Rationale: Markers of inflammatory activity are important for assessment and management of many respiratory diseases. Markers that are currently unrecognized may be more valuable than those presently believed to be useful.

Objectives: To identify potential biomarkers of suppurative and inflammatory lung disease in induced sputum samples.

Methods: Induced sputum was collected from 20 healthy control subjects, 24 patients with asthma, 24 with chronic obstructive pulmonary disease, 28 with cystic fibrosis (CF), and 19 with bronchiectasis. Twelve patients with CF had sputum sampled before and after antibiotic therapy for an infective exacerbation. The fluid phase of induced sputum was analyzed by surface-enhanced laser desorption/ionization time-of-flight (SELDI-TOF) mass spectroscopy on three protein array surfaces. Some protein markers were selected for identification, and relevant ELISA assays sought. For 12 patients with CF, both SELDI-TOF and ELISA monitored changes in inflammatory responses during infective exacerbations.

Measurements and Main Results: SELDI-TOF identified potential biomarkers that differentiated each of the disease groups from healthy control subjects: at a significance of \( P < 0.01 \), there were 105 for asthma, 113 for chronic obstructive pulmonary disease, 381 for CF, and 377 for bronchiectasis. Peaks selected for protein identification yielded calgranulin A, calgranulin B, calgranulin C, Clara cell secretory protein, lysozyme c, proline rich salivary peptide, cystatin s, and hemoglobin \( \alpha \). On treatment of an infective CF exacerbation, SELDI-TOF determined falls in levels of calgranulin A and calgranulin B that were mirrored by ELISA-measured falls in calprotectin (heterodimer of calgranulins A and B).

Conclusions: Proteomic screening of sputum yields potential biomarkers of inflammation. The early development of a clinically relevant assay from such data is demonstrated.

Keywords: biomarkers; calprotectin; cystic fibrosis

Although there is considerable clinical and research need for good assessment of inflammation in airway diseases, the requirement of relatively invasive procedures, such as bronchoscopy and bronchoalveolar lavage (BAL), precludes sampling from a wide range of patients and on repeated occasions. This has encouraged investigators to use techniques of exhaled gases and exhaled breath condensate (EBC) analysis, although there may be limitations of value (1–10). The cellular phase of induced sputum is helpful in evaluating and monitoring asthma (11), but its use in suppurative lung diseases such as cystic fibrosis (CF) is less clear, although in children induced sputum measurements have been shown to correlate with lung function (12). The fluid phase of sputum has been rather underutilized; however, potential problems of variability in noncellular material may be encountered and as yet there is no agreed-upon correction factor as in BAL sampling.

Surface-enhanced laser desorption/ionization time-of-flight (SELDI-TOF) technology has been used to discover biomarkers of disease groups in direct comparison to control groups. The primary focus of much of this research has been to determine diagnostic biomarkers in a wide range of diseases, such as rheumatoid arthritis, HIV infection (13), prostate cancer (14), ovarian cancer (15, 16), motor neurone disease (17), ischemic heart disease (18), and infectious diseases (19, 20), using a diverse range of body fluids, such as serum, urine, cerebrospinal fluid, and joint fluid. The generated protein profiles demonstrate differences between disease groups and individual proteins may be identified to act as biomarkers either alone or in tandem with others. SELDI-TOF data may be used to generate serum “protein fingerprints” using bioinformatics to provide a potential diagnostic test in cancer and infectious diseases (15, 19, 20). This approach relies on interpretation of mass spectral patterns rather than identifying individual proteins and has been widely criticized in the literature and, in some cases, such as the work of Petricoin and colleagues (15), analysis of data sets independently have led to completely different conclusions (21). Baggerly and coworkers (21) demonstrated that inconsistent data processing can alter the ultimate outcome of these sorts of experiments. A different, and more direct, approach is to identify the actual proteins responsible for the mass spectral peaks provided by mass spectrometry (MS) data. Indeed, the identification of key biomarkers after initial proteomic studies may provide a more viable route to clinical application (22) and furthermore may allow the functional importance of such markers to be investigated. Therefore, using SELDI-TOF as a screening tool...
to identify individual potential biomarkers for further characterization may be a more valid and robust approach. SELDI-TOF has been applied to a small number of patients with chronic obstructive pulmonary disease (COPD) demonstrating a protein profile in BAL (23), and in pulmonary sarcoidosis (24). SELDI-TOF has been used to demonstrate protein expression in BAL in CF (25, 26). There is as yet a lack of published data using SELDI-TOF to profile sputum, although complementary proteomic approaches of two-dimensional electrophoresis (27), shotgun protein sequencing (28), and affinity immunoproteomics (29) have been applied to sputum in an effort to discover new biomarkers.

We applied SELDI-TOF to induced sputum from patients with inflammatory (asthma, COPD) and suppurative (CF, bronchiectasis) airway diseases to determine whether a high-throughput proteomics screening method can reveal differential protein expression between disease groups and therefore candidate biomarkers, which may then be developed into point assays of inflammation with potential application in the clinic. This work was driven by the need to develop biomarkers to monitor suppurative lung disease, in particular CF.

**METHODS**

**Subjects and Selection**

The Lothian Ethics Committee (Edinburgh, UK) granted ethical approval and all participants gave written consent. Sputum was obtained from 28 patients with CF, 19 with bronchiectasis, 24 with asthma, 24 with COPD, and 20 healthy control subjects. All patients were clinically stable and recruited from the outpatient clinic of a respiratory medicine unit. All subjects were recruited from specialist respiratory clinics at a major teaching hospital where they were attending with a diagnosis of their respective illness. In the absence of preexisting data on intersubject variability, formal statistical powering was not possible, but we approached this problem by recruiting similar numbers of subjects to previous biomarker studies. The patient groups represented respiratory diseases with clinical features that both complement and contrast with each other.

**Longitudinal Data**

To determine the usefulness of discovered markers in the assessment of lung disease during a changing level of inflammation, longitudinal samples were collected from 12 patients with CF during an infective exacerbation. Samples were collected at onset and completion of intravenous antibiotic therapy (duration, 14–21 d). The biomarkers were assessed both by SELDI-TOF and by ELISA.

**Induced Sputum**

Sputum induction was performed as previously described (30). Subjects inhaled nebulized hypertonic saline at concentrations of 3, 4, and 5%. Sputum was processed within 2 hours of collection (31). Sputum plugs were selected and processed with 4 × weight/volume of 0.1% dithiothreitol after which 4 × weight/volume of phosphate-buffered saline was added. Samples were filtered through 48-μm mesh and centrifuged to remove the cells. Supernatants were stored at −80°C until further analysis. Cytospins were stained with May-Grünwald-Giemsa for differential cell counting.

**SELDI-TOF MS**

Three different chromatographic chip surfaces were used to cover a wide range of protein characteristics: a weak cation exchange at pH 4 (CM10), an anion exchange at pH 10 (Q10), and an immobilized metal affinity surface activated with nickel (IMAC nickel [IMACNi]). Induced sputum samples were adjusted to contain 1 mg/ml protein at a concentration of 0.1% dithiothreitol. Ten microliters of sample were added to CM10 and Q10 surfaces in a bioprocessor unit (Ciphergen, Freemont, CA) and 1 μl of sample was added to the preactivated IMACNi surface for on-spot incubation (more reproducible for IMACNi surfaces). All chips were treated with sinapinic acid (SPA) matrix (2 × 0.8 μl/spot) and allowed to air dry. Samples were analyzed on the Protein Biology System 2 SELDI-TOF mass spectrometer (Ciphergen). Chips were read with a laser intensity of 205 with the deflector set at 4,000 D and a focus mass of 7,500 D. Profiles were then exported to Ciphergen Express (Ciphergen) for clustering and data analysis. Intra- and interassay coefficients of variation were assessed with the CM10 surface as outlined in the online supplement.

**Data Analysis**

Data were normalized to total ion current to take into account any spot to spot variability in chip surface binding. Data with high normalization coefficients were identified and individual spectra examined. Poor-quality spectra or absent signal were excluded from further analysis. Protein peaks were automatically clustered to identify biomarkers of similar molecular weight. Mann-Whitney rank testing was performed to demonstrate statistical differences in clusters between disease groups. Values of \( P < 0.01 \) were taken to be significant. For longitudinal data, paired \( t \) test analysis was performed using Prism 4 software (GraphPad, San Diego, CA). For a further description of methods, please refer to the online supplement.

**Protein Identification**

Highly significant peaks (\( P < 0.0001 \)), assessed on relative abundance and quality of original spectral data, were selected for protein identification. Pooled sputum was used. Samples were applied to 18% trisene/glycine acrylimide gel for electrophoresis. Bands of appropriate molecular weight were excised and destained and the protein eluted. The eluate was digested with proteomics grade trypsin (Sigma, Gillingham, UK), applied to a normal-phase chromatography chip surface and analyzed with SELDI-TOF MS or MALDI (matrix-assisted laser desorption/ionization)-TOF MS using a Voyager DE STR mass spectrometer (Applied Biosystems, Foster City, CA). The resultant mass fingerprint was used to identify the parent protein through reference to protein identification databases (MS-Fit; University of California, San Francisco, San Francisco, CA). Proteins were identified based on molecular weight search (MOWSE) score and cross-checked with online MASCOT (http://www.matrixscience.com), which allows a probability-based MOWSE score. Protein identification was only accepted when \( P \) values were less than 0.05. Tandem MS/MS analysis was performed to confirm identification of proteins using a Q-star tandem MS instrument (Applied Biosystems) with SELDI chip interface (Ciphergen). This approach provided additional protein identification based on amino acid sequence from the MS/MS analysis.

Commercial antibodies, when available, were used for protein confirmation by Western blotting. For calgranulins A and B, we were also able to use an in-house quantitative immunooassay measuring calprtecin (calgranulin A/B complex). An in-house double antibody sandwich ELISA (antibodies and calprotectin standard courtesy of E. Sundrehagen, M.D.) was developed for use in the 12 patients with CF monitored longitudinally, and results compared with data from SELDI-TOF.

**RESULTS**

**Demographics**

Patient demographics are given in Table 1. Of the patients with CF, 18 were chronically infected with *Pseudomonas aeruginosa*.

**Reproducibility of SELDI-TOF Assay in Sputum**

This was assessed as described in the online supplement. The intraassay coefficient of variation ranged from 11.5 to 44% with an average coefficient of variation (CV) of 22.4%. Interassay variation demonstrated a CV ranging from 6 to 43%, with an average CV of 16.2%.

**Protein Profiles Generated by SELDI-TOF**

An example of generated data is demonstrated in Figures 1 and 2. Cluster analyses determined a large number of peaks present in
relative proportions that differentiated between disease groups and healthy controls (Table 2). Three chip surfaces yielded a total of 621 clustered protein peaks, 105 of those significantly different at $P < 0.01$ and 56 at $P < 0.001$ for asthma versus healthy controls. For COPD compared with healthy controls, the respective figures were 625 protein peaks, with 113 significant at $P < 0.01$ and 50 at $P < 0.001$ for three surfaces. For bronchiectasis versus healthy controls, the respective figures were 671 protein peaks, with 377 significant at $P < 0.01$ and 314 at $P < 0.001$ for three surfaces. For the CF group, the respective figures were 660 protein peaks, with 381 significant at $P < 0.01$ and 315 at $P < 0.001$ for three surfaces (Table 2; for full summary, see Table E1 in the online supplement).

Although all four groups of patients yielded a substantial number of differential proteins compared with healthy control subjects, there were much closer similarities between the two obstructive airway diseases and between the two suppurative airway diseases. For asthma and COPD, only 16 proteins, on the three chip surfaces, were different at $P < 0.01$, and one protein was different at $P < 0.001$ (Table 2; Table E1). This latter protein, molecular weight (MW) 29 kD, was seen on the IMACNi surface (Figure 3A) but had low signal intensity. This marker has not yet been identified. CF and bronchiectasis demonstrated relatively similar biomarker profiles compared with each other, albeit the abundance of protein in the CF group was greater. Fifty-eight proteins differentiated the groups at $P < 0.01$ and 9 proteins differentiated the groups at $P < 0.001$ (Table 2; Table E1) over the three surfaces. Five of these nine proteins were observed on the IMACNi surface, two on CM10 and two on Q10. The marker that showed the greatest separation between the disease groups and had the highest signal intensity had an MW of 12.246 kD (Figure 3B). We have not identified this protein.

![Figure 1](image-url)  
*Figure 1.* This figure demonstrates spectra from individual subjects on the immobilized metal affinity chromatography (IMAC) nickel surface in the mass range of 5 to 15 kD. Even in this mass range, there is an abundance of peptide peaks, and noticeable differences are observed between groups. BIectasis = bronchiectasis; CF = cystic fibrosis; COPD = chronic obstructive pulmonary disease.

### Table 1: Demographic, Lung Function, and Sputum Neutrophil Data of the Patients Studied

<table>
<thead>
<tr>
<th>Group</th>
<th>Age (yr)</th>
<th>Sex</th>
<th>FEV$_1$ (% pred)</th>
<th>Sputum Neutrophil (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Asthma</td>
<td>47.8 (2.9)</td>
<td>15 F/8 M</td>
<td>82.3 (4.4)</td>
<td>50.6 (5.0)</td>
</tr>
<tr>
<td>COPD</td>
<td>65.2 (1.2)</td>
<td>10 F/14 M</td>
<td>57.7 (4.1)</td>
<td>79.1 (2.7)</td>
</tr>
<tr>
<td>Bronchiectasis</td>
<td>61.8 (3.5)</td>
<td>15 F/3 M</td>
<td>71.7 (11.6)</td>
<td>83.2 (5.6)</td>
</tr>
<tr>
<td>CF</td>
<td>28.8 (1.7)</td>
<td>8 F/19 M</td>
<td>59.2 (3.9)</td>
<td>91.6 (2.0)</td>
</tr>
<tr>
<td>Control</td>
<td>36.4 (2.1)</td>
<td>11 F/9 M</td>
<td>101.4 (3.1)</td>
<td>54.2 (5.3)</td>
</tr>
</tbody>
</table>

*Definition of abbreviations:* CF = cystic fibrosis; COPD = chronic obstructive pulmonary disease; F = females; M = males.

*Values are mean (SEM).*
The top 20 protein peaks that showed the most statistically significant differences between individual disease groups and healthy controls are shown in Table E2, indicating their molecular weight and the chip surface on which they were identified. There are common peaks found in asthma and COPD, and CF and bronchiectasis.

**Biomarker Identification**

The proteins we have formally identified from the induced sputum samples are calgranulin A, calgranulin B, calgranulin C, Clara cell secretory protein (CCSP, CC16, CCSP10, uteroglobin), lysozyme c, proline rich salivary peptide, cystatin s, and hemoglobin A (Table 3).

Calgranulins A, B, and C were highly abundant in CF and bronchiectasis, but were also seen in the other disease groups. These proteins are represented by mass spectral peaks at 10.831 kD, 12.700 kD, and 10.100 kD, respectively. These peaks were preferentially expressed on the IMACNi surface but were also bound to the weak cation exchange surface. Some binding of calgranulin A was also demonstrated on the anion exchange surface. A separate form of calgranulin A with a lower molecular weight (10.574 kD) was seen in the CF and bronchiectasis groups. This was identified by mass fingerprinting of both molecular weights after purification on sodium dodecyl sulfate-polyacrylamide gel electrophoresis (data not shown). The relative expression of calgranulin A across all groups is shown in Figure 4A and the relative expression of the smaller form of calgranulin A in Figure 4B. The presence of calgranulin A was confirmed with Western blotting in pooled sputum samples from all disease and control groups (Figure 4C).

In contrast to the calgranulins, CCSP was in lower abundance in all disease groups compared with controls, but to the greatest extent in CF and bronchiectasis. This had a mass spectral peak of 7.900 kD on the CM10 surface. The relative expression of CCSP across all groups is shown on Figure 5. CCSP was confirmed on Western blot (data not shown).

**Correlation of Biomarkers with FEV₁ and Sputum Cytology**

In the CF group, the most abundant biomarker, calgranulin A, showed a weak inverse correlation with FEV₁% predicted, which failed to reach statistical significance (Spearman $r = 0.63$, $P = 0.08$). There was no correlation of calgranulin A with the percentage of sputum neutrophils (data not shown).
TABLE 2. THE TOTAL NUMBER OF PEAKS DETECTED BY SELDI-TOF

<table>
<thead>
<tr>
<th></th>
<th>IMAC Nickel</th>
<th>Weak Cation Exchange (CM10)</th>
<th>Anion Exchange (Q10)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Asthma vs. control</td>
<td>4 (n = 22 vs. 20)</td>
<td>16 (n = 20 vs. 19)</td>
<td>36 (n = 16 vs. 18)</td>
</tr>
<tr>
<td>COPD vs. control</td>
<td>11 (n = 23 vs. 20)</td>
<td>13 (n = 22 vs. 19)</td>
<td>26 (n = 21 vs. 18)</td>
</tr>
<tr>
<td>CF vs. control</td>
<td>108 (n = 28 vs. 19)</td>
<td>111 (n = 22 vs. 19)</td>
<td>96 (n = 22 vs. 19)</td>
</tr>
<tr>
<td>Bronchiectasis vs. control</td>
<td>112 (n = 19 vs. 20)</td>
<td>103 (n = 17 vs. 19)</td>
<td>99 (n = 18 vs. 19)</td>
</tr>
<tr>
<td>CF vs. bronchiectasis</td>
<td>19 (n = 28 vs. 19)</td>
<td>18 (n = 28 vs. 17)</td>
<td>21 (n = 22 vs. 19)</td>
</tr>
<tr>
<td>Asthma vs. COPD</td>
<td>10 (n = 23 vs. 22)</td>
<td>5 (n = 21 vs. 20)</td>
<td>1 (n = 21 vs. 25)</td>
</tr>
</tbody>
</table>

Definition of abbreviations: CF = cystic fibrosis; COPD = chronic obstructive pulmonary disease; SELDI-TOF = surface-enhanced laser desorption/ionization time-of-flight.

Significantly different (P < 0.001) for disease versus control are shown. The two suppurative airway diseases, CF and bronchiectasis, and the two inflammatory airway diseases, asthma and COPD, were also compared in terms of numbers of differentiating proteins (P < 0.01). Proteins could be increased or decreased in their abundance. Spectra that failed normalization were excluded from analysis (see text). n = number of normalized spectra used in each analysis, as per order in table legend. The spectra failing to normalize were not the same for each analysis. The data are displayed for each of the three chip surfaces that were used in the study.

Longitudinal Assessment of Identified Biomarker and Application of Specific ELISA

There were decreases in the levels of both calgranulin A and B, as assessed by SELDI-TOF, during treatment of an exacerbation of CF (Figures 6A and 6B; P < 0.01). Calprotectin (heterodimer of calgranulins A and B), measured by ELISA, similarly showed significant reduction (Figure 6C).

DISCUSSION

The use of SELDI-TOF MS has allowed us to generate sputum protein profiles of several diseases. We have shown there are substantial numbers of potential protein biomarkers that differentiate patients with inflammatory airway diseases from healthy subjects. We have proceeded to identify several of these candidate biomarkers and, using the example of calgranulins A and B, have demonstrated clinical relevance with longitudinal evaluation during altered disease activity. We have demonstrated that, having identified a biomarker by primary screening rather than empirical preselection, a standard assay, ELISA, can then be applied to the same clinical samples and deliver quantitative measures of the biomarker. Thus, screening with high-technology SELDI-TOF proteomics can be converted into clinically applicable measures.

Because this is the first study of its kind to investigate the proteomics of induced sputum using SELDI-TOF, we sought to determine the reproducibility of this assay. The average intraassay CV was 22.41% and average interassay CV was 16.25%. This comes well published to display for serum (intraassay CV of 15.6%, interassay variation of 24.4% [19], urine (intraassay CV, 8–30%) (32), and saliva (intraassay CV, 18%, and interassay CV, 31%) (33).

Previous studies have demonstrated the utility of SELDI-TOF in the diagnosis of illnesses in a range of body fluids. We used the fluid phase of sputum as a noninvasive means of assessing the airways. Although previous work has suggested a correlation of induced sputum with bronchial washings but not BAL (34), and therefore suggests that it reflects the pathophysiology of more central airways, induced sputum has been demonstrated useful in inflammatory respiratory diseases (11, 12, 35–46). Therefore, we would conclude that the proteomic assessment of induced sputum is a valid means of assessing airway disease.

As described previously, SELDI-TOF technology has been used as a technique in a number of inflammatory and neoplastic diseases, mainly using serum as a sampling medium. It was suggested as a diagnostic platform using bioinformatic algorithms to separate subjects with ovarian cancer from healthy control subjects (15). This initial approach, however, was widely criticized in the literature because the results were not reproducible when the same data sets were examined by independent investigators (21). Baggett and colleagues (21) drew particular notice to the impact of different modes of data preparation after acquisition, such as baseline subtraction in some groups of data and not in others. They also suggested that inaccuracies in sample collection protocol and mass calibration could lead to misinformed results in these sorts of experiments. To ensure uniformity in this study, all data in our analysis underwent the same steps of preparation before analysis. We also ensured that all sample preparation was uniform as outlined in Methods and below. Mass calibration was performed before each experiment using protein and peptide standards (Ciphergen All in One protein and peptide standards; Ciphergen). Furthermore, the widely criticized Petricoin study also used mass spectral data at less than 1,000 D, a mass range at

TABLE 3. PROTEIN IDENTIFICATIONS CONFIRMED FROM PRESENT STUDY

<table>
<thead>
<tr>
<th>Protein</th>
<th>Molecular Weight (D)</th>
<th>Accession Number</th>
<th>Corresponding SELDI Peak (D)</th>
<th>Confirmed by PMF</th>
<th>Confirmed by MS/MS</th>
<th>Confirmed By Antibody</th>
<th>Direction of Change in Suppuration</th>
</tr>
</thead>
<tbody>
<tr>
<td>Calgranulin A</td>
<td>10,834</td>
<td>P05109</td>
<td>10,834, 10,596</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Increased</td>
</tr>
<tr>
<td>Calgranulin B</td>
<td>12,960</td>
<td>P06702</td>
<td>12,960, 13,200</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Increased</td>
</tr>
<tr>
<td>Calgranulin C</td>
<td>10,100</td>
<td>P80511</td>
<td>10,100</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
<td>Increased</td>
</tr>
<tr>
<td>Clara cell secretory protein</td>
<td>7,900</td>
<td>P11684</td>
<td>7,900</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Decreased</td>
</tr>
<tr>
<td>Proline rich salivary peptide</td>
<td>8,188</td>
<td>P02814</td>
<td>8,119</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>Decreased</td>
</tr>
<tr>
<td>Lysozyme C precursor</td>
<td>16,537</td>
<td>P61626</td>
<td>14,600</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
<td>No Change</td>
</tr>
<tr>
<td>Cystatin s</td>
<td>16,204</td>
<td>P01036</td>
<td>16,079</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No Change</td>
</tr>
<tr>
<td>Hemoglobin alpha</td>
<td>15,117</td>
<td>P69905</td>
<td>15,080</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No Change</td>
</tr>
</tbody>
</table>

Definition of abbreviations: PMF = peptide mass fingerprinting; SELDI = surface-enhanced laser desorption/ionization.

Proteins were identified by trypsin digest and PMF and well as tandem MS/MS. When available, antibodies were used to confirm protein identification by Western blot. Molecular weight refers to the theoretical molecular weight of each protein as derived from sequence. Corresponding SELDI peak refers to the protein peak seen on SELDI analysis of sputum fluid phase (differences in molecular weight may represent post-translational modifications).
which the discrimination of genuine protein peaks from background noise may be difficult. Our approach of excluding data from below 4,000 D from analysis further strengthens the findings of this study. Other studies using SELDI-TOF as a diagnostic platform in infectious diseases using more robust sample collection and data preparation algorithms have proved more promising (19, 20). However, in this study, we used SELDI-TOF as a high-throughput screening platform for biomarker discovery rather than as a primary diagnostic assay. Although, on initial observation, our data demonstrate a large number of differentiating peaks when we look at the combined data set, the actual clinical significance of this observation is less clear and should be interpreted with caution.

Previous work by our group suggested an optimum mass range for respiratory biomarker discovery with this platform to be 3 to 20 kD (25). This may lead to biomarkers of a higher mass being ignored. Other proteomics methods, such as two-dimensional gel electrophoresis and immunocapture, may be better in assessing larger molecular weight proteins, and has been applied to CF (27, 29).

The largest numbers of differentiating proteins between disease and control were found in CF and bronchiectasis. This may suggest greater numbers of inflammatory and disease-related proteins in these conditions. Further studies are needed to confirm these findings and to determine the clinical relevance of these proteins.

Figure 3. (A) The protein that gave the statistically greatest differentiation between asthma and chronic obstructive pulmonary disease (COPD) (P < 0.001) had a MW of 29 kD, and was detected on the IMAC nickel (IMACNi) surface. The individual data and the median bars are shown. The data are for 22 patients with asthma and 23 patients with COPD (see explanation in footnote to Table 2). Other disease groups are demonstrated for comparison. (B) The protein that gave the statistically greatest differentiation between cystic fibrosis (CF) and bronchiectasis (B’lectasis) (P < 0.001) had an MW of 12.246 kD, and was detected on the IMACNi surface. The individual data and the median bars are shown. The data are for 28 patients with CF and 19 patients with bronchiectasis (see explanation in footnote to Table 2). Other disease groups are demonstrated for comparison.

Figure 4. Calgranulin A in all disease groups: (A) shows the relative expression, on surface-enhanced laser desorption/ionization time-of-flight (SELDI-TOF), of the 10,831-D protein, which was elevated (P < 0.001) in all disease groups versus control; (B) shows the expression of the 10,576-D form of calgranulin A, which was specifically elevated in patients with cystic fibrosis (CF) and bronchiectasis (B’lectasis) (P < 0.001 vs. all other groups) (the data in A and B were obtained using the IMAC Nickel surface); (C) is a Western blot of calgranulin A. The blot represents the overall abundance of calgranulin A from both 10,834- and 10,576-D forms in the sample groups (lanes are labeled as Bl = blank, A = asthma; Br = bronchiectasis; CF = cystic fibrosis; Con = control; CO = COPD). Note the strong staining for CF and bronchiectasis groups and very weak staining in asthma. (D) demonstrates the effect of bacterial colonization on the signal intensity of the 10,576-D form of calgranulin A; the majority of patients were colonized with Pseudomonas aeruginosa (shown in the first column labeled “Pseudomonas’’); other organisms included Staphylococcus aureus, Haemophilus, Stenotrophomonas, and Burkholderia (labeled “Other’’).
related proteins in these groups or, alternatively, samples obtained from these groups had greater abundance of protein overall. Sputum samples obtained from subjects with CF and bronchiectatic subjects were of greater volume and better quality subjectively. However, selected mucus plugs were used for analysis and processed with equal ratios of buffer as previously described (46). Moreover, to ensure samples were comparable, supernatants were adjusted to a total protein concentration of 1 mg/ml before applying to chip surfaces. Also, mass spectral data were normalized to total ion current before data analysis, thus minimizing effects of possible variation in protein binding to chip surfaces. After normalization, samples were excluded because they had absent or very poor signals. We would therefore suggest that the differences expressed between groups represent real differences in protein expression rather than confounding due to issues of sample quality and abundance.

Our data revealed large numbers of potential biomarkers. The preset parameters we created may have limited the possible number. A lower limit cutoff of 4 kD may have been too harsh. Indeed, our spectra showed the human α defensins in all groups, but these were excluded from analysis because their average mass was less than 4 kD. Alternatively, we may have overestimated true numbers of differentiating markers because some proteins were detected on more than one chip surface. Also, a proportion of some proteins may be doubly protonated. Because the “time of flight” is related to both MW and charge, such proteins will appear “twice,” once with half the MW of the original protein. Furthermore, different peaks may represent cleavage products of the same (higher molecular weight) proteins and this may in part explain the high numbers of potential differentiating peaks we recorded, particularly when comparing our CF and bronchiectasis groups with control subjects.

Despite finding a large number of peaks differentiating CF and bronchiectasis from controls, we were surprised not to have identified any of these as known biomarkers of inflammation, such as IL-8 or neutrophil elastase. In our study, SELDI-TOF was most efficient at demonstrating peaks in the 5–20 kD range and this may in part explain these findings as the predicted molecular weights of neutrophil elastase is 29.5 kD (47). Furthermore, the chip surface chemistry and binding conditions may simply not have favored preferential selection of IL-8.

Despite the limitations in this technique, we found large numbers of potential biomarkers from which we have so far identified a modest number, although further identification of other peaks is ongoing. We used one of these biomarkers, calprotectin (calgranulin A and B), to show that it was possible to monitor disease activity and to apply an ELISA to the same clinical samples, obtaining the same results as with MS.

In all disease groups, we identified calgranulin A at higher levels than in control subjects. This S100 protein is produced by neutrophils, macrophages, and epithelial cells. In the CF and bronchiectasis groups, a protein at 10.574 kD was also identified as calgranulin A. Theoretical removal of glutamic acid and lysine from the N terminus of calgranulin A would result in an identical mass shift from 10.831 to 10.574 kD. Calgranulin 10.574 may therefore be the product of specific cleavage or post-translational modification in CF and bronchiectasis. Whether this finding may be of functional significance is as yet unclear. Calgranulins A and B have also been described recently in BAL fluid from subjects with CF (25, 26); a similar finding of a separate peak representing calgranulin A at a lower molecular weight was also described in BAL fluid by McMorran and colleagues (26).

Calgranulin has previously been described in the sera of subjects who were homo- and heterozygous for CF mutations and was referred to as CF antigen (48). Calgranulin A may
provide a robust and sensitive marker of airway inflammation in CF. The likely source of calgranulin A is from neutrophils in the airway, although in this study we failed to demonstrate any significant correlation between neutrophil % in sputum and calgranulin A as measured by SELDI-TOF. This may in part be explained by the finding that the majority of the patients with CF had a profound neutrophilia in sputum in excess of 95%, regardless of other clinical details. Furthermore, there was a weak relationship between decreasing FEV1 and higher levels of calgranulin A, implying that calgranulin A may be a marker of disease severity in CF. We replicated our SELDI-TOF results using an ELISA to calprotectin, the heterodimer of calgranulins A and B. Thus, a more practical assay can be applied to induced sputum and yield the same clinical information. Calprotectin has previously been recognized as a marker in other inflammatory disorders, including inflammatory bowel disease and arthritis (49–51). Calgranulin C, another S100 protein, was also identified in our samples. This is in keeping with previous work demonstrating its presence in sputum and its potential usefulness as a biomarker in serum for CF (52).

A second abundant protein identified showed opposite effects to calgranulin A. CCSP was reduced in all disease groups compared with controls. CCSP is an antiinflammatory protein mainly expressed in the epithelial cells of the airways. Low serum levels have been reported in patients with asthma (53), and low nasal lavage levels in patients with allergic rhinitis (54). CCSP has a number of possible activities, including inhibition of phospholipase A2, chelation of calcium, and down-regulation of IFN-γ, IL-1, and tumor necrosis factor-α. Our findings are consistent with these previous studies, and raise the possibility that resolution of inflammatory processes might be monitored by rising levels of CCSP.

In conclusion, we have demonstrated the utility of SELDI-TOF MS as a tool for biomarker discovery in induced sputum. We have positively identified proteins of biological significance in the fluid phase of sputum. These proteins may form the basis of clinical point assays to assess the presence and activity of inflammation in a variety of lung diseases, with a particular emphasis being placed on disease monitoring.

**Conflict of Interest Statement:** None of the authors has a financial relationship with a commercial entity that has an interest in the subject of this manuscript.

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**References**


