Pilot-Scale Study Of Human Plasma Proteomics Identifies ApoE And IL33 As Markers In Atopic Asthma

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Background: The pathobiology of atopic asthma is complex and the symptoms similar to other respiratory diseases. As such, identification of biomarkers of atopic asthma is of prime importance for better diagnosis and control of the disease.

Objectives: We sought to study the changes in plasma proteome and cytokine-expression profile across healthy and atopic asthmatics for identifying biomarkers and exploring aberrant pathways for atopic asthma.

Methods: A pilot-scale study in humans was performed to identify differentially expressed proteins in blood plasma of healthy controls (n=5) and treatment-naïve atopic asthma patients (n=5) using quantitative label-free liquid chromatography–tandem mass spectrometry proteomics and ELISA.

Results: Mass spectrometry–based proteomic analysis revealed ApoE to be significantly downregulated in atopic asthmatics compared to healthy volunteers. Decreased expression of ApoE in atopic asthmatics was validated by immunoblotting (50.74% decrease). Comparison with atopic asthmatics and COPD patients showed that ApoE was decreased (36.33%) in atopic asthma compared to COPD. IL33 was significantly upregulated in atopic asthmatics compared to healthy subjects (3.84-fold).

Conclusion: ApoE was downregulated and IL33 upregulated in atopic asthma patients compared to healthy volunteers. These two proteins’ profiles were distinct in atopic asthma from healthy and COPD plasma samples. Differential expression of these proteins could serve as a probable candidate for a two-protein classifier–based prognostic biomarker of atopic asthma.

Keywords: atopic asthma, LC-MS/MS, apolipoprotein E, ApoE, interleukin 33, IL33, biomarkers

Introduction

Asthma is a chronic inflammatory disease of the airways which has recently been recognized as heterogeneous in nature. Other general features of this complex disease are reversible airflow obstruction, airway hyperresponsiveness, wheezing, shortness of breath, and chest tightness.1,2 International guidelines define asthma by the presence of reversible airflow obstruction on spirometry. Asthma is considered severe when high-dose inhaled corticosteroids and other additional controllers, such as long-acting inhaled β2 agonists, montelukast, and/or theophylline or prolonged oral corticosteroid therapy, fail to control asthma symptoms or exacerbation rates.3 Apart from stratification of asthma based on severity of the disease, it can also be broadly classified as atopic and nonatopic asthma.4 Atopic or allergic asthma is generally
triggered by extrinsic allergens, whereas specific external trigger factors are not usually associated with nonatopic asthma. The prevalence of atopic asthma was found to be much higher than nonatopic asthma in a population-based study. The complexity and heterogeneity of asthma has made its diagnosis and treatment difficult. Th2 and Th1 inflammatory response have both been associated with asthma pathogenesis. The Th1/Th2 paradigm of asthma further complicates the overall scenario. Besides this, the striking similarity of symptoms (such as cough, shortness of breath, airflow obstruction, and sputum production) between asthma and COPD may further mislead proper diagnosis. Therefore, asthma and COPD have emerged as major health concerns in both developed and developing countries. In such circumstances, recent asthma research has focused on discovering asthma biomarkers, pathways, and novel targets for drug discovery.

Biomarkers serve as dynamic tools for understanding pathogenesis and prognosis of a wide variety of diseases, including neurological diseases, cancers, cardiovascular diseases, and genetic, allergic, and immunological disorders. Biomarkers are measurable characteristics that act as indicators of biological and pathogenic processes and response to therapeutics. In closely related diseases like leiomyosarcoma and gastrointestinal stromal tumour, diagnostic biomarkers based on relative expression of two genes often yield better accuracy as biomarkers than expression profile of a single gene. Different omics approaches, including liquid chromatography–tandem mass spectrometry (LC-MS/MS)-based proteomics, are widely employed for identification of biomarkers. The Database of Allergy and Asthma Biomarkers, a manually curated repository, enlists differentially expressed genes/proteins in different allergic diseases like asthma and allergic rhinitis. These molecules can serve as potential candidates for allergy and asthma biomarkers.

The present study focuses on quantitative LC-MS/MS-based proteomic study of human plasma samples from healthy volunteers and adult atopic asthma patients from the Indian subcontinent to identify biomarkers. Cytokine profiling for IL33, the alarmin cytokine, and other Th2 cytokines like IL4, IL5 and IL13 was performed to assess the pattern of expression of these cytokines in atopic asthma. The outcome of high-throughput proteomic data finding of differential expression of ApoE was validated by immunoblotting. Cytokine profiling and Western blotting was also performed with plasma samples from COPD patients to study the expression patterns of our findings in symptomatically similar respiratory disease. The overall goal of this study was to identify probable biomarker candidates, explore pathways, obtain mechanistic insight, and find therapeutic targets for atopic asthma with a view to differentiate asthma from other overlapping airway diseases based on these identified biomarkers.

Methods

Sample Collection

Venous blood samples (10 mL) were collected from healthy subjects (n=5), atopic asthma patients (n=5), and COPD patients (n=5) of east Indian origin. Asthma was defined by the presence of history of such symptoms as wheeze, shortness of breath, chest tightness, and cough that varied over time and in intensity, together with the presence of reversible airflow obstruction on spirometry. COPD was diagnosed by the presence of cough with expectoration, breathlessness, exposure to risk factors (eg, tobacco smoke, biomass fuels, occupational dust) along with irreversible airflow obstruction on spirometry. Reversible airflow obstruction on spirometry was defined as the presence of airflow obstruction (FEV1/FVC <0.7) associated with an improvement in FEV1 by at least >200 mL and >12% 15–20 minutes after inhalation of 200 µg of salbutamol. Spirometry was done according to the American Thoracic Society–European Respiratory Society Task Force guidelines. Patient samples were obtained from the Department of Pulmonary Medicine, BR Singh Hospital, Kolkata, India with prior consent following all ethical issues. In order to work with human blood samples, Human Ethics Committee clearance (BIHEC/2014-15/4) from Bose Institute, Kolkata was also obtained. We included adult atopic asthma patients who were treatment-naïve. Total IgE levels, allergen-specific IgE profilling (for common ragweed, molds, and house dust mites) and spirometry (% predicted FEV1) for patients were conducted at BR Singh Hospital. Allergenicity of atopic asthma patients toward common pollen allergens like Helianthus annuus and Cocos nucifera were assessed by dot-blot assays. Total IgE counts and spirometry (% predicted FEV1) for healthy volunteers were evaluated at a local pathology laboratory.

Isolation Of Blood Plasma

Blood samples were collected in K2-EDTA tubes (BD vacutainers). Blood samples were centrifuged immediately after collection at 350 g for 15 minutes at 4°C for plasma separation. Plasma samples were further subjected to centrifugation at 15,000 rpm for 15 minutes at 4°C for lipid
removal. On centrifugation, the lipid scum generated on top was removed by gentle pipetting. Lipid-free plasma samples were then stored at −20°C until use.

Dot-Blot Assays

Dot blot is a method widely used to detect allergenicity of samples to specific allergens. Antigen (pollen allergen, namely *H. annuus* and *C. nucifera*) extracts (1 µg) were dotted on polyvinylidene difluoride (PVDF) membranes (GE Life Sciences). Membranes were then blocked with 0.5% BSA in TBS-T buffer for 3 hours at room temperature. Patients’ sera were then added in 1:5 ratios in TBS-T buffer and membrane pieces incubated overnight at 4°C. Unbound IgE was removed by washing the membrane twice. Membranes were then incubated with secondary antibody in 1:1,000 dilution (Mouse monoclonal antihuman IgE–AP conjugate; Sigma-Aldrich) at 4°C for 3 hours. Washing was performed thrice with TBS-T. Dots were developed with nitro-blue-tetrazolium-5-bromo-4-chloro-3’-indolylphosphate (Abcam). Dot-blot assays were performed for atopic asthma patients and just one healthy subject. They were not done for any other healthy subjects, since the IgE observed for healthy individuals was low and within normal range. The immunodot blot was performed by spotting 10 µL antigenic extract right at the center of the circular disks of PVDF membranes. In cases of positive IgE binding, the substrate reaction occurs in the antigenic spots and the colored spots quickly develop at the center of the membrane disks. The appearance of prominent colored spots at the center of the disks denotes sensitivity to allergens. No sensitivity to allergens is denoted by lack of spot development at the center of the disk. In turn, the entire disk turns purplish, owing to prolonged exposure to photosensitive substrate solution.

Depletion Of High-Abundance Proteins From Blood-Plasma Samples

Six high-abundance proteins (albumin, transferrin, IgG, IgA, haptoglobin, and antitrypsin) were removed from the collected human plasma samples (healthy subjects, n=5; atopic asthma patients, n=5; COPD patients, n=5) using an Agilent Multiple Affinity Removal System following the manufacturer’s protocol. A snapshot of unidimensional SDS-PAGE of depleted and undepleted plasma samples is shown in Figure S1.

Unidimensional Gel Electrophoresis And Sample Processing For LC-MS/MS Analysis

Each of the depleted plasma samples (50 µg) of healthy subjects (n=5) and atopic asthma patients (n=5) were fractionated by unidimensional 12% SDS-PAGE. The gel was stained with Coomassie brilliant blue. Pieces of equal size were excised from each lane of the gel and outsourced to C-CAMP for subsequent LC-MS/MS run.

LC-MS/MS

In-gel digestion was performed using trypsin in C-CAMP using the standard protocol. Trypsin-digested peptides were vacuum-dried and reconstituted with 0.1% formic acid and then vortexed for 1 hour, followed by centrifugation at 10,000 rpm for 5 minutes. The supernatant containing the digested peptide mix was then collected and separated using Agilent Nano 1200 high-performance liquid chromatography, present in C-CAMP. An Advion NanoMate Triversa (automated nano–electrospray ionization tool) was then used to generate ions for subsequent MS analysis. An LTQ Orbitrap Discovery was used for accurate determination of m/z ratios and data acquisition.

LC-MS/MS Data Analysis

Proteins were identified by the Mascot search engine after LC-MS/MS runs. The raw data files in .dat file format were uploaded to Scaffold software for protein quantification. Protein identification was accepted at a 5% false-discovery rate and contained at least two identified peptides. Total unique peptides and total spectral count for each protein across the different healthy and patient samples were obtained. Run-to-run variation was normalized using the normalized spectral abundance factor (NSAF). The NSAF of a protein (p) was calculated:

\[ \text{NSAF} = \frac{(\text{SpC}/\text{Length})_p}{\sum (\text{SpC}/\text{Length})} \]

where the spectral count (SpC) identified for protein p by the protein length (Length) is divided by the sum of SpC/Length for all proteins in the experiment to obtain the NSAF. Heat maps were constructed using Morpheus software (Broad Institute). Box plots were constructed using R packages for visual representation of the differential expression of target proteins across healthy and clinical samples.
Western Blot For Validation Of LC-MS/MS Results
Western blotting was performed to validate the outcome of LC-MS/MS using depleted blood-plasma samples from healthy subjects (n=5) and atopic asthma patients (n=5). Immunoblotting was also done with plasma samples from COPD patients (n=5). An equal amount of protein (150μg) was loaded on each lane followed by separation using SDS-PAGE and then transferred onto PVDF membrane (Merck Millipore). The membrane was blocked for 1 hour at 37°C with 5% BSA solution. Then, the membrane was incubated with anti-ApoE mouse monoclonal antibody (ab1907, 1:1,000; Abcam) overnight at 4°C, followed by incubation with horseradish peroxidase-conjugated antimouse secondary antibody (ab205719, 1:3,000; Abcam) for 1 hour. After washing, the membrane was developed using an electrochemiluminescent Western blotting detection buffer and detected with Chemidoc (GE Amersham Imager 600). β-Actin (1:5000, Abcam, ab8226) was used as a loading control. Band intensity was quantified using ImageJ 1.51j8.

Cytokine Profiling
Quantification of Th2 cytokines IL4, IL5, and IL13 in healthy subject (n=5) and atopic asthmatic (n=5) plasma samples was conducted using ELISA kits (RayBio) following the manufacturer’s protocol. Cytokine profiling of IL33 was performed thrice for freshly collected healthy volunteer (n=5), atopic asthmatic (n=5) and COPD (n=5) plasma samples with ELISA using a Quantikine ELISA kit for human IL33 (P118763 for healthy and asthma patient samples and P177042 for COPD samples; R&D systems) as per the manufacturer’s protocol. For each kit, a standard curve was generated separately and concentration of IL33 measured from the OD values at 450 nm using standard curve equations. OD values at 450 nm were obtained using an ELISA plate reader (Thermo Fisher Scientific).

Ingenuity Pathway Analysis Of Proteomic Data And Cytokine Profiling
Pathway analyses were done using ingenuity pathway analysis (IPA) after merging the different apolipoproteins from LC-MS/MS data and IL33 cytokine profiling. The other proteins identified by LC-MS/MS did not show significant variation in expression patterns and hence were not included in the IPA to reduce signal:noise ratio.

Statistical Analysis
Student’s t-test was used for experiments conducted, including LC-MS/MS, Western blot, and cytokine profiling.

Results
Baseline Study-Population Characteristics
Five atopic asthma patients, five COPD patients, and five healthy volunteers of east Indian origin were studied. Table 1 presents the baseline clinical characteristics of the asthmatics, COPD patients, and healthy volunteers from whom blood samples were collected. Of the five atopic asthma patients, two were male and three female. All asthma patients were sensitive to the two pollens — *H. annuus* and *C. nucifera* — as evident in Figure 1. The asthmatics were also allergic to other common environmental allergens like molds and house dust mites (Table 1 and Table S1). All five COPD patients were male. Neither patients nor the healthy cohort had any other major health complications as per individual health record.

Association Between Apolipoproteins And Atopic Asthma
Plasma levels of some apolipoproteins has been found to be altered in patient samples in comparison with healthy individuals. LC-MS/MS data obtained were analyzed by Scaffold version 4.8.7 using the XTagend search engine for protein quantification. The 167 different protein hits (Table S2) obtained from Scaffold were analyzed to study expression patterns. The proteins identified by LC-MS/MS can be classified into groups: blood-clotting factors, proteases, apolipoproteins, complement factors, and serum amyloid proteins. Since apolipoproteins have been linked with lung health and diseases, further study was focused on apolipoproteins.25,26 A total of 14 apolipoproteins, including ApoA1, ApoB, ApoE, ApoD, and ApoA4, were identified from LC-MS/MS across control and atopic asthma patient samples (Figure 2A and Table S3, A–C). ApoA4 was downregulated in atopic asthmatics (34.99% decrease, P<0.05), as shown in Figure 2B. Significant downregulated expression of ApoE (26.14% decrease, P<0.01) was also observed in atopic asthma patients (n=4) when compared with healthy controls (n=4) in terms of NSAF (Table S4). One healthy sample and one asthma patient sample in assessing the decreased expression of ApoE were ignored, because they appeared to be outliers in terms of computed NSAF values, as evident from the box plot in Figure 2C. As per the Human Protein...
Atlas, ApoA4 is not expressed in airway and lung tissue, but rather produced largely by the gastrointestinal tract. Since asthma primarily affects the airway tract, validation studies by Western blotting were not conducted for ApoA4. However, ApoE mRNA is expressed in lung tissue, so we validated the reduction in ApoE in allergic asthma patients by Western blotting (Figure S2).

Tables that appeared as outliers in LC-MS/MS analysis did not get reflected in low-throughput study performed with Western blot. Densitometric analysis of immunoblot data (Table S5) showed significant reduction in the expression of ApoE in atopic asthma patients (50.74% decrease, \( P<0.05 \); Figure 3, A and B). In COPD patients, ApoE showed reduction (22.63% decrease, \( P=0.12 \)). The reduced expression of ApoE in COPD patients was not as marked as in atopic asthma patients. The expression pattern of ApoE in atopic asthmatics was also significantly reduced (36.33% decrease, \( P<0.05 \)) compared to COPD patients.

Plasma IL33 Concentration Is Increased In Atopic Asthma Patients
IL33 concentration was significantly higher (3.84-fold, \( P<0.01 \)) in atopic asthma patients’ plasma samples in comparison to healthy individuals (Figure 4). For IL33, the minimum detectable dose of the kit was 0.519 pg/mL. The plasma-expression levels of IL33 in healthy subjects (\( n=5 \))
and atopic asthmatics (n=5) were 2.35±0.308 pg/mL and 9.033±0.585 pg/mL, respectively (Table S7A). IL33 levels were also assessed in COPD patients. The plasma concentration of IL33 in COPD patients was found to be 5.92±0.089 pg/mL (Table S7B). Concentrations of the other cytokines studied (IL4, IL5, and IL13) were below the detection limit for both healthy and patient plasma samples (Table S6).

Pathway Analysis Reveals Involvement Of Immunopathways
Pathway analysis was performed by IPA after merging the five most predominantly observed apolipoproteins (ApoA1, Apo B, ApoD, ApoE, and ApoA4) and IL33. Figure 5 reveals the different probable pathway hits obtained. Most of the pathways with ApoE as a component were found to be

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**Figure 2** Apolipoproteins involved in atopic asthma.
**Notes:** (A) Heat map showing expression patterns of the 14 apolipoproteins across healthy subjects and atopic asthmatics. Only ApoA4 and ApoE expression were altered significantly among the healthy volunteers and atopic asthmatics. The heat map was generated using Morpheus software. (B) ApoA4 was reduced in atopic asthmatics. The box plot (generated using R packages) shows the expression trend of ApoA4 in blood plasma of healthy volunteers (n=5) and atopic asthma patients (n=5) obtained from liquid chromatography–tandem mass spectrometry (LC-MS/MS) in terms of normal spectral abundance factor (NSAF). (C) ApoE may render protection in atopic asthmatics. The box plot (generated using R packages) shows the expression trend of ApoE in blood plasma of healthy volunteers (n=5) and atopic asthma patients (n=5) obtained from LC-MS/MS based proteomics study in terms of NSAF.
involved in human immune response, i.e., acute-phase response signaling, IL12 signaling and production in macrophages, clathrin-mediated endocytosis signaling, and production of nitric oxide and reactive oxygen species in macrophages. ApoE and IL33 were both found to be involved in several pathways, such as LXR–RXR activation, FXR–RXR activation, atherosclerosis signaling, and LPS/IL1–mediated inhibition of RXR function. These pathways are involved in lipid metabolism and inflammatory response.

**Discussion**

All the atopic asthma patients used in this pilot study were found to be sensitive to almost all or a majority of the most common allergens, i.e., pollens (*H. annuus, C. nuifera* [common ragweed]), house dust mites (*Dermatophagoides farinases, Dermatophagoides pteronyssinus*) and molds (*Aspergillus fumigatus, Penicillium notatum, Candida albicans, Mucor racemosus*). Blood-plasma samples obtained from the patients and healthy subjects were used for the study. Modern clinical proteomics suggests the use of plasma for identification of diagnostic and therapeutic biomarkers, as the circulating bloodstream harbors tissue-leakage proteins and inflammatory proteins that may serve as biomarker candidates. Here, we showed that plasma levels of ApoE were significantly downregulated in atopic asthmatics compared to healthy volunteers and COPD patients. ApoE is a 34 kDa protein that plays a crucial role in cholesterol transport and lipid metabolism. It is expressed by several organs, including the lungs. In the airways, ApoE is expressed primarily by CD68$^{+}$ alveolar macrophages. Apart from its role in cholesterol metabolism, ApoE is known to modulate normal lung health. In our study, high abundance of ApoA1 was observed in both healthy subjects and asthmatics, but significant differences in
expression profiles were not noted. However, the level of ApoE was found to be reduced in cases of atopic asthmatics, suggesting a probable protective role of ApoE in asthma. Other studies have shown that ApoE functions as a negative regulator of airway hyperresponsiveness and goblet-cell hyperplasia, features commonly noted in asthma. In COPD patients, there was a reduction in expression of ApoE with respect to healthy subjects, but the reduction was not as marked as in atopic asthmatics. ApoM has been reported to be significantly overexpressed in COPD. ApoM levels were not significantly altered in atopic asthma patients, as evident from LC-MS/MS data (Table S3). ApoA4 was significantly downregulated in atopic asthmatics (34.99% decrease, \( P<0.05 \)) in our study. Apolipoproteins are released as tissue-leakage products into the bloodstream. However, ApoA4 mRNA is not expressed in the lungs, but rather produced primarily in the gastrointestinal tract. As such, further validation study was not conducted for ApoA4 in atopic asthmatics using Western blotting.

Apart from label-free plasma proteomics, cytokine profiling was done for IL4, IL5, IL13, and IL33 from blood-plasma samples. Concentrations of IL4, IL5, and IL13 were below the detection limits of the ELISA kits. Similar ELISA-based study of these cytokines has reported a lack of detection in plasma. However, IL4, IL5, and IL13 have been noted to be upregulated in sputum, bronchoalveolar lavage fluid, and peripheral blood lymphocytes of allergic asthma patients. For IL33, plasma levels of the cytokine were found to be significantly upregulated in atopic asthma patients compared to healthy individuals in this study. Similar trends in IL33 expression have been noted in bronchoalveolar lavage fluid and lungs of asthma patients in other studies, pointing toward its significant role in asthma pathogenesis.

IL33 belongs to the IL1 family of cytokines, and is expressed by a variety of cells, including epithelial cells, endothelial cells, and fibroblasts. It promotes Th2-type immune response by inducing the expression of inflammatory cytokines like IL4, IL5, IL13, IL1β, IL6, and TNFα. It has been noted that IL33 is associated with different types of inflammatory disease, such as rheumatoid arthritis, atopic allergy, asthma, and cardiovascular diseases, and can be targeted for therapeutic purposes.

In this study, the increased levels of IL33 support its possible role in promoting systemic inflammation in asthma. IL33 levels were found to increase significantly in COPD patients; however, the rise was not as marked as in atopic asthmatics. Similar trends in IL33 expression have also been noted in exhaled breath condensate from COPD and asthma patients.

An indirect interaction between ApoE and IL33 was observed through IPA of proteomics and cytokine data sets. IL33 is known to regulate expression of genes involved in cholesterol metabolism. However, the exact connection of ApoE and IL33 is not clearly known for asthma. Further experiments are required to study the role of ApoE in asthma and to understand the actual regulatory mechanisms between ApoE and IL33 involved in asthma pathogenesis. ApoE and IL33 were both found to be involved in several pathways from IPA: LXR–RXR activation, FXR–RXR activation, atherosclerosis signaling, and LPS/IL1-mediated inhibition of RXR function. Pathway analysis in IPA also showed ApoE to be involved in immunopathways: acute-phase response signaling, IL12 signaling and production in macrophages, clathrin-mediated endocytosis signaling, and production of nitric oxide and reactive oxygen species in macrophages. This suggests that apart
from its role in lipid metabolism, ApoE has significant roles in immunity.

The upregulation of IL33 and the simultaneous downregulation of ApoE serve as marker pairs, which mimics the two-gene relative expression–based biomarker or classifier system. Such two gene–based classifier systems have been highly successful in accurate distinction of clinically distinct but symptomatically similar diseases like leiomyosarcoma and gastrointestinal stromal tumors. As such, the identified ApoE and IL33 pair could serve as probable candidates for distinguishing atopic asthma from similar airway inflammatory disorders like COPD. The two-protein biomarker concept based on ApoE and IL33 and the ratio of these protein biomarkers can be implemented for testing asthma patients in future studies.

Despite interesting findings, the main limitation of our study was its relatively small sample and inclusion of atopic asthmatics only. Since the study participants were treatment-naïve asthmatics and COPD patients without any other health complications, recruitment was very difficult. Another limitation of this study was that the IgE count of the atopic asthmatics was highly variable, ranging from 385 to 1,993 IU/mL. Also, the study was not extended to nonatopic asthmatics, due to the unavailability of an adequate number of patients in the clinics during the study. Also, the study did not include nonatopic asthma patients, which is another limitation. Another important limitation was the depletion of six high-abundance proteins from blood plasma, which might have resulted in the removal of several other plasma proteins that form complexes with those six highly abundant plasma proteins. We considered ApoE as a tissue-leakage product in blood from lungs, but apolipoproteins are also secreted by other organs like the liver and intestine. However, the human lung is also known to release huge amounts of cholesterol and apolipoproteins in the bloodstream.

**Conclusion**

In this study, a subset of patients was chosen based on their history, spirometry, and total blood-IgE levels. It can be concluded that in spite of the diversity among patients, distinct trends were found in expression levels of ApoE and IL33 in asthma patients’ samples compared to healthy subjects and COPD patients. Expression levels of ApoE were downregulated, whereas IL33 was highly expressed in asthmatics. Therefore, simultaneous downregulation of
ApoE and upregulation of IL33 identified from blood plasma may serve as probable candidates for a two protein–based diagnostic and prognostic biomarker of atopic asthma and can be explored for therapeutic purposes.

**Abbreviations**

COPD, chronic obstructive pulmonary disorder; FEV1, forced expiratory volume in 1 second; ApoE, apolipoprotein E; IL33, interleukin 33; LC-MS/MS, liquid chromatography–tandem mass spectrometry; NSAIF, normalized spectral abundance factor; IPA, ingenuity pathway analysis.

**Ethics Approval And Informed Consent**

Human Ethics Committee clearance (BIHEC/2014-15/4) from Bose Institute, Kolkata was obtained to work with human blood samples. All human blood samples were collected with written informed consent from patients and healthy subjects. The methods used were in accordance with relevant guidelines and regulations. All the experimental protocols were approved by the aforementioned committee.

**Availability Of Data And Materials**

Supporting figures and tables are included in the Supplementary file and Table S2.

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**Author Contributions**

SS, SGB, and AD conceptualized and designed the study. MB, SM, and AD were involved in acquisition of data. SM, MB, SGB, and SS were involved in data analysis. All authors contributed toward drafting and critically revising the paper, gave final approval of the version to be published, and agree to be accountable for all aspects of the work.

**Disclosure**

The authors report no conflicts of interest in this work.

**References**


33. Martin K, Viera K, Petk C, Marie N, Eva T. Simultaneous analysis of cytokines and costimulatory molecules concentrations by ELISA technique and of probabilities of measurable concentrations of interleukins IL-2, IL-4, IL-5, IL-6, CXCL8 (IL-8), IL-10, IL-13 occurring in plasma of healthy blood donors. Mediat Inflamm. 2006;2006(5):65237.


