Biomarkers in chronic obstructive pulmonary disease: confusing or useful?

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Abstract: The field of biomarker research has almost reached unmanageable proportions in chronic obstructive pulmonary disease (COPD). The developments of new technology platforms have generated a huge information data base, both cross sectionally and increasingly, longitudinally. The knowledge emerging provides an enormous potential for understanding the disease pathophysiology, for developing markers specific for long-term outcomes, and for developing new therapeutic strategies. However, the excitement must be tempered with an understanding of the limitations of the data collection techniques, and of the variations in disease state, activity, impact, and progression. Nevertheless, the most crucial aspect in interpreting the current literature is the recognition of the relatively superficial characterization of what is a complex group of pathological processes with a common end point of airflow limitation. The current review explores some of these issues together with those areas where real progress appears to have been made, and provides caution on interpretation.

Keyword: emphysema, inflammation, secretions, technology platforms

Introduction

“Biomarkers” have become a hot topic in the study and treatment of chronic obstructive pulmonary disease (COPD). In simple terms, a biomarker is a measurable characteristic that reflects the presence, severity, or state of a disease. For instance, whilst spirometry reflects the presence and dyspnea the severity or impact of COPD, temperature may reflect an infectious state. Furthermore, a change in the expression of a biomarker may reflect progression, the risk of progression, or the response to treatment, whilst genes and their products reflect the likelihood of disease emergence. Thus biomarkers can be specific cells, molecules, genes, gene products, organ functions, or general clinical characteristics that reflect the damage done, or the process that the damage sets in motion or the process that leads to the damage, or can indicate the prognosis and/or response to therapy. More recently, the use of general technology platforms has led to an exponential increase in information, accompanied by uncertainty as to whether this reflects cause or effect.

The current article attempts to review (at least in part) this enormous field together with some guidelines, based on potential pitfalls and on what we need to know. In general the “hot” questions we need answers to are:

- Can we identify the susceptible individual before the disease state develops?
- Can we identify “early” disease whilst it is still relatively asymptomatic?
- Can we predict the rate of progression of the disease?
• Can we identify or confirm the key pathophysiological processes, leading to new drug discovery?
• Can we identify factors that predict the response to current treatment or short Phase II clinical trials that provide confidence to undertake more extensive and definitive Phase III studies?
• Can we be sure of the impact of our new strategies on the patient, in the present or future?

Defining COPD status
COPD is a slowly progressing disease, although the rate of progression varies. This is an important concept as it impacts upon three features of COPD, namely, the disease activity, its severity, and the impact on the patient. The activity of the disease process will influence the current or future severity in a time dependent manner, ie, a highly active disease process will produce more severe disease at an earlier age. Similarly, the severity of the disease will have a variable impact on the patient, dependent upon the rate at which it developed. Small changes that develop rapidly are more likely to have a greater impact on the older patient than the same change developing slowly will have on a more elderly patient (see Figure 1).

In addition, it is increasingly recognized that COPD, although defined by major changes in spirometry, has several distinct phenotypes (clinically and pathologically) as well as additional features, such as the exacerbation complex (also consisting of many pathophysiological processes) and the recognized comorbidities, which may have some common physiological yet distinct pathological mechanisms. 1 “Omics” is a neologism that aims at the collective characterization and quantification of pools of biological molecules that translate into the structure, function, and dynamics of an organism. Trying to understand the complexities of the enormous data generated by the “omics” platforms in COPD (proteomics, genomics, metabolomics, transcriptomics etc) is almost impossible without a similar high-quality and detailed patient characterization, and all biomarker studies need to be interpreted in this light.

Clinical biomarkers
Chest hyperinflation, low body mass index (BMI), the use of accessory muscles of respiration, and prolonged expiration have always been the physical markers of the presence of airflow obstruction, especially that occurring with a predominant emphysema phenotype. However the objective measures of lung physiology have been the “gold standard” biomarkers to date. In particular, not only has spirometry been the lynchpin of diagnosis but forced expired volume in 1 second (FEV1) has been central to our description of severity. It is a marker of disease progression and response to therapy, as well as a sound predictor of mortality 2 and a weak indicator of future exacerbations. 3 Nevertheless, it only measures one aspect of COPD and relates poorly to the presence of emphysema 4 or to patient-reported quality of life, 5 and its deterioration is not linear with time. 6

However spirometry remains central to our thinking of COPD, requires little equipment, is relatively simple to perform and interpret (with appropriate training), and hence is a useful tool to detect “early” disease (at least when sufficiently advanced to fulfill the cross-sectional diagnostic criterion).

Figure 1 Line graph representation of disease progression in 3 idealized patients. Patient (Pt) A has slow disease progression and with age may notice or report few symptoms (ie, low impact/low disease activity/mild disease). Patient B has greater progression and symptoms become noticeable in the 60s (moderate impact/moderate disease activity/moderate disease). Patient C has rapid disease progression leading to symptoms in the 30s (major impact/high disease activity/mild disease).
Yet, even the term “early disease” is interpreted in different ways. To a patient, this may be understood as the stage in the disease when the symptoms or their impact is mild, whereas to physiologists, it is understood to occur when the spirometry is abnormal and yet is relatively close to the normal range. This concept is summarized in Figure 1, showing that the speed with which airflow obstruction develops will influence the symptoms perceived by the patient, which will clearly be dependent upon the activity of their lifestyle and expectations. Furthermore, the presence of comorbidity and other physiological impairment (gas trapping and defects of gas transfer) will also influence the overall impact on the patient. Indeed, the impact, as determined by health status tools, is also a fair guide to long-term mortality. However, the poor correlation of health status with physiology suggests such tools only partly measure the same thing. Mortality is related to many factors, including the degree of breathlessness, exercise limitation, and low BMI. These markers predict the same end point but do not necessarily reflect the same thing. For this reason, composite scores, such as the BODE index (Body mass index, airflow Obstruction, Dyspnoea and Exercise), may be better predictors of long-term mortality. Because these relatively insensitive tests have been studied for many years, their use in assessing patients’ prognosis, progression, and response to treatment has been well documented. However, although useful as cross-sectional measures of cohort risks, these composite markers are less robust in assessing individual patients, and a progression of health status can occur independent of spirometry, even though progression of both also predict mortality. However, the composite markers do fulfill two other criteria, namely, that changes, both by decline or improvement, also relate to mortality, although may not be predictive of progression per se.

**Patient-reported outcomes**

Symptoms are included in many patient-reported outcomes and can be scored, in their own right, to provide both an indicator of the disease impact in individual patients but also, of progression and, importantly, the response to treatment. This alone consists of a large literature of assessment and validation. A good start for the interested reader is the validation of the COPD Assessment Test and its comparison with other patient-reported outcomes (which again highlights the weak relationship to spirometry). Nevertheless, these subjective tests do act as a biomarker of the impact of the disease on the patient, even if this is not entirely (or even only partly) explained by the current objective measures (a disjunction between disease severity and impact).

**Radiology**

More recently, the widespread use of high-resolution computed tomography (CT) scans and other radiological techniques has been applied to COPD populations. The CT scan is now the marker of choice to identify and quantify the presence and type of emphysema. This patient phenotype also relates to health status (though also weakly) as shown in Figure 2. Emphysema is a good predictor of mortality, can be present

![Figure 2](https://www.dovepress.com/)

**Figure 2** Relationship between health status recorded as the SGRQ score and decreasing lung density as a marker of emphysema. Individual patient data points are shown and the significance of the correlation is given.

**Abbreviation:** CT, computed tomography.
with normal spirometry, and is the outcome of choice in treatments aimed at improving or slowing down the progression of emphysema. CT scanning is recognized as the most sensitive test of progression in patients with emphysema, being significantly more sensitive than spirometry and even gas transfer, in such patients. More recently, further refinements of CT analysis have enabled the relationship between small airways disease and emphysema to be characterized in more detail, and the preliminary data suggest that small airways disease may precede the development of radiological emphysema. The implications of this for our understanding of pathophysiology clearly requires further study.

Radiology can also study the airways, indicating responses to bronchodilator therapy and the presence and distribution of accompanying bronchiectasis, which is also a predictor of mortality (although the reasons are currently unclear).

**Chronic bronchitis**

The chronic bronchitis phenotype received little attention as a marker until recently and is now recognized as a predictor of disease progression and recurrent exacerbations. The latter defines a further clinical phenotype (and hence marker) likely to respond to anti-inflammatory therapy, both by the inhaled route and oral administration.

Despite the well-tried and accepted roles of these “clinical” biomarkers, in more recent research, the term has come to mean a fluid sample test that is specific, particularly in terms of the pathophysiological processes taking place. Such research was initially based on a hypothesis-testing strategy, but the development and ready availability of technology platforms has caused this to become a data-collecting and possibly, a hypothesis-generating strategy. The latter relies on extensive data collection, which is then probed for patterns (generally the “omics” strategy) that may become informative about the potential process(es) taking place.

**Biomarkers based on pathophysiology**

The generally accepted concept is that the development of COPD occurs in susceptible individuals as a result of an enhanced inflammatory response to inhaled substances, like cigarette smoke. However, not only is COPD a generically defined syndrome but, inflammation itself is a highly complex, interrelated network of cells, cytokines, and other proteins that both amplify and dampen the process – in part, this represents a balance between the destructive elements and the reparative ones. However, once damage has occurred to the delicate structures of the lung, repair cannot completely restore function, especially in the alveolar regions. This has been best demonstrated in animal models of emphysema, where a single challenge has been shown to result in the loss of lung elastin and, although it reaccumulates, the development of emphysema. To date, the majority of biomarker studies in COPD have concentrated on the damage process.

Our understanding of the mechanisms involved in the pathophysiology of COPD relates back to the single observation that alpha-1-antitrypsin (AAT)-deficiency (AATD) resulted in an increased susceptibility to the development of early-onset basal emphysema. The cascade of events that followed this observation resulted in many studies of AAT phenotypes in COPD, but apart from the recognition of the Pi ZZ severe form of deficiency and some very rare null genes, little was found. However, even the deficiency itself is not a highly sensitive marker since many affected patients remain healthy and with normal lung function. The true incidence of the deficiency and disease and hence, the sensitivity of even this genetic/biochemical marker remains unknown.

The mechanism involved became relatively easy to define. AAT is a specific inhibitor of serine proteinases, and elastase released by activated neutrophils, have both been shown to produce emphysematous lesions in experimental animals. Since neutrophil recruitment to the lung results in the release of these enzymes, elastin digestion during transit should be excessive in the presence of low concentrations of AAT. Completion of the pathophysiological axis required a chemoattractant signal and cell activation, adhesion, and migration, as summarized in Figure 3. The figure also summarizes the relevance of fluid sampling related to the parts of the process being studied.

It was strongly believed that monitoring the destructive process would identify a biomarker that would be increased with progression and decreased by effective therapy. Assays were developed to detect elastin-breakdown products/peptides and the cross-linking peptides, desmosine and isodesmosine. Lung elastin is a very long-lived connective tissue, and hence, breakdown products should not be a feature of health. Unfortunately elastin is not confined to the lung, and these elastin-breakdown products, despite being elevated in patients with neutrophilic lung diseases, do not help with the management of even AATD. There is some evidence that these elastin-breakdown products increase with disease progression but do not mark any therapeutic benefit. Furthermore a variety of other inflammatory lung diseases and variable clinical states also affect the levels of those markers, and for many reasons, their validity remains controversial. It is possible that only specific elastase-generated fragments
would be useful and hence results may depend on both enzyme-cleavage preparation of antigen and monoclonal antibody. More recently, renewed interest in the role of desmosine (an elastin cross-linking peptide) has occurred, reviewed extensively by Luisetti et al and shown to relate to lung physiology. The relationship with spirometry was weak but that to gas transfer was stronger, consistent with elastin breakdown being more a feature of emphysema. However, studies of elastin breakdown may be more relevant if assessed in airways secretions. So despite the studies that evaluated lung elastin-breakdown products as far back as the 1970s and strong evidence that this is a key process in the development of emphysema, its validation as a biomarker remains an unresolved problem.

**Elastase and other proteinases**

The release of elastase from activated neutrophils within the lung tissues is necessary to cause the proteolytic damage implicated, at least, in emphysema. This would seem, therefore, to be a much more relevant and direct biomarker of this clinical/pathological phenotype. The problem is that although elastase release in the lung interstitium is likely to be critical for matrix degradation at this site, the enzyme is rapidly inactivated (even in AATD), especially by local inhibitors, even if already bound to elastin. An alternative approach has been to measure the enzyme activity in the lung secretions. Sputum elastase activity is detectable in usual COPD but only during neutrophilic exacerbations, again reflecting the capacity of the local inhibitors. However, although elastase activity is more easily detectable in AATD patients, even in the stable state, it is also present in the secretions of bronchiectasis patients and without cystic fibrosis. Since these latter diseases do not develop emphysema, the data suggest neutrophil recruitment more locally (bronchial circulation), sparing the interstitium, in response to bacterial load, and overwhelming local- and serum-derived inhibitors play a role, potentially damaging the airway epithelium and local host defenses.

Indeed, other enzymes are often detected in these same secretions, including proteinase 3, metalloproteinases, and cysteine proteinases, all of which have been implicated in emphysema but which also form part of an independent proteinase cascade. Whatever the final pathway, all these enzymes and their inhibitors may become both specific and nonspecific biomarkers. Thus, whether these markers provide insight into airways disease above and beyond simple associations or merely reflect sputum purulence (see Exacerbations section) remains to be determined.
For the emphysema process, bronchoalveolar lavage (BAL) would be the only viable option for the study of markers in the airway. Although this approach has been used for more general inflammation markers (see General inflammation markers section), it has added little to the data because of the difficulty in obtaining a true alveolar sampling. However, in AATD, lavage, AAT, and elastase inhibitory capacity have been used to support the benefit of AAT augmentation therapy,\(^5\) which has also been reflected in sputum studies.\(^5\)

More recently, an alternative approach to the detection of lung elastase activity has been developed and at least partly validated. As enzyme is released from neutrophils, there is a finite area and time within which the enzyme remains active. Specific cleavage of the surrounding substrate can produce a biochemical footprint of enzyme activity. A specific cleavage product of fibrinogen (A\(_\alpha\)- Val360) has been shown to relate to lung physiology, be reproducible, increase during exacerbations, and to decrease with augmentation therapy, in AATD.\(^5^2\) Concentrations were also found to be higher than normal in usual COPD and to be related to physiological and radiological measure of emphysema as well as radiological progression of emphysema in the lower zone,\(^5^3\) this suggests a similar elastolytic process is at least partly responsible even in the presence of normal AAT. These latter findings may well relate to the abnormal neutrophil function seen in usual COPD (see The neutrophil section). Whether studies of similar markers of other elastolytic enzymes will also be informative remains to be seen. However, of interest, it should be noted that as part of the proteinase cascade, neutrophil elastase (NE) is responsible for the activation of the cysteine proteinase cathepsin B.\(^5^4\) Although this enzyme is not inhibited by AAT, the inhalation of AAT, which would inactivate NE, was also shown to result in the inactivation of cathepsin B,\(^5^5\) suggesting the latter is also a downstream biomarker of local NE activity. In the same vein NE activates matrix metalloproteinases (MMPs),\(^5^6\) and these enzymes are responsible for tumor necrosis factor (TNF)\(\alpha\) processing,\(^5^7\) which, in its own right, drives neutrophilic inflammation. Thus, understanding these parts of the inflammatory cascade provides, not only potential therapeutic targets but also, potential biomarker readouts of efficacy (Figure 4).

**The neutrophil**

Since the identification of NE as a potential mediator of emphysema, studies of this cell have become central to our understanding of the end point of tissue destruction in this subtype of COPD. There is extensive literature documenting evidence of the activation of neutrophils and their adhesion molecules in COPD. However, few of these studies have determined whether this is COPD-specific or reflects lung neutrophilic disease states in general. However, airway neutrophils have been shown to relate to the decline in lung function\(^5^8\) and emphysema quantified by CT scan.\(^5^9\) Nevertheless, the findings of such studies (as with all inflammatory mediators) beg the question of whether the observations are physiological, caused by the “severity” of the disease, or pathological, causing the severity of the disease itself. The data implicating the neutrophil is extensive but remains circumstantial. Although it is recognized that the degree of emphysema relates to the amount of NE in the lung\(^6^0\) and that NE destroys elastin, for this to be pathological would require that there be an abnormal response but not in those not demonstrating the phenotype (ie, healthy age-matched smokers). AATD provides another control since the pathological mechanism (low AAT) is well established, and disease severity (to obviate any physiological response) and treatment (to overcome any influences of drugs) can be matched to patients with usual COPD.

![Figure 4](https://www.dovepress.com/)

**Figure 4** Interrelation of proteases and TNF. Cigarette smoking leads to TNF release and sequential events leading to emphysema (1). Serine proteases released by recruited neutrophils activate MMP12 and inactivate its’ cognate inhibitor’s (2). MMP12 inactivates αAT facilitating its own activation by serine proteases (3). MMP12 then leads to extracellular processing of the interaction of TNF with its receptor (4) facilitating the main pathway (1).

Abbreviations: α1 AT, α1-antitrypsin; CT, computed tomography; MMP12, matrix metallopeptidase 12; TIMP, tissue inhibitor of metalloproteinase; TNF\(\alpha\), tumour necrosis factor α.
It was recognized in 1987 that the polymorphonuclear leukocytes (PMN) from emphysema patients with COPD behaved abnormally. The cells demonstrated a rapid motility and an excessive ability to destroy connective tissue.\(^{61}\) This was related to increased endothelial adhesion under-flow\(^{62}\) and reflects a defective chemotactic response to routine chemoattractants.\(^{63}\) These features are not found in healthy smokers and matched patients with AATD. The defect has been linked to the abnormal expression of phosphoinositide 3 (PI3) kinase and may reflect excessive production by phosphatidylinositol (3,4,5)-tri-phosphate (PIP\(_3\)), a genetic defect of regulation, or defective breakdown by the phosphates, phosphatase and tensin homolog (PTEN) and SH2 domain-containing inositol 5'-phosphatase 1 (SHIP1). Thus further studies are required to determine the exact mechanism, but it does reflect a pathologically relevant trait that needs further validation, including determining whether this is a feature of susceptible smokers alone and relates to the rate of progression. Nevertheless, recent data has shown that activated neutrophils in usual COPD colocalize to the area of the lung where the emphysema develops.\(^{59}\) The signal is also clearly different to healthy controls and AATD patients and provides a platform for early Phase II studies of neutrophil-modifying therapies.

This whole pathophysiological hypothesis of the role of the neutrophil in the development and progression of disease (at least of emphysema) has therefore a long and well-established history based on a human genetic model. With all this knowledge, the potential for biomarker discovery would seem relatively straightforward, and yet, no such validated biomarker that fulfills all the necessary requirements (marks the disease process, reflects the activity of the disease process, is stable, predicts progression, and identifies patients amenable to therapy) has emerged. With this lesson as an example, all other markers need similar understanding and background before true advances in specific disease management can occur.

The eosinophil
The eosinophil is more classically associated with asthma. However, one study in COPD has shown that an increase in the proportion of eosinophils in induced (or spontaneous) sputum is indicative of a good spirometric response to inhaled corticosteroids.\(^{64}\) Furthermore a recent publication has also indicated that this marker (reflected in blood eosinophils) also identifies patients who are likely to benefit from oral steroid therapy for exacerbations.\(^{65}\) Importantly, the study also suggested that the noneosinophilic group fared less well with steroid therapy. These studies would be consistent with an asthma/COPD overlap syndrome and provide guidance for a more steroid-orientated treatment strategy. Hence, eosinophils can be seen to fulfill one of the requirements for a biomarker. Although bronchodilator reversibility, per se, influences spirometric decline,\(^{66}\) it remains unknown whether it is the eosinophilic subset that marks the susceptible, yet steroid-responsive patient or predicts future progression.

Oxidative stress
The other key process implicated in the pathophysiology of COPD is oxidative stress. Cigarette smoke itself contains high concentrations of oxygen-free radicals with the potential to damage tissues and proteins, and stimulate inflammation. Despite an efficient antioxidant defense, there has been clear and repeated evidence of oxidant stress in COPD. Again, whether this is pathological or physiological (a general response-to-smoking) remains unclear. This topic was extensively reviewed in 2008 and more recently in 2013,\(^{67}\) and the oxidant and proteinase story has been brought together in a further recent review.\(^{59}\) However, to date, the markers of oxidative stress have not been widely used other than to reflect the process itself. Validation of these markers and their use in supporting effective antioxidant therapy is awaited.

Genetic studies have, however, cast some light on the potential pathological, rather than physiological, nature of the ability to deal with oxidative stress (see Technology platforms section).

Inflammation markers
Although the preceding sections addressed relatively specific marker studies in COPD that were based on clinical and pathophysiological processes, the rapidly expanding literature on biomarkers has taken a more generic stance. It is widely accepted that the majority of diseases in the COPD syndrome have inflammation as the key underlying mechanism, and studies of a vast array of mediators have been undertaken, addressing three distinct questions.

1. Is it/are they elevated in COPD?
2. Does it/do they relate to the physiological markers (usually spirometric) of severity?
3. Does it/do they reflect a specific patient phenotype or a therapeutic target?

Using the search terms “biomarkers and COPD”, in excess of 1,000 papers can be found that have been published since 2008. It is clearly beyond the scope of the current review to discuss such a large body of literature, so selective aspects have been chosen to illustrate the general principles required for interpretation (as few have emerged
to answer most of the questions outlined in the introduction). C-reactive protein (CRP) is probably the most nonspecific marker of inflammation and yet, has been widely studied in COPD. At present, over 300 articles can be identified via PubMed, using the search words of “CRP and COPD”. CRP levels, even those within the accepted normal range, predict future cardiac events in a general population, and they are implicated in the pathophysiology of vascular disease.1 There is an inverse relationship between plasma CRP levels and lung function, even in subjects assumed to be healthy.71 The levels are elevated in COPD patients without evidence of ischemic heart disease; these are stable and are reduced with inhaled corticosteroids, although this does not reflect a disease-modifying effect.

There has been no clear mechanism to implicate CRP in the pathophysiology of COPD, nor does it reflect known genetic polymorphisms, suggesting it is merely a general marker of the underlying inflammatory process associated with COPD.73 However, the data did suggest that high CRP was associated with an increased odds ratio for hospitalization, although probably reflecting disease severity. The conclusions were based on a large population cohort and as such, only indicate trends and will have little impact on managing individual patients.73

CRP levels rise during exacerbations particularly when there is an increased neutrophilic influx due to a bacterial cause.74 In addition, a raised CRP in the stable state predicts recurrent exacerbations due either to a failure to completely resolve the first episode or a persistent underlying airway colonization that predisposes to further episodes.75

Fibrinogen has also been widely studied in COPD and has been well covered in a recent review.76 Fibrinogen is also an acute-phase protein and is raised in COPD (as part of the systemic effects of the disease), increases during exacerbations, and is higher in those with frequent exacerbations and those with more severe airflow obstruction. It may also play a role in comorbidities, especially vascular disease and potentially cardiovascular death due to an increased clotting tendency. Again, no genetic polymorphisms have been identified, indicating fibrinogen is also more likely to be a reflective feature of COPD. Although studies indicate that therapy can reduce fibrinogen levels, it remains unknown whether this affects progression or other outcomes. If so, fibrinogen could act as an ideal biomarker at least for some aspects of COPD, and its management and further studies are awaited. The references related to these data are included in the review article.76

Many other inflammatory markers have been measured in COPD, and some seem to be persistently raised. The nonspecific nature of these changes provides little insight into management or pathophysiology. Surfactant protein D (SPD) is a lung-derived protein. SPD is elevated in the plasma in COPD and falls with corticosteroid therapy,77 suggesting that steroids reduce systemic leak from the lungs as part of a general anti-inflammatory effect. However, in one study, the changes in SPD following inhaled therapy were also reflected in the degree of improvement of health status and FEV1, in the short term,78 suggesting that lung inflammation itself affects these features. Of interest, the CRP and interleukin (IL)-6 did not change, suggesting that the effects were acting locally in the lung and not influencing other aspects of generic systemic inflammation (although the SPD effect could be measured systemically).

Studies are beginning to address the requirements of an ideal biomarker, with therapeutic interventions for Phase II and Phase III studies in mind. For example, fibrinogen was used as a marker in a Phase II study of p38 mitogen-activated protein kinase (MAPK) inhibitors,79 which effectively block the proinflammatory pathway. The treatment clearly had an effect, as determined by reducing fibrinogen levels. However, whether this pathway is merely physiological, rather than pathological, remains unknown, and hence, fibrinogen still remains uncertain as a therapeutic marker option in the long-term or short-term management of COPD. Nevertheless, this study represents a first step toward the development and assessment of new therapies using a biomarker, perhaps with a cardiovascular benefit (in those with this comorbidity), rather than a pulmonary one.

Other inflammatory markers are measurable and elevated in the plasma, and although none have proven to be specific, an alternative has been to measure several to develop a multicomponent approach.80 Using the Evaluation of COPD Longitudinally to Identify Predictive Surrogate Endpoints (ECLIPSE) cohort,80 the authors measured several blood biomarkers that were increased in COPD patients compared with healthy smokers and nonsmokers. The white blood cell (WBC) count, CRP, and fibrinogen levels fitted this criterion, as did IL-6, IL-8, and TNFα. The proportion outside the 95th percentile of nonsmokers was networked at baseline and at 1-year follow-up to determine consistency. A subgroup (16% of the patients) with persistently elevated inflammatory markers (two or more) was identified who had a higher incidence of exacerbations (cause or effect?) and worse survival.

This approach of selecting certain markers or traits to identify patient groups with common features is the basis for “cluster analysis”, which may be informative but clearly depends on the traits or factors selected and always needs validation in a second patient cohort; such studies are awaited.
The ECLIPSE cohort has been used to study a large number of other potential biomarkers. In 2011 Dickens et al reported on the stability of 34 blood biomarkers over 3 months. The importance of stability is that single measurements could be informative of prognosis and particularly, may reflect the underlying disease activity that leads to progressive deterioration. Fifteen of the markers were different in COPD compared with the controls, and four reflected patients who had exacerbations between baseline and follow-up (CRP, fibrinogen, IL-6, and SPD), but few were classified as stable and hence useful as markers of baseline disease activity (fibrinogen, Clara cell protein [CC]16, and SPD). Although there were weak correlations of the other markers with spirometry, fibrinogen, which did not correlate, emerged as the most promising as it was stable and was a marker of a frequent exacerbation phenotype, exercise impairment, the severity of the BODE index, and dyspnea.

However, of these “stable” markers, only CC16 showed both a relationship to baseline spirometry and the subsequent annual decline. At present, only this protein and the Ato-Val360 (see Elastase and other proteinases section) biomarker have shown such characteristics. CC16 is an immunosuppressant and antioxidant produced in the airways. The fact that it is elevated in COPD suggests it is reflective of the disease, rather than causative; however, it is possible that its measurement in serum (like SPD) reflects leakage from the lung due to local inflammation, which may be the generic process associated with, rather than causing, decline. Nevertheless, this series of studies demonstrates several of the steps needed to validate a biomarker, which if subsequently implicated in the pathophysiology, will also direct thinking to new specific therapeutic targets.

SPD levels in blood are also raised in COPD and respond to steroid therapy as indicated above. Since low SPD levels in the lung have been implicated in the pathophysiology of COPD, are reduced in BAL in COPD, and relate to spirometry, it fulfills more of the pathophysiological role than does CC16. However, although SPD responds to oral steroids with a decrease in plasma levels, it is unknown what happens in the lung in the same patients. Nevertheless, in a separate study, COPD was associated with low lung levels of SPD, and inhaled steroid therapy was associated with higher levels, suggesting that SPD is a good marker of disease and response to treatment. Subsequently, a polymorphism of the SPD gene was identified that also relates to COPD and that is reflected in the level of SPD – at least in the serum (transcription and lung levels are currently unknown).

This latter point is of importance: whereas this appears to “close the loop”, it should be noted that a functional polymorphism of SPD that is of pathophysiological importance should result in low lung levels of SPD, but the presence of airway inflammation and leak may, counterintuitively, lead to high blood levels. Nevertheless, the studies do suggest the potential for local therapy with SPD in COPD and that it is certainly worth pursuing.

**Sampling issues**

Studies of the pathophysiology of COPD are made all the more complex because of the different pathological phenotypes and the fact that the site of damage/repair is in the local tissues. Clearly this is not an issue if the biomarker is either genetic (as in AATD), or systemically reflects the disease process or specifically samples the disease area. To date, biomarkers have been studied in exhaled breath condensate, spontaneous or induced sputum, BAL, or serum/plasma. All of these have inherent problems in interpretation, especially when using high-throughput methodology.

Firstly, many commercially available assays have not been validated using the biological medium collected and processed for the measurement (especially airway secretions) and can produce inaccurate results. Secondly, the mediator measurement (especially in exhaled breath condensate) may be detectable and often below the lower limit of quantification. Thirdly, the biological medium may not specifically sample the area where the disease process is occurring. For instance, sputum or induced sputum is unlikely to represent samples derived from the small airways or alveoli, and BAL does not exclusively sample the small airways or alveolar region. In addition, the variability between samples from the lung is often wider than the differences between subjects or clinical states. For instance, markers in spontaneous sputum may have a day to day variability of up to 200%, for many reasons. Obtaining samples sequentially and averaging the results drastically reduces this variability, if the patient is stable; however, this approach cannot easily be applied to invasive procedures, such as BAL or induced sputum, since the introduction of saline or hypertonic saline itself is proinflammatory, and several days are needed between sampling to enable a return to the stable state. However, serum or plasma stability is much more reliable, at least for some biomarkers. Pathological specimens provide an opportunity to sample individual cells and their nature/metabolic activity, by laser capture methodology, but other sampling techniques provide varying information from normal and abnormal tissue and may not reflect the pathological area unless performed on lung resection material within defined pathological areas. Finally, mediators measured in blood samples may reflect processes occurring in the interstitium,
recirculated from lymph but not necessarily the airways. The subsequent volume of distribution, not only dilutes mediators but also, clears them by appropriate receptor binding. Even then, studies like proteomics and metabolomics will differ in plasma compared with serum, due to coagulation and also, by sample collection, distribution, processing, and storage. Careful matching with appropriate controls and interpretation of data is therefore essential in all studies but is rarely done or reported, making the comparison between studies relatively tenuous.

**Exacerbations**

Exacerbations are an important contributor to health status, health care utilization, and death, particularly if leading to hospitalization. The definition of this symptom-led event, its complexity, and the broad approach to treatment and prevention has generated a search for an objective measure of an exacerbation and therapeutic guidance.

In general, exacerbations are events that are associated with an increase in inflammation, both in the lung and systematically. However, in general, the events may reflect a bacterial or viral cause or be related to other causes of increased airway obstruction or may even reflect the destabilization of other organ systems, such as in associated cardiovascular decompensation. The main decision required for such events is whether to increase bronchodilator therapy alone, to treat with corticosteroids or antibiotics, or both. Of these, the use of antibiotics has received most attention because overuse is likely to lead to the continued development of bacterial resistance.

Serum procalcitonin is normally produced by neuroendocrine cells of the thyroid and lung but is also produced elsewhere in the body, in response to bacterial infection. For this reason, it has been used to predict bacterial infections and outcome in patients with sepsis. However in general this approach has been unsuccessful in COPD. Serum amyloid A has also been suggested for this as it is increased during exacerbations with a viral or bacteriological aetiology; however, it failed to differentiate them, despite reflecting severity of the episode, and hence would not facilitate treatment guidance.

To date, the best two markers that influence therapy are symptom deterioration with known or current eosinophilia, which likely benefit from steroid therapy, or sputum color. The latter reflects the presence or absence of a large neutrophilic load, imparting various shades of green. A change in color reflects an increase or decrease in sputum neutrophilia. The color can be characterized at least semiobjectively by both the patient and health care worker, using a standard color chart. In the initial study using this approach in primary care, there was high sensitivity to bacteria isolated from sputum and in particular, an increase relating to the colonizing microbial load, even in the stable clinical state. This approach, using sputum color, enabled antibiotics to be withheld without detriment from patients with white or colorless sputum and also enabled monitoring of the antibiotic response by the patients. Subsequent studies have confirmed the robust nature of this approach in identifying patients with significant colonization and also, that withholding antibiotics from those without yellow or green sputum had no adverse effects. Interestingly the authors found that CRP and not procalcitonin was increased in patients with purulent sputum, suggesting that CRP may be a better discriminator for bacterial infection. Clearly, further clinical trials are indicated to consolidate this approach.

**Technology platforms and the “omics” revolution**

In recent years, the ability to obtain multiple assay kits commercially and the development of the proteomics, metabolomics, transcriptomics, and genomics approaches to patient assessment has led to an overwhelming plethora of data. This does lead to some technical issues: the assay may not have been validated for the biological sample or the processing methodology used may lead to spurious and unrecognized effects on the result, both within and between patients. In addition, the data generated has a sensitivity that far outweighs our current ability to phenotype patients sufficiently from a clinical, physiological, and radiological viewpoint. It may be that this approach will lead to a pathophysiological classification of COPD that negates clinical phenotyping, although this remains to be determined and would have to be validated against the clinical “gold standard”.

Proteomics (with 75 citations in the past year) is based on the identification of COPD proteins in biological fluids by a multitude of techniques, which produce a two-dimensional spectrum of data (Figure 5) to determine the similarities or differences between subjects. This demonstrates the amount of data that can be generated by the “omics” approach. The benefit is that this is not a “hypothesis-led” approach, and new data can emerge as “hypothesis generating”. Nevertheless, such an approach can lead to false-positive and false-negative results, and new statistical approaches to the data are needed.

Indeed it has even been suggested that combining the data from different “omics” platforms, as part of an integrated
Figure 5 A two dimensional proteomics read-out. Each dot/blot represents a single peptide. Changes in density/presence are considered as biomarkers of the disease progress or activity.

system biological approach to understanding lung diseases, may be necessary to make progress. The methodology is advancing at great pace. The review by Nicholas et al defines the principles and aims of the methodology and is worth reading, for those interested.

Less has been published on metabolomics, but in a small study of smokers with emphysema, the assessment of 3,534 metabolite markers was able to separate smokers with and without emphysema, and the top 12 were used for predictive models, some of which produced clear separation (as good as CT scan?).

Genetic studies have dominated the COPD world since the early 2000s, with both a candidate gene approach and genome-wide association studies. The data have aimed at detecting the genetic markers of COPD predisposition, thereby leading to new therapies based on gene function (over- or underactive) or a preventative approach to the disease(s). Many of the genetic markers failed to identify polymorphisms that influence the targeted gene function, but associations with COPD or its features suggested the marker was in linkage disequilibrium, probably with a nearby functional abnormality. Such susceptibility markers known to be associated with functional abnormalities include matrix metallopeptidase 9 (MMP9) and apical emphysema, TNFα and bronchitis, superoxide dismutase and emphysema, iron-responsive element binding protein 2 (IREB2) and COPD, and IL-13 promoter polymorphisms and an interaction with cigarette smoking on FEV1. Nevertheless, such associations represent only minor contributors to the COPD complex that could reflect either the superficial phenotyping of patients or indeed the complexity of the disease(s).

Such studies do reflect potential susceptibility genes and hence, pathophysiological pathways, but how or whether they reflect the rate of progression or long-term outcomes has yet to be determined. Nevertheless, in other diseases, treatments have been influenced by genetic studies, suggesting the same may follow for COPD, at least for those associated with a functional contribution. It may be that a panel of genes or a panel of gene transcripts may provide further insight into progression or the response to treatment, although far from personalized medicine. Interestingly, a recent study has suggested that a polymorphism in the a disintegrin and metalloprotease domain 33 (ADAM33) gene is associated with all-cause and cardiovascular mortality in COPD. The group of proteins encoded by this gene domain play important roles in cell adhesion, migration, and proteolysis, and ADAM33
Table 1  Biomarkers in context

<table>
<thead>
<tr>
<th>Sample</th>
<th>Measures</th>
<th>Context</th>
<th>Problems</th>
<th>Benefit</th>
</tr>
</thead>
<tbody>
<tr>
<td>Plasma/serum</td>
<td>Cytokines/peptides/products</td>
<td>Systemic inflammation</td>
<td>Coagulation may activate cells/proteins/peptides</td>
<td>Easy to obtain</td>
</tr>
<tr>
<td></td>
<td>Acute phase proteins</td>
<td>Systemic response</td>
<td></td>
<td>Repeated sampling acceptable</td>
</tr>
<tr>
<td></td>
<td>Cell activation markers and secreted/shed products</td>
<td>Inflammatory cells</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Tissue damage products</td>
<td>Tissue damage</td>
<td>Distant from the lung</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Lung leakage</td>
<td>Lung inflammation</td>
<td>May not be lung specific</td>
<td>No dilutional problems</td>
</tr>
<tr>
<td>Exhaled breath</td>
<td>Cytokines</td>
<td>Airways/oropharyngeal inflammation</td>
<td>Many measures are below the lower limit of quantification/detection</td>
<td>Repeated sampling</td>
</tr>
<tr>
<td>condensate</td>
<td>Markers of oxidant stress</td>
<td>pH as a marker of inflammation</td>
<td>Site not localized</td>
<td></td>
</tr>
<tr>
<td>Lung lavage</td>
<td>Cytokines/peptides</td>
<td>More localized to the lung</td>
<td>Variable dilution by condensate</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Markers of oxidant stress</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Cells number and function</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Protein leakage from plasma</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Damage markers</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sputum/induced</td>
<td>Cytokines/peptides/proteins</td>
<td>More localized to the lung</td>
<td>Samples bronchial secretions</td>
<td>Spontaneous sputum is readily repeatable</td>
</tr>
<tr>
<td>sputum</td>
<td>Markers of oxidant stress</td>
<td></td>
<td>Oropharyngeal contamination</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Cell number and function</td>
<td></td>
<td>Variable dilution</td>
<td>Variability can be reduced by sequential sampling</td>
</tr>
<tr>
<td></td>
<td>Damage markers</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>DNA</td>
<td>Variation in genetic sequence</td>
<td>Underlying susceptibility and pathophysiology</td>
<td>Induced is proinflammatory</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Physiology</td>
<td>End-organ damage</td>
<td>Assesses the change in lung function as a result of damage</td>
<td>Does not determine the degree, site, or pathological damage with precision</td>
<td>Reasonably well understood</td>
</tr>
<tr>
<td>CT scan</td>
<td>Lung density</td>
<td>Localizes abnormalities that influence clinical outcome</td>
<td>Radiation exposure</td>
<td>Correlates with health status, activity, and mortality</td>
</tr>
<tr>
<td></td>
<td>Changes in architecture</td>
<td></td>
<td></td>
<td>Identifies pathology</td>
</tr>
<tr>
<td>Biopsy</td>
<td>Cells</td>
<td>Lung tissue</td>
<td>Variability between samples</td>
<td>Sensitive to emphysema progression</td>
</tr>
<tr>
<td></td>
<td>Architecture</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Cytokines</td>
<td></td>
<td>Not easy to repeat</td>
<td>It is the most direct way of studying the process in situ</td>
</tr>
<tr>
<td></td>
<td>Damage products</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Abbreviations: CT, computed tomography; DNA, deoxyribonucleic acid.

has also been implicated in the rapid decline in lung function and development of COPD.112,113 Whether this reflects a functional abnormality and hence, a treatment target needs to be determined.

Summary

Biomarker discovery has become a rapidly expanding field of research primarily to develop easy and early readouts for delivery of Phase II clinical studies and a shortening of pivotal Phase III studies that displace conventional markers like FEV1. A variety of methodologies and sample collections have been undertaken, measuring things as diverse as the mechanics of the lung, patient symptomatology, markers of exposure (such as inhaled oxidants from cigarette smoke), markers of tissue damage (connective tissue degradation products), number and function of implicated cells, generalized inflammatory markers, cell-activation products, underlying proteins (their function, metabolism, and source), and the genetic background of the patients. These have involved a variety of biological samples collected in different ways, all with both benefits and drawbacks and all requiring caution in interpretation, as summarized in Table 1. However, the identification and understanding of such markers, either in a hypothesis-testing or hypothesis-generating approach, may prove sufficiently informative for understanding the disease process and for developing new therapeutic strategies. Nevertheless, without accurate patient characterization, biomarker validation, and an understanding of the issues that likely influence the measurements (including current therapy and dissecting cause and effect), this process may prove to be both fruitless and confusing.

Disclosure

The author has lectured widely as part of pharmaceutical sponsored symposia, sat on numerous advisory boards for drug design and trial implementation, and received
noncommercial grant funding. The author reports no other conflicts of interest in this work.

References


