterozygotes versus MM individuals. Box sizes are proportional to inverse-variance weights (random effects model). Lines represent 95% confidence intervals. Adapted with permission from Dahl M, Hersh CP, Ly NP, Berkey CS, Silverman EK, Nordestgaard BG. *Eur Respir J*. 2005;26:67–76. Copyright © 2005 European Respiratory Society Inc, and from Hersh CP, Dahl M, Ly NP, Berkey CS, Nordestgaard BG, Silverman EK. *Thorax*. 2004;59:843–849. Copyright © BMJ Publishing Group.

**Abbreviation:** PI, protease inhibitor.

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**Study** | **Odds ratio (95% CI)** | **% Weight**
---|---|---
Fagerhol, 1969 | 1.10 (0.36 to 3.36) | 6.3
Talamo, 1972 | 1.49 (0.32 to 6.86) | 4.3
Kueppers, 1974 | 16.78 (2.16 to 130.58) | 2.7
Barnett, 1975 | 4.66 (0.99 to 21.89) | 4.1
Cox, 1975 | 2.60 (1.16 to 5.80) | 8.3
Cox, 1976 | 2.73 (1.00 to 7.27) | 7.3
Kueppers, 1977 | 1.52 (0.52 to 4.42) | 6.6
Matzen, 1977 | 2.96 (0.62 to 14.20) | 4.1
Chan-Young, 1978 | 0.15 (0.01 to 2.41) | 1.6
Abboud, 1979 | 6.86 (0.36 to 129.15) | 1.4
Gulsvik, 1959 | 1.20 (0.53 to 2.73) | 8.6
Bartmann, 1985 | 5.27 (2.40 to 11.57) | 8.9
Lieberman, 1986 | 3.42 (2.21 to 5.28) | 12.3
Poller, 1990 | 2.17 (0.66 to 5.44) | 7.3
Sandford, 1999 | 10.17 (0.59 to 174.22) | 1.5
Dahl, 1992 | 1.34 (1.65 to 1.71) | 13.9
Overall, (95% CI) | 2.31 (1.60 to 3.35) |

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**Study** | **Odds ratio (95% CI)** | **% Weight**
---|---|---
Fagerhol, 1969 | 1.99 (0.12–31.98) | 10.1
Abboud, 1979 | 1.87 (0.07–47.63) | 7.8
Gulsvik, 1959 | 3.78 (0.34–42.04) | 12.8
Bartmann, 1985 | 24.46 (3.25–183.94) | 16.8
Lieberman, 1986 | 0.82 (0.16–4.22) | 22.3
Dahl, 1992 | 3.78 (1.06–13.41) | 30.1
Overall | 3.26 (1.24–8.57) |
The meta-analyses suggest that risk of COPD is marginally elevated in MS and MZ individuals (in cross-sectional studies), whereas the more severe SZ genotype is an important risk factor for COPD. It is possible that MS, MZ, and SZ have greater risk estimates in subgroups of patients or in individuals with other additional risk factors for COPD, but further studies of very large populations are needed to determine if this is the case. We now offer an analysis in COPD patients that detects the S and Z alleles of the \( \alpha_1 \)-antitrypsin gene with a simultaneous measurement of plasma \( \alpha_1 \)-antitrypsin.\(^{22}\) This will help clinicians to detect \( \alpha_1 \)-antitrypsin-deficient individuals with high risk of COPD, and opens up for early smoking prevention counselling and potential therapy to individuals identified with \( \alpha_1 \)-antitrypsin deficiency through this new analysis.

### Cystic fibrosis F508del and COPD

Cystic fibrosis is characterized by progressive obstructive lung disease in homozygous individuals, and it is possible that heterozygotes for cystic fibrosis are at increased risk of less severe forms of obstructive lung disease. To examine this, we screened 9141 individuals for the common F508del allele, and we compared lung function and prevalences of COPD and asthma in F508del heterozygous individuals versus noncarriers.\(^{23,24}\)

Two hundred and fifty individuals were heterozygous and none homozygous for the cystic fibrosis F508del allele.\(^{23,25}\) Among heterozygotes, none had previously been admitted to a hospital due to cystic fibrosis or carried an additional 394delTT or N1303K allele in the CFTR gene.\(^{24}\) Second to the F508del allele, the 394delTT and N1303K alleles are the two most common cystic fibrosis alleles in Denmark with allele frequencies of about 1% or more in the Danish population.\(^{26}\)

Based on triplicate measurements of pulmonary function, we found that heterozygous individuals had lower FEV\(_1\) and FVC than noncarriers (Figure 3).\(^{24}\) Consistent with this, heterozygous individuals also reported that they had asthma more often than noncarriers, whereas the prevalence of COPD did not differ between the two groups (Figure 4).\(^{23,24}\)

These findings were independent of influence from the common but less severe polythymidine tract variants in intron-8 of the CFTR gene.\(^{27}\) The shorter this polythymidine tract is, the more often exon-9 is skipped from CFTR mRNAs leading to a CFTR protein without chloride activity.\(^{28}\) Studies of asthma risk in cystic fibrosis heterozygotes, however, have not convincingly shown increased asthma among heterozygotes.\(^{23,24,27,29–39}\) Combining all the available data from the literature in a meta-analysis,\(^ {23,29,31,34–37,39}\) the summary odds ratio for asthma in cystic fibrosis/F508del heterozygotes is 1.3 (1.1–1.6; \( p = 0.01 \) ) (Figure 5).

In conclusion, cystic fibrosis F508del heterozygotes are not at greater risk of COPD, but may be overrepresented among people with asthma and have poorer lung function than non-carriers. Combining recent reports from the literature, the risk for asthma in cystic fibrosis heterozygotes is roughly 30% elevated. Thus, F508del cannot be used as a biomarker for COPD or asthma in clinical settings. It is possible that F508del can be used as a clinical marker for asthma in certain subgroups of patients in the future, when other susceptibility markers for asthma are available.

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**Figure 3** Levels of FEV\(_1\) and FVC by cystic fibrosis F508del carrier status. Values are means ±SEM, based on 10-year age groups. Number of measurements: F508del, \( n = 270 \); and noncarriers, \( n = 10,002 \). \( P \)-values are by general linear repeated-measures analysis comparing F508del heterozygotes versus noncarriers. Adapted with permission from Dahl M, Nordestgaard BG, Lange P, Tyljaerg-Hansen A. J Allergy Clin Immunol. 2001;107:818–823.\(^ {24}\) Copyright © Elsevier.
Mannose-binding lectin (MBL) is an acute phase protein secreted from the liver.\textsuperscript{40,41} It has antibody-like function in the innate immune system, where it binds sugar moieties on microorganisms and on apoptotic cells. Bound MBL can clear its targets through promoting opsonophagocytosis or by activating the complement system through MBL-associated protease 1 and 2. Deficiency of MBL levels/activity in plasma due to polymorphisms in the MBL-2 gene is thought to weaken normal innate immune functions against microorganisms and apoptotic cells in the body.\textsuperscript{42} Genetically reduced MBL in plasma raises the susceptibility for disease in COPD and asthma.\textsuperscript{13,14,43} Thus, we determined the risk of COPD and asthma in MBL deficient individuals versus controls in our study of the population at large.\textsuperscript{44} We did not find an increased risk of COPD or asthma with MBL deficiency (Figure 6). The risk of death due to respiratory disease was unaffected by MBL deficiency status.

Only few other studies have examined the potential relation between MBL deficiency and obstructive lung disease.\textsuperscript{13,14,43} Our study had ample statistical power to test the hypothesis that MBL deficiency is associated with risk of obstructive

\textbf{Mannose-binding lectin and COPD}

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\textbf{Study name} & \textbf{Odds ratio} & \textbf{Lower limit} & \textbf{Upper limit} & \textbf{p-value} & \textbf{Asthma/Total} & \textbf{Relative weight} \\
\hline
Dahl & 1.60 & 1.03 & 2.48 & 0.03 & 23/250 529/8891 & 23.65 \\
Lowenfels & 1.30 & 0.92 & 1.83 & 0.13 & 107/1113 52/688 & 37.89 \\
Munthe e-kaas & 1.22 & 0.60 & 2.48 & 0.58 & 13/34 223/663 & 8.99 \\
Castellani & 1.16 & 0.46 & 2.90 & 0.74 & 12/261 8/201 & 5.42 \\
de cid & 0.77 & 0.42 & 1.43 & 0.41 & 25/47 222/374 & 12.22 \\
Lazar & 1.44 & 0.85 & 243.83 & 0.06 & 21/21 12/164 & 0.57 \\
Mennie & 0.99 & 0.46 & 2.12 & 0.99 & 11/186 20/337 & 7.88 \\
Tzets & 4.50 & 1.41 & 14.34 & 0.01 & 9/17 11/55 & 3.37 \\
Overall & 1.31 & 1.06 & 1.62 & 0.01 &  &  \\
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\textbf{Figure 5} Cross-sectional and case-control studies of asthma risk in cystic fibrosis/F508del heterozygotes. Box sizes are proportional to inverse-variance weights (random effects model). Lines represent 95% confidence intervals.
lung disease. In conclusion, genetic MBL deficiency is not a major risk factor for COPD in the Danish general population. At present, genetic variants in the MBL-2 gene cannot be used as biomarkers of obstructive lung disease.

In children, MBL could have an important function in pulmonary host defence during the vulnerable period when the adaptive immune system is immature. We tested whether adult MBL deficient individuals more often than controls were hospitalized or died due to pulmonary infection, but found no increased frequency of pulmonary infection with MBL deficiency. These results suggest that MBL does not have a major role in pulmonary defence against microorganisms in adult Danish individuals.

**Other genetic markers of COPD**

We have used a candidate gene approach to evaluate genetic markers in the α1-antitrypsin, CFTR, and MBL genes, and their relation to COPD. Other approaches include linkage studies, transcript-expression analyses, and animal model genetics, all of which can be used to identify certain genetic candidates for COPD. To translate results from such studies into clinical medicine, an estimation of risk of disease is needed. This can be achieved through association studies of the population at large such as ours.

In COPD recent focus has been on the enzyme serine protease inhibitor E2 (SERPINE2), as this enzyme is a family relative of α1-antitrypsin (SERPINA1), the most important risk factor of inherited COPD. Supporting that SERPINE2 is of importance to COPD susceptibility, polymorphisms in the SERPINE2 gene have been shown to associate with features of COPD. The enzymes epoxide hydrolase 1 (EPHX1) and superoxide dismutase 3 (SOD3) are also of potential interest as markers in COPD, as these enzymes may inhibit increased oxidative stress in the lungs due to cigarette smoke inhalations. Comprehensive reviews on other candidate genes in COPD are readily available.

### Plasma fibrinogen and COPD

Pulmonary inflammation in COPD is associated with increased levels of acute phase reactants in plasma, and these reactants could potentially be used to predict risk of future COPD events. To test whether the acute phase reactant fibrinogen predicts future COPD, we determined lung function and relative risk of COPD hospitalization in individuals with increased levels of plasma fibrinogen. We found that individuals with baseline plasma fibrinogen of more than 3.3 g/L versus less than 2.7 g/L had reduced lung function, increased cumulative incidence of COPD hospitalization (Figure 7), and an increased relative risk of 1.7 (1.1–2.6) for COPD hospitalization during 6 years of follow-up. These findings suggest that plasma fibrinogen is a significant predictor of future COPD in the population at large.

A later study confirms plasma fibrinogen as a marker of COPD prognosis: Gan et al estimated a 0.37 g/L difference in plasma fibrinogen between COPD patients and controls. The clinical utility of plasma fibrinogen as a biomarker of COPD seems limited based on its low predictive value for COPD. IL-6 regulates fibrinogen production from the liver. Thus, the results point to an important function for fibrinogen, IL-6, or factors upstream of IL-6 in the pathogenesis of COPD. Further studies are required to elucidate which part, if any, fibrinogen or IL-6 has in the progression of COPD.

### Plasma C-reactive protein and COPD

COPD is associated with elevated C-reactive protein (CRP) concentration in serum which equals concentrations in plasma, and it is possible that CRP as a marker of pulmonary inflammation could be used to predict risk of future COPD events. To test whether plasma CRP predicts future COPD events, we measured CRP in a subgroup of individuals with high risk of clinical COPD, that is, individuals with spirometry-defined airway obstruction. During 8 years of follow-up, we recorded admissions and deaths due to COPD, and relative risks for COPD hospitalization or deaths were determined. We found that individuals with serum CRP of more than 3 mg/L versus ≤3 mg/L at baseline had increased cumulative incidence of COPD hospitalization and death (Figure 8), an increased hazard ratio of 1.4 (1.0–2.0) for...
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COPD hospitalization, and an increased hazard ratio of 2.2 (1.2–3.9) for COPD death. Absolute risks for COPD hospitalization and death increased with baseline serum CRP levels (Figure 9). The highest absolute 10-year risks for COPD hospitalization and death were 54% and 57% among individuals with baseline CRP above 3 mg/L, above 70 years of age, with a tobacco consumption of more than 15 g/day, and an FEV1% predicted less than 50%.

The data suggests an important role for CRP, or its main regulatory cytokine IL-6, in COPD progression and development. Supporting a role for IL-6 in COPD pathogenesis, IL-6 may stimulate inflammatory cell recruitment to the lung, and lead to emphysema-like airspace enlargement and respiratory muscle wasting in animal models of lung disease. Thus, our own and previous data seem to suggest that IL-6 or factors upstream of IL-6 are involved in the pathogenesis of COPD. In the future, CRP levels could be used clinically to assess prognosis in patients with airway obstruction. High absolute risk as depicted in Figure 9 may motivate patients at the highest risk to quit smoking or to receive medication.

Other biochemical markers of COPD

In COPD, biochemical biomarkers represent a new field that we have just begun to explore. The potential is great. COPD is a multicomponent disease for which airway inflammation plays an imperative role. This component, among several others, needs descriptive measures to provide a more comprehensive and clinically accurate assessment of COPD.

Biochemical COPD markers can be measured in exhaled breath, induced sputum, bronchoalveolar lavage fluid, lung biopsies, and plasma. Compared to biomarkers measured in other sample material, biochemical markers determined in plasma are reliably measured using equipment that is cost-effective and readily available in clinical settings. However, biochemical markers, and in particular those measured in plasma, may be modulated by morbidities other than COPD, and thus may in some circumstances represent epiphenomena unrelated to the COPD phenotype.

The validation of a biomarker has been proposed to involve the following three phases: 1) demonstration that the marker frequency/magnitude is associated with a clinical outcome. 2) Phase I/II trials demonstrating effects on the marker with therapeutic intervention. Are there dose-dependent effects of treatment on the marker? 3) Demonstration that treatment-related changes in the marker are associated with positive changes in clinical outcomes. Is the marker applicable to all disease stages and all interventions?

In COPD the biochemical markers are many, but these have been less well investigated than biochemical markers for plasma fibrinogen. P = 0.003 for >3.3 g/L versus <2.7 g/L, P = 0.31 for 2.7–3.3 g/L versus <2.7 g/L on log-rank test. Adapted with permission from Dahl M, Tybjærg-Hansen A, Vestbo J, Lange P, Nordestgaard BG. Am J Respir Crit Care Med. 2001;164:1008–1011. Copyright © 2001 American Thoracic Society.
in asthma. When evaluating therapies for COPD, markers that could surrogate for critical pathological processes in COPD would be highly warranted. For example desmosine as part of mature elastin indicates elevated turnover of elastic fibres in COPD.\textsuperscript{66} Novel lung-specific markers also appear promising,\textsuperscript{67,68} in particular surfactant protein-A, which is elevated in lung tissue and sputum in COPD patients when compared with controls in a proteomics study.\textsuperscript{69} The exploration and validation of potential markers for COPD is currently ongoing and no specific marker has so far been implemented for clinical use.\textsuperscript{7–9,70–72}

Conclusions

Using data from up to 9245 adult participants from the Copenhagen City Heart Study and information from the national Danish Hospital Discharge Registry and the national Danish Causes of Death Registry, we explored new potential markers of COPD.

The risk of COPD is marginally elevated for the genetic biomarkers, MS and MZ, in the $\alpha_1$-antitrypsin gene, whereas the SZ biomarker is an important risk factor of COPD. The ZZ genotype is already in clinical use as a genetic marker for early-onset COPD. We have set-up an analysis for diagnosing the SZ and ZZ genotypes together with measurement of plasma $\alpha_1$-antitrypsin levels to be used broadly, including in COPD patients. This will help clinicians to detect $\alpha_1$-antitrypsin-deficient individuals with a high risk for COPD, and opens up for early smoking prevention counseling and potential therapy for individuals identified with $\alpha_1$-antitrypsin deficiency through this new analysis.

The genetic biomarker, F508del, in the cystic fibrosis gene was not associated with COPD, but with increased asthma risk. Aggregating results from previous reports, the risk of asthma is only marginally elevated. Thus, F508del cannot at present be used as a marker of COPD or asthma in clinical settings.

Genetic deficiency of MBL is not a major risk factor for COPD or asthma, and it appears that genetic variants in the MBL-2 gene cannot at present be used as biomarkers for obstructive lung disease in the Danish population.

Elevated plasma fibrinogen and CRP levels were associated with increased COPD risk and may be good for prognostication once a diagnosis of COPD has been established. In the future after further validation, it is possible that these markers, particularly CRP, can be used to categorise individuals with low, medium, or high risk of COPD, and thus point out patients in need of intensified prevention and treatment for COPD. Furthermore, the observed association between COPD and elevated fibrinogen and CRP levels suggests that either factor or upstream factors regulating fibrinogen and CRP expression are important players in the pathogenesis.

![Figure 8](https://www.dovepress.com/)

Figure 8. Cumulative incidence of COPD events according to baseline serum CRP levels. Cumulative incidences of COPD hospitalization and death were increased in individuals with baseline CRP $>3$ mg/L versus $\leq3$ mg/L. Adapted with permission from Dahl M, Vestbo J, Lange P, Bojesen SE, Tybjaerg-Hansen A, Nordestgaard BG. Am J Respir Crit Care Med. 2007;175:250–255. Copyright © 2007 American Thoracic Society.
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of COPD. To determine whether fibrinogen or CRP are causally related to COPD, we are currently investigating if genetically increased levels of fibrinogen and CRP are associated with increased risk of future COPD. We anticipate that fibrinogen and CRP-regulating factors (IL-6, TNF-α, IL-1) rather than fibrinogen and CRP themselves are involved in COPD pathogenesis, as IL-6, TNF-α, and IL-1 are already factors with established functions in COPD. IL-6 is involved in inflammation, vascular permeability, and cell proliferation, while TNF-α, and IL-1 initiate and maintains inflammation, and activates endothelial and epithelial cells.5

The best known early disease markers for COPD remain genetic variations in alpha-1-antitrypsin. They cause lifelong reduced levels of alpha-1-antitrypsin in plasma and increase the risk for COPD. As such, future studies on markers in COPD should ideally demonstrate association between a marker and COPD, as well as change in the COPD risk with changes in the marker concentration or function (genetically or medically induced). Randomized pharmacological trials or mendelian randomization designs are useful in this regard and will enable us to assess the importance of early markers for COPD much better. The study of disease markers in COPD can improve the phenotyping of different types of COPD. Linking the markers to critical pathologic events in COPD can provide novel insights into the pathogenesis of COPD. With the recent focus on heterogeneity in COPD and the mounting efforts in identifying novel markers for COPD this research field seems in progress.

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Disclosures

Neither author has conflicts of interest to disclose.
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