MMP-2 and TIMP-1 predict healing of WTC-lung injury in New York City firefighters

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Abstract

Rationale: After 9/11/2001, most FDNY workers had persistent lung function decline but some exposed workers recovered. We hypothesized that the protease/anti-protease balance in serum soon after exposure predicts subsequent recovery.

Methods: We performed a nested case–control study measuring biomarkers in serum drawn before 3/2002 and subsequent forced expiratory volume at one second (FEV1) on repeat spirometry before 3/2008. Serum was assayed for matrix metalloproteinases (MMP-1,2,3,7,8,9,12 and 13) and tissue inhibitors of metalloproteinases (TIMP-1,2,3,4). The representative sub-cohort defined analyte distribution and a concentration above 75th percentile defined elevated biomarker expression. An FEV1 one standard deviation above the mean defined resistance to airway injury. Logistic regression was adjusted for pre-9/11 FEV1, BMI, age and exposure intensity modeled the association between elevated biomarker expression and above average FEV1.

Results: FEV1 in cases and controls declined 10% of after 9/11/2001. Cases subsequently returned to 99% of their pre-exposure FEV1 while decline persisted in controls. Elevated TIMP-1 and MMP-2 increased the odds of resistance by 5.4 and 4.2 fold while elevated MMP-1 decreased it by 0.27 fold.

Conclusions: Resistant cases displayed healing, returning to 99% of pre-exposure values. High TIMP-1 and MMP-2 predict healing. MMP/TIMP balance reflects independent pathways to airway injury and repair after WTC exposure.

Keywords: Biomarkers, Lung disease, Occupational exposure

Introduction

The collapse of the World Trade Center (WTC) on 9/11/2001 produced a massive exposure to dust and products of combustion [1-3]. The Fire Department of New York (FDNY) bureau of health services rapidly responded to this atrocity, initiating a medical monitoring program in October of 2001. Over 13,000 exposed rescue workers have been longitudinally followed by the FDNY-WTC-Medical Monitoring and Treatment Program. Approximately 7,000 exposed workers had serum samples drawn, stored and FEV1 and FVC measured within six months of 9/11/2001. A vast majority of those exposed had an acute decline in lung function in the first six months followed by stabilization. There was no recovery in lung function for the group as a whole, but a minority of those exposed recovered lung function over the following six and a half years.

We used serum obtained soon after the exposure to measure serum biomarkers of lung injury during the process of disease evolution. We reported that inflammatory cytokines, lipids and other measure of metabolic syndrome as well as biomarkers of cardiovascular risk predict abnormal lung function years later [4-6].

The balance of increased protease activity and reduced anti-protease activity are components of many diseases including cigarette-induced chronic lung disease and other causes of accelerated lung function decline [7-10]. Genetic association studies with matrix metalloproteinases (MMPs) demonstrate a strong association with the development of lung disease [7]. MMP-1 is induced in smokers with COPD and its overexpression in mice causes emphysema [11,12].

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The destructive effects of MMPs are inhibited by tissue inhibitors of matrix metalloproteinases (TIMPs). Since most clinical investigation focuses on disease, there is little data on the role of MMPs and TIMPs in the resistance to the damaging effect of dust exposure [13-16]. One carefully done pathologic study demonstrated increased MMP-2 and TIMP-1 mRNA expression in surgically removed lung and predicted improved FEV1 in COPD patients [17]. Serum MMP and TIMP expression reflects the severity of COPD supporting an investigation of a link between serum MMP/TIMP balance and lung function in the WTC exposed cohort [18].

This study investigates serum expression of MMPs and TIMPs soon after damaging particulate matter exposure and tests if protease/anti-protease balance is associated with a subgroup that demonstrates an above average healing potential after WTC-LI. We report that elevated TIMP-1 and MMP-2 predicts recovery of lung function while elevated MMP-1 reduces the odds of recovery years after WTC exposure.

Methods

Study participants and design

The Institutional Review Boards (IRB) of NYU and Montefiore Medical Center approved this study and patients signed consent at the time of serum draw within 6 months of 9/11/2001 (Montefiore Medical Center IRB; #07-09-320 and New York University IRB; #11-00439). The parent cohort for this investigation consists of 1,720 exposed workers who required subspecialty pulmonary evaluation SPE, when spirometry was performed and serum drawn, was 2 months in cases and controls. Time to subspecialty pulmonary evaluation SPE, when spirometry that defined case status was performed, was 32 months in cases and controls. Time to medical monitoring entry (MME), when spirometry was performed and serum drawn, was 2 months in cases and controls. Time to subspecialty pulmonary examination (SPE), when spirometry that defined case status was performed, was 32 months in cases and controls.

A nested case–control study tested the association of serum biomarkers and FEV1 at SPE. The baseline cohort N = 801 was assembled to exclude patients with abnormal pre-9/11 FEV1 and tobacco use to eliminate these confounders of post exposure lung function [19]. Resistant cases (N = 100/801) had a FEV1% one standard deviation above the mean (>107%) at SPE. Controls (N = 171/801) were randomly selected after stratification of the baseline cohort for BMI and FEV1. Serum was available for N = 137/171 of the cohort controls, N = 77/100 of the resistant cases. For this case–control study, the control subjects are all individuals in the random sample cohort control who did not meet criteria to be resistant cases. Analyte cutoff values were the 75th percentile cohort control expression. Analyte cutoff values were the 75th percentile cohort control expression. The Hosmer-Lemeshow goodness-of-fit test was used to assess calibration of the final model. The model discrimination was evaluated using the receiver operating characteristic area under the curve (AUC).

Serum biomarker assays

Processing of blood has been previously described [6,20]. Biomarkers were assayed with a TIMP panel (R&D Systems, Minneapolis, MN), and MMP panel (Procarta/Affymetrix) using a Luminex 200-IS (Luminex Corporation, Austin, TX). Each plate contained 1:2 ratio of resistant cases to controls to account for batch effect.

Chest computed tomography (CT)

We have reported on lung function and computed tomography (CT) findings of this group. High-resolution chest CT scans were obtained with 7-mm thick mages reconstructed at 6-mm intervals. Images were read for airway and parenchymal abnormalities by dedicated, board certified radiologists who had no knowledge of the subjects’ exposure status or clinical findings. Bronchial wall thickening and air-trapping were interpreted qualitatively and analyzed as a yes/no variable.

Statistical analysis

Database management and statistics used SPSS 20 (IBM, Armonk, NY). Odd ratios were modeled with multivariate binary logistic regression with case status as the outcome. Analyte cutoff values were the 75th percentile cohort control expression. The Hosmer-Lemeshow goodness-of-fit test was used to assess calibration of the final model. The model discrimination was evaluated using the receiver operating characteristic area under the curve (AUC).

Results

Participants

Derivation of cases and controls is depicted in Figure 1. N = 111/171 cohort controls with serum available for study were compared to N = 77/100 resistant cases. Controls were not significantly different from the baseline cohort in arrival time, years of service, age, time to all pulmonary examinations, or BMI, Table 1. Controls had higher BMI than resistant cases. Time to medical monitoring entry (MME), when spirometry was performed and serum drawn, was 2 months in cases and controls. Time to subspecialty pulmonary evaluation SPE, when spirometry that defined case status was performed, was 32 months in cases and controls.

Longitudinal lung function in resistant cases and controls

Cases and controls underwent three longitudinal measures of lung function. The first spirometry documented pre-exposure lung function; the second was immediately after exposure at MME and the third was later at SPE, Table 2 and Figure 2. Resistant cases had higher median pre-9/11 FEV1 than controls (117% vs 98% p < 0.001).

FEV1 declined in cases and controls soon after exposure. To test if group data represented individuals’ response the ratio of FEV1 pre-9/11 to FEV1 at MME was calculated for each case and controls. The mean (SD) of MME/Pre-911 ratio was 0.91(0.10) for resistant cases and 0.91 (0.12) for controls (p = 0.95 for cases vs control). Cases recovered most of the lost FEV1 (117% to 113%). Controls had less improvement than cases in their median FEV1 (98% to 93%). To test if group data represented individuals’ response the ratio of FEV1 pre-9/11 to FEV1 at SPE was calculated for each case and controls. The SPE/Pre-911 ratio...
was 0.99+/−0.09 for resistant cases and 0.91+/−0.12 for controls (p < 0.001 for cases vs control). Raw FEV₁% predicted has also been shown in Figure 2.

Both groups had a decline in FEV₁/FVC ratio SPE/MME, with resistant cases having the less than controls (0.85 to 0.81 p < 0.001). Both cases and controls had a high proportion of individuals with bronchodilator response (22% vs 28% NS). Cases had less methacholine reactivity when measured as a continuous slope (0.64 vs 0.04 p < 0.01); 9% of cases reactive to a 10 mg/ml dose while 20% of controls were reactive (p = 0.11). Cases had less air trapping than controls (111% vs 121% p < 0.02) when residual volume was used to measure air trapping Table 3. At SPE, resistant cases had the higher TLC, DLCO and alveolar volume than controls (p < 0.001 for all comparisons), Table 2.

Chest CT of resistant cases and controls
Chest imaging was used to investigate if resistant cases and controls had structural differences. In those with chest CT images available, 14% of the resistant cases had bronchial wall thickening whereas 35% of the controls had this evidence of airway inflammation (p < 0.03), Table 3. There was no significant difference between cases and controls in air trapping defined by mosaic attenuation (28% vs 46% p = 0.09). Both cases and controls have a high proportion of pulmonary nodules (38% vs 36% p = 0.8).

Biomarker models
Analyte levels were compared between controls and resistant cases, Table 4. We used logistic regression with analyte expression above a pre-defined 75th percentile cut points

**Table 1 Demographics**

| Date/event          | Baseline cohort | Sub-cohort controls | Resistant cases | p    |
|---------------------|-----------------|---------------------|-----------------|------|
| WTC Exposure, n(%)  | High            | 197 (25)            | 21 (19)         | 14 (18) | 1.000 |
|                     | Intermediate    | 604 (75)            | 90 (81)         | 63 (82) | 0.755 |
| 9/11 to PFT, months | MME             | 2.7 (2.0-3.8)       | 2.5 (2.0-3.2)   | 2.6 (2.2-3.2) | 0.219 |
|                     | SPE             | 33.8 (24.8-57.0)    | 32.7 (24.6-57.4) | 32.5 (28.3-52.2) | 0.755 |
| BMI, kg/m²          | MME             | 28.0 (26-30)        | 28.1 (26-31)    | 27.3 (26-29) | 0.018 |
|                     | SPE             | 28.9 (27-31)        | 29.1 (27-31)    | 27.6 (26-30) | 0.010 |
| Years of service    | 9/11/01         | 13 (7-19)           | 14 (7-18)       | 14 (10-19) | 0.249 |
| Age                 | 9/11/01         | 40 (36-45)          | 41 (36-44)      | 42 (38-46) | 0.159 |

WTC World trade center, PFT Pulmonary function test, BMI Body mass index, MME Medical monitoring entry, SPE Subspecialty pulmonary exam. Expressed as median (interquartile range.)
to test if protease or anti-proteases expression predicted resistance to WTC-LI [21]. Models were adjusted for BMI, age, exposure group and pre-9/11 FEV1. Reduced models examined the ability of a single analyte to predict each case definition. Analytes with significant odds ratios in single biomarker models were used to develop the final multi-analyte model.

Using 75th percentile of cohort expression, we calculated OR with FEV1 > 107% at SPE as the outcome. Elevated MMP-1 reduces the odds of FEV1 > 107% by 31%, elevated MMP-2 increases the odds of FEV1 > 107% by 218% and elevated TIMP-1 increases the odds by 231%. After adjusting for BMI, age, exposure group and pre-9/11 FEV1 elevated MMP-1 reduces the odds of resistance to WTC-LI by 68% while individuals with high MMP-2 and TIMP-1 were 300% and 350% more likely to resist WTC-LI, Table 4. Combining MMP-1, MMP-2 and TIMP-1 in a multi-analyte model improved the OR for each of these biomarkers, Table 5. Elevated MMP-1 was a risk factor reducing the odds of resistance to lung injury to 0.27 (95% CI 0.09-0.82 p = 0.02). Elevated MMP-2 and TIMP-1 were protective factors improving odds of resistance by 4.2 fold (95% CI 1.6-10.8 p = 0.003) and 5.4 fold (95% CI 1.9-14.9 p = 0.001), Table 5. The area under the receiver operator curve for the multi-analyte model was 0.90 (95% CI; 0.86-0.94). (Figure 3) The sensitivity and specificity of the model was 74% and 86% respectively.

**Discussion**

We report that elevated MMP-1, MMP-2 and TIMP-1 in serum shortly after WTC exposure predicts subsequent return of FEV1 to pre-exposure values. All subjects in this nested case control investigation had significant WTC dust exposure, arriving at the collapse site within 2 days of 9/11/2001. The FDNY measured serial FEV1 pre and post 9/11. We focused our current study on a subgroup of highly exposed individuals who did not suffer persistent

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**Table 2 Longitudinal lung function assessment of cohort**

| Time     | Variable          | Controls    | Resistant | p   |
|----------|-------------------|-------------|-----------|-----|
| Pre-9/11 | FEV1%             | 98 (90–108) | 117(108–124) | <0.0001 |
|          | FEV1/FVC          | 84 (81–87)  | 87 (84–89) | <0.0001 |
| MME      | FEV1%             | 91 (81–96)  | 105(97–113) | <0.0001 |
|          | FEV1/FVC          | 84 (79–87)  | 85 (83–89) | <0.0001 |
| SPE      | FEV1%             | 93 (83–98)  | 113 (109–119) | <0.0001 |
|          | FEV1/FVC          | 77 (73–79)  | 81 (79–84) | <0.0001 |
| BD response |                  | 5 (2–10)   | 5(2–9)    | 0.776  |
| BD response > 12% N (%) |           | 13/46 (28)  | 5/23 (22) | 0.772  |
| MCT slope |                  | 0.64 (0.31-0.15) | 0.04(0.02-0.07) | 0.010  |
| PC20 < 10 mg/mL MCT, N (%) |           | 17/86 (20)  | 6/66 (9)  | 0.108  |
| TLC      |                  | 103 (95–109) | 106 (103–116) | 0.016  |
| RV/% Pred|                  | 125 (113–141) | 111 (103–123) | 0.015  |
| DLCO     |                  | 106 (99–114) | 116 (108–131) | 0.001  |
| VA       |                  | 93 (85–98)  | 100 (95–105) | <0.0001 |
| DLCO/VA  |                  | 120 (107–130) | 110 (107–126) | 0.358  |

**FEV1** Forced expiratory volume in one second, **FVC** Forced vital capacity, **MCT** Methacholine, **BD** Bronchodilator, **PC20** concentration of methacholine causing a 20% fall in FEV1, **DLCO** Diffusion capacity of the lung for carbon monoxide, **VA** Alveolar volume.

**Figure 2** FEV1% predicted of cases and controls over time. Mean (SD) expressed for cases n = 77 (green) and controls n = 111 (grey) at Pre-9/11, MME and SPE.

**Table 3 Chest CT abnormalities in cases and controls**

| CT Finding            | Controls | Resistant cases | p† |
|-----------------------|----------|-----------------|----|
| Bronchial wall thickening | 23/65 (35) | 5/36 (14) | 0.022 |
| Air trapping          | 30/65 (46) | 10/36 (28) | 0.090 |
| Nodules               | 25/65 (38) | 13/36 (36) | 0.834 |

†Significance calculated by Chi-Squared.
FEV₁ decline. This resistant subgroup had greater than average reduction in FEV₁ immediately after exposure but returned to pre-exposure FEV₁ over the next 6.5 years. Because serum was drawn well before the pulmonary function test that demonstrated recovery, the biomarker information reflected the evolving response to injury. MMP-2 and TIMP-1 expression above the 75th percentile are protective biomarkers, significantly increasing the odds of resistance between 4.2 and 5.4 fold. Alternately, elevated MMP-1 is a risk factor, reducing the odds of resistance by 73%. The biomarker model using serum MMP-1, MMP-2 and TIMP-1 concentration predicted resistance with a sensitivity of 74%, a specificity of 86% and a receiver operator characteristic of 0.90.

As expected in this highly exposed group, both cases and controls suffered an acute reduction in FEV₁ as a result of WTC exposure. Resistant cases differed from controls because they returned to 99% of their pre-exposure FEV₁. Over the 6.5 years post 9/11 FEV₁ returned to only 91% of their pre-exposure FEV₁. The return of FEV₁ to pre-exposure levels provides evidence that resistant cases have above average capacity to heal after an acute injury.

Table 4 Serum biomarkers

| Analyte | 75th percentile cutpoint | Controls | Resistant | OR (95% CI) |
|---------|--------------------------|----------|-----------|-------------|
|         | pg/mL                    | Median (IQR) | Proportion | Median (IQR) | Proportion |
| Proteases |                         |           |           |             |
| MMP-1   | 1239                     | 689 (264–1210) | 27/111 | 549 (233–1013) | 14/77 | 0.691 (0.335-1.425) |
| MMP-2   | 4949                     | 2933 (1570-4608.25) | 22/111 | 3340 (2108.19-5682.13) | 27/77 | 2.185 (1.128-4.231) |
| MMP-3   | 13345                    | 7494 (2940–13358) | 28/111 | 6906 (3457–11509) | 17/77 | 0.840 (0.422-1.671) |
| MMP-7   | 355                      | 217 (100–333) | 25/111 | 237 (95–375) | 22/77 | 1.376 (0.707-2.77) |
| MMP-8   | 157                      | 10 (10–156) | 27/111 | 10 (10–116) | 16/77 | 0.816 (0.405-1.645) |
| MMP-9   | 62086                    | 22559 (10596–40806) | 24/111 | 23003 (12043–64398) | 21/77 | 1.359 (0.692-2.670) |
| MMP-12  | 306                      | 66 (18–306) | 28/111 | 55 (27–277) | 19/77 | 0.971 (0.496-1.902) |
| MMP-13  | 138                      | 75 (10–133) | 26/111 | 71 (10–140) | 20/77 | 1.147 (0.585-2.248) |
| Anti-proteases |               |           |           |             |
| TIMP-1  | 155003                   | 125491 (100396–151151) | 23/111 | 126292 (100483–164661) | 29/77 | 2.312 (1.206-4.430) |
| TIMP-2  | 115905                   | 102470 (89901–113738) | 23/111 | 103199 (88959–119820) | 22/77 | 1.530 (0.780-3.005) |
| TIMP-3  | 31922                    | 7750 (7750–31922) | 29/111 | 7750 (7750–33680) | 21/77 | 1.060 (0.550-2.044) |
| TIMP-4  | 1585                     | 1242 (9288-1562) | 26/111 | 1342 (970-1717) | 27/77 | 1.765 (0.929-3.354) |

MMP matrix metalloproteinase, TIMP Tissue inhibitor of matrix metalloproteinase, OR Odds RATIO (unadjusted), IQR Interquartile range.

Elevated TIMP-1 and MMP-2 expression increases the odds of being resistant 4.1 and 5.3 fold respectively. These results remain significant after adjusting for multiple comparisons. Although TIMP and MMP over expression have been observed in COPD and are probably affected by chronic injury secondary to cigarette smoke or other damaging processes, interpreting the cause and effect relationship between lung function and MMP/TIMP balance in humans has been challenging [22-24]. A carefully done study of 63 patients who had surgery for lung cancer or lung transplantation demonstrated that increasing TIMP-1 and MMP-2 mRNA in the small airways and/or

Table 5 Model of resistance to WTC-lung injury

| Serum biomarker | 75th percentile cutpoint | Adjusted§ |
|-----------------|--------------------------|-----------|
| Single analyte  |                          |           |
| MMP-1 ≥ 1239    | 0.33 (0.11-0.93)         |           |
| MMP-2 ≥ 4949    | 3.00 (1.25-7.18)         |           |
| TIMP-1 ≥ 155003 | 3.52 (1.41-8.81)         |           |
| Multi-analyte   |                          |           |
| MMP-1 ≥ 1239    | 0.27 (0.09-0.82)         |           |
| MMP-2 ≥ 4949    | 4.16 (1.61-10.76)        |           |
| TIMP-1 ≥ 155003 | 5.38 (1.94-14.94)        |           |

§Models Adjusted for: BMI at SPE, exposure Group, Age at 9/11 and Pre-9/11 FEV₁% Predicted; 2 Log Likelihood 144.081; Hosmer and Lemeshow (Goodness of Fit) Sig 0.743.

Figure 3 Receiver operator characteristics (ROC) of final multi-analyte model. Area Under the Curve (AUC) = 0.901; 95% CI (0.860-0.943).
the parenchyma surrounding the small airway was positively associated with FEV$_1$ [17]. Our findings on particulate matter induced lung injury are consistent with these findings in smoking related COPD patients. The mechanisms involved in the protective effects of TIMP-1 and MMP-2 in response to particulate matter induced lung injury require further study.

Since resistant cases were defined by a FEV$_1$ of > 107%, starting close to this threshold should increase the likelihood of crossing this defining value. After adjusting the logistic model for pre-9/11 FEV$_1$, the association of MMP-1 MMP-2 and TIMP-1 with better than average post exposure FEV$_1$ markedly improved as quantified by their ORs. This suggests that while pre-existing differences in lung function is a confounder including them in the regression allowed the biomarker-lung function association to remain robust. We briefly examined if excluding individuals who failed their MCT by the 3rd level would alter our results. We found that the ORs of MMP-1, MMP-2 and TIMP-1 remained significant and of the same magnitude (data not shown). Furthermore, we repeated the analysis in a population with their Pre-9/11 FEV$_1$% predicted constrained; cases n = 58 had an FEV$_1$ of 96–146 and controls n = 58 had an FEV$_1$ of 96–143. Using this constrained population OR (95% CI) were as follows: MMP-1 0.318 (0.098-1.039), MMP-2 4.993 (1.664-14.983) and TIMP-1 8.061 (2.423-26.825). Importantly, patients with resistance to WTC-LI had at subspecialty pulmonary evaluation increased TLC measured on plethysmography, increased alveolar volume measured by methane dilution and increased diffusion of carbon monoxide than controls. In animal models, MMP-2 is required for normal lung development and failure to produce MMP-2 leads to emphysema and collagen deposition around the bronchioles [25-27]. Our data is consistent with the hypothesis that MMP-2 and TIMP-1 are biomarkers of an individual’s intrinsic capacity to heal after irritant induced lung injury.

The resistant cases also had significantly less bronchial wall thickening on chest CT. Accumulation of inflammatory cells in the bronchovascular bundle likely accounts for this radiographic finding [28]. Our Chest CT findings suggest that the resistant group accumulates fewer inflammatory cells around the bronchovascular bundle after dust exposure. Interestingly, in rodent models of lung injury, MMP-2 expression reduces bronchovascular inflammatory cells and enhances repair [29,30]. Since the chest CT was performed years after the insult, the mechanism that produced the bronchial wall thickening persists for years after the original exposure that precipitated the inflammation.

Conversely, elevated MMP-1 reduced the odds of being resistant to WTC-LI to 0.27. This association, however, becomes insignificant when adjusted for multiple comparisons. We have maintained MMP-1 in the model because of the overwhelming evidence that this protease is an important is disease pathogenesis. Over expression of MMP-1 leads to emphysema in animal models [31]. In humans, MMP-1 is expressed in type II pneumocytes of patients with COPD but not controls [11,12]. The 73% reduction in the odds of having above average FEV$_1$ is consistent with the damaging effects of this protease on lung integrity.

Since we excluded ever smokers and individuals with pre-9/11 lung disease, our results are not confounded by these common causes of FEV$_1$ decline that are unrelated to inhalation of WTC dust. Another advantage of this study is the SPE PFT was performed during the initial pulmonary evaluation prior to treatment initiation. Therefore our results are not confounded by treatment effect. In spite of our objective case definition, using FEV$_1$ as a single measure did produce misclassification of disease. The resistant group with FEV$_1$ > 107% at SPE did have individuals with evidence of airway injury. Up to 22% of the resistant cases had airway reactivity on PFT or radiographic evidence of bronchial wall thickening. This misclassification of disease should bias toward the null. In spite of this bias, we observed highly significant associations between biomarkers in serum drawn soon after exposure and above average lung function years later.

This nested case control study has several limitations. The cohort was highly unusual, suffering an acute overwhelming exposure to PM that overwhelmed normal protective mechanisms. The results therefore have limited generalizability. The findings require replication in independent particulate matter exposed cohorts. Even though the serum biomarkers were expressed years before the FEV$_1$ that defined resistance to lung injury, the results are correlations and do not imply causation. It is possible that exposure lead to an alteration in these biomarkers due to different mechanisms controlling lung injury. Alternately, pre-existing attenuation of these biomarkers may lead to differential healing. Further, investigation in model systems and longitudinally followed cohorts is required to better understand the role, if any, of MMP-1 and TIMP-1 in healing after PM induced lung injury. Finally, this study had no unexposed control group because the few unexposed workers were markedly different from the exposed group with poor health that prevented them from working at the WTC collapse site. This restricts our ability to assess the impact of WTC exposure to the observed biomarker disease relationship.

This report documents the serum biomarkers that predict better than average FEV$_1$ after massive dust exposure. This group had evidence of healing with return to pre-9/11 FEV$_1$ after a significant drop immediately post exposure. The processes initiated by WTC exposure impacted multiple distinct injury and repair pathways. One interpretation of the findings is that biomarkers of resistance reflect biological processes leading to healing after particulate matter induced injury.
Competing interests
The authors report no financial or competing interests.

Authors’ contributions
AN, SK and MDW participated in study conception and design. AN, SK, ALC, BN and MDW were responsible for data collection. AN, SJC and SK were responsible for data validation. AN, ALC and SK participated in data analysis. AN, SK undertook the statistical analysis. All authors participated in data interpretation, writing and revision of the report and approval of the final version. All authors read and approved the final manuscript.

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