

A transcriptome-driven analysis of epithelial brushings and bronchial biopsies to define asthma phenotypes in U-BIOPRED

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CHSK, DL, BDM and SP performed the analysis. CHSK, SP, ML, FB, AR, CA, IP and YG designed the analytical approaches taken and analysed the results. RD, PJS, PH, UH, SD, PC, DS, BD, NK, TS and KFC participated in the clinical characterisation of the patients. SJW and PH participated in the immunohistochemical studies. CR participated in experimental work and analysis. SF, LF, AS and JC were part of the data curation team. IMA, RD, PJS and KFC conceived & designed the study, and CHSK, IMA & KFC coordinated the data and drafted the manuscript. All have read the final version.

At a glance commentary

Scientific knowledge on the subject

Unbiased clustering of asthma using clinical features has derived phenotypes of limited pathobiological relevance. It is not known whether semi-biased clustering on transcriptomic data will provide additional information on underlying asthma biology.

What the study adds to the field

We show that semi-biased clustering of transcriptomic profiles from asthmatic airway samples produces biologically relevant clusters that have implications for directed or personalized therapy. We report subgroups of eosinophilic asthma driven by T-helper Type 2 cells and corticosteroid insensitivity signatures that featured either high submucosal eosinophils or sputum eosinophils. An inference scheme using currently-available inflammatory biomarkers can be used to predict the subtypes of gene expression.

Abstract (245 words)

Rationale and objectives: Asthma is a heterogeneous disease driven by diverse immunologic and inflammatory mechanisms. We used transcriptomic profiling of airway tissues to help define asthma phenotypes.

Methods: The transcriptome from bronchial biopsies and epithelial brushings of 107 moderate-to-severe asthmatics were annotated by gene-set variation analysis (GSVA) using 42 gene-signatures relevant to asthma, inflammation and immune function. Topological data analysis (TDA) of clinical and histological data was used to derive clusters and the nearest shrunken centroid algorithm used for signature refinement.

Results: 9 GSVA signatures expressed in bronchial biopsies and airway epithelial brushings distinguished two distinct asthma subtypes associated with high expression of T-helper type 2 (Th-2) cytokines and lack of corticosteroid response (Group 1 and Group 3). Group 1 had the highest submucosal eosinophils, high exhaled nitric oxide (FeNO) levels, exacerbation rates and oral corticosteroid (OCS) use whilst Group 3 patients showed the highest levels of sputum eosinophils and had a high BMI. In contrast, Group 2 and Group 4 patients had an 86% and 64% probability of having non-eosinophilic inflammation. Using machine-learning tools, we describe an inference scheme using the currently-available inflammatory biomarkers sputum eosinophilia and exhaled nitric oxide levels along with OCS use that could predict the subtypes of gene expression within bronchial biopsies and epithelial cells with good sensitivity and specificity.

Conclusion: This analysis demonstrates the usefulness of a transcriptomic-driven approach to phenotyping that segments patients who may benefit the most from specific agents that target Th2-mediated inflammation and/or corticosteroid insensitivity.

Key words: severe asthma, bronchial biopsies, bronchial brushings, corticosteroid insensitivity, T-helper Type 2(Th2), exhaled nitric oxide, gene-set variation analysis, asthma phenotype

Introduction

Asthma is a heterogeneous disease that presents with varying degrees of inflammation in the airways in association with airway wall structural changes. Both infiltrating and resident airway structural cells participate in the inflammatory and remodeling processes of asthma and various mechanisms have been proposed that could underlie the asthmatic inflammatory process (1). T-helper Type 2 (Th2) cell activation characterized by the secretion of IL-4, IL-5 and IL-13 is a key mechanism of allergic asthma with these cytokines being overexpressed in the bronchial submucosa of asthmatic patients (2, 3). Th2-high asthmatic patients are characterized by a greater degree of bronchial hyperresponsiveness, higher serum IgE levels, greater blood and airway eosinophilia, subepithelial fibrosis and airway mucin gene expression (4).

Th1 and Th17 cells, and their products are also important in asthma. Th1 cells are characterised by IFN γ production which plays a role in dealing with viral infections and in autoimmunity and both Th1 cells and IFN γ are overexpressed in asthma subjects (5-7). These have also been shown to be important in severe asthma (8, 9). Th-17 cells express IL-17A, IL-17E, IL-17F and IL-22, and may mediate corticosteroid-resistant airway inflammation and airway hyperresponsiveness in mice (10). IL-17A and IL-17F, have been localised in the airways of patients with severe asthma (11).

Oxidative stress is another important feature of asthma, particularly those with severe asthma, and is associated with increased levels of reactive oxygen species (ROS) and compromised antioxidant responses (12, 13). ROS may mediate various features of severe asthma such as the chronic inflammatory response, the hypercontractility of the airways and corticosteroid insensitivity (14-16). Finally, structural abnormalities can alter airway mechanics and contribute to bronchial hyperresponsiveness, while structural cells can also contribute to inflammatory processes through release of cytokines, chemokines, growth factors and extracellular matrix elements (17-19).

Therefore, asthma is a syndrome with many potential endotypes and defining these is an essential step towards providing personalized treatments (20, 21). Defining asthma phenotypes from clinical and physiological parameters, including currently-used inflammatory biomarkers, is limited in expressing the heterogeneity of underlying mechanisms (20, 22, 23). This may not be surprising considering the complexity of immunoinflammatory mechanisms and the diversity of cell types implicated in asthma (24-27). Deriving asthma clusters according to differentially-expressed genes may be a better approach to capture the diverse pathways of asthma pathobiology that could lead to the identification of important targets for therapy and offer insights towards achieving personalized medicine (20, 28, 29). This approach has been used to identify genes from sputum cells that linked pathway-based transcriptomic clusters to clinically-important features of asthma (30).

Conventional gene set enrichment (GSE) methods have limitations owing to the assumption of two classes (e.g. case versus control) (31). Gene Set Variation Analysis (GSVA) addresses this limitation by calculating GSE scores as a function of genes inside and outside a specifically-defined gene set (32) and can indicate functional activity across individual samples (33). Topological data analysis (TDA) that recognises the invariant shape of complex data sets (34, 35) has been used to overcome the challenges that stochastic gene expression presents on conventional hierarchical or k-means clustering (36).

In the current study, we analysed transcriptomic data from bronchial biopsies from asthma participants from the Unbiased BIOMarkers in PREdiction of respiratory disease outcomes (U-BIOPRED) project (37) and performed supervised clustering using 42 gene sets associated with asthma and immune/inflammatory pathways by GSVA. We then explored the relationship between these GSVA signatures and their unique clinico-histopathological features. This was then further investigated in the transcriptome from epithelial brushings of the same participants with the aim of examining whether transcriptomic signatures may help define subsets of Th2-high eosinophilic and non-Th2 asthma from distinct airway

compartments. Some of the results of these studies have been previously reported in the form of an abstract (38).

Materials and methods (500 words)

Details of analytical methods are provided in the *On-line Supplement*.

Clinical data

107 participants (**Table 1**) with moderate-to-severe asthma from the U-BIOPRED cohort underwent fiberoptic bronchoscopy for epithelial cell brushings and bronchial biopsies(39). Pre-bronchodilator spirometry, exhaled nitric oxide (FeNO), sputum differential cell count, skin prick tests, serum total IgE, and differential blood count were measured(39). Oral corticosteroid dependence was assessed by determining the lowest maintenance dose of daily prednisolone. The study was approved by the Ethics Committees for each of the 16 clinical recruiting centres. All participants gave written and signed informed consent.

Immunohistochemistry for submucosal cells in the bronchial biopsies

Samples from glycolmethacrylate embedded sections (2 μ m) were stained with monoclonal antibodies against CD3, CD4, CD8, neutrophil elastase and EG2. Cell counts were performed in a blinded fashion and expressed as positive cells/mm². These results have been presented previously in an abstract form [43].

Microarray analysis of mRNA

Expression profiling was performed using AffymetrixU133 Plus 2.0 microarray (Affymetrix, Santa Clara, Calif) on total RNA extracted from bronchial biopsies and epithelial brushings. RNA purity (RIN >9.5) was measured by Agilent Bioanalyser (Agilent, Santa Clara, Calif). Raw data were quality assessed and pre-processed by robust multi-array average normalization using Almac Pipeline and Pre-processing Toolbox (Almac, Craigavon, United Kingdom). Probes of low expression were filtered according to robust multi-array signal values <5 and batch/technical effects were adjusted as covariates using a linear model.

Gene Set Variation Analysis (GSVA)

42 gene sets containing 2431 genes were manually curated: 6 for six key immune cell type-specific gene expression associated with their differentiation and activation, 6 for evolutionarily-conserved transcriptional signatures for Th1 and CD8 memory T cell differentiation, 5 of effector CD4+ T cell differentiation for Th2, Th17 and Treg cell subset, 6 for chronic effect of oxidative stress in response to ozone from a mouse model and genes associated with COPD from human lung, 7 from peripheral blood mononuclear cells (PBMCs) in four autoimmune diseases, 7 for asthma-specific mechanism driven by T cells and 5 from human lung biopsy, airway smooth muscle cells and PBMCs in response to corticosteroid treatment (Supplementary TableS1).

TDA and machine learning algorithm

TDA was used to explore clusters driven by gene signatures utilizing the Ayasdi Core™ (Ayasdi, Inc. California, USA) program and the nearest shrunken centroid algorithm was used to further define the optimal number of classifiers. Details of the methods are provided in Supplementary file.

Statistical analysis

All datasets needed for this analysis were uploaded from the transSMART system, an open-source knowledge management platform for sharing research data(40) supported by the European Translational Information and Knowledge Management Services (eTRIKS) project. All categorical variables were analyzed using Fisher's exact test. Student's t test was used for continuous variables with normal distribution, otherwise Wilcoxon rank-sum test was used. A linear model for microarray data (Bioconductor R package limma) with false discovery rate (FDR) correction was used for differential expression gene analysis, and recursive partitioning (R package PARTY) for decision tree learning. An FDR<0.05 was considered

statistically significant.

Results

Transcriptomic profiles from 91 bronchial biopsies and 99 epithelial brushings were obtained, of which 83 samples were paired from the same subject.

Differentially-expressed gene and gene set analysis

We performed differential expression analysis of individual genes and of gene sets comparing moderate asthma versus severe asthma in both epithelial brushings and bronchial biopsies. None of the individual genes or gene sets were significantly different between epithelial brushings from severe and non-severe asthma after multiple testing correction for FDR (**Fig 1A & B**). In addition, only 2 genes with marginally significant expression and three GSVA gene sets with a trend toward significant expression were found when comparing expression profiles in bronchial biopsies between severe and non-severe patients (**Fig 1C & D**). Therefore, the clinical classification of moderate versus severe asthma provides a limited framework for the identification of differential gene expression profiles.

There was a significant correlation between the ES of the Th2 signature and the relative steroid insensitivity signature in both biopsy and brushings (Fig S1), together with a significant relationship between the oxidative stress signature and T-cell, Th1 and Th2 signatures (Fig S2), and between the oxidative stress signatures and relative corticosteroid insensitivity signatures in both biopsy and brushings (Fig S3). These correlations indicate potential mechanistic interactions between Th2 and Th1 genes with oxidative stress and relative corticosteroid insensitivity which was defined by the signature of genes that are usually down-regulated by corticosteroids showing no change or being up-regulated.

In both biopsy and brushings, *CD44* as a constituent of corticosteroid insensitivity signature had the most extensive association/interaction with a subset of Th2 signature genes including *CCL26*, *IL1R2* and *CST2*. In addition, a highly correlated association between

NELFE (within the corticosteroid insensitivity signature) and *ATP5J* (within the Th2 signature) was also seen in both biopsy and brushing compartments.

Clinical clusters driven by bronchial biopsy GSVA analysis

We performed an analysis of the bronchial biopsy transcriptome using GSVA in order to investigate the immune cell-specific characteristics and distinctive pathophysiological processes in asthma. Unsupervised clustering of GSVA gene sets applied to biopsies using TDA identified a cluster characterized by high submucosal eosinophils (5.6 vs. 1.2/mm², $p=6.6 \times 10^{-5}$), high FeNO (34.0 vs. 21.0 ppb, $p=0.028$) and high oral corticosteroid use (41.0 vs 15.4%, $p=0.006$) (**Fig 2 & Table 2**). Comparing Cluster A to Cluster non-A, 26 out of 42 gene sets were found to be differentially expressed ($p<0.05$) (**Fig 3**). A similar clustering was obtained when hierarchical clustering was performed using hierarchical clustering (**Fig 4**).

Finding signatures for optimal cluster classification

In order to determine the GSVA signatures that best distinguished Cluster A from Cluster non-A, the shrunken centroid algorithm was applied to these 26 GSVA gene-sets and produced a classification accuracy of 82.4% (Fig 5A). After feature reduction, 9 GSVA gene-sets of non-zero value which were expressed in the opposite direction in Cluster A and non-A were retained as centroid classifiers of discrimination (Fig 5B, Table 3). ROC curve analysis demonstrated a high performance of this 9-gene-set signature model (AUC: 0.866; 95%CI: 0.796-0.927, $p=1.3 \times 10^{-9}$) in differentiating membership of Cluster A, providing a sensitivity and specificity of 84.6% and 90.4%, respectively (Fig 5C).

Clusters from epithelial brushings and from combined epithelial brushings and bronchial biopsies

In order to test whether these 9 GSVA signatures were also able to discriminate clusters

in bronchial brushings, a similar analysis was performed on the 99 epithelial brushing samples. The 9 GSVA signatures from Cluster A determined in bronchial biopsies was found in 31 (31.3%) of subjects as calculated using the nearest shrunken centroids of the signatures. This subgroup reproduced the clinical characteristics noted in biopsies in terms of significantly higher submucosal eosinophils (5.8 vs. 1.5/mm², $p=1.3\times 10^{-5}$), higher exhaled NO (51.5 vs. 21.0 ppb, $p=4.2\times 10^{-4}$) and higher oral corticosteroid use (40.7 vs 17.9%, $p=0.032$; **Table 4**). In addition, higher eosinophil counts in blood and sputum and higher numbers of submucosal CD3, CD4 and CD8 T cells in bronchial submucosa were also noted.

The relative expression of these 9 GSVA signatures in bronchial biopsies and epithelial brushings is shown in **Table 5**: Group 1 (19 subjects, 23%) expressed these 9 signatures in both bronchial samples and epithelial brushing cells, Group 2 (17 patients, 20%) expressed these signatures in bronchial biopsies only, Group 3 (8 patients, 10%) in epithelial brushing cells alone and Group 4 (39 patients, 47%) expressed these 9 GSVA signatures in neither compartment.

Compared to Group 4 participants, two subtypes of patients with eosinophilic inflammation and relative steroid insensitivity were noted (Groups 1 and 3; **Table 6**). Group 1 had the highest submucosal eosinophils (6.1 vs. 1.0/mm², $p=7.64\times 10^{-6}$), CD3 (45.3 vs. 33.3/mm², $p=0.048$), CD8 (26.4 vs. 13.9/mm², $p=0.020$) T cells, FeNO (56.5 vs. 19.0 ppb, $p=0.001$), acute exacerbation rate (3.0 vs. 1.5 times/year, $p=0.022$), oral corticosteroid use (47.7 vs 12.8%, $p=0.010$) and intermediate-high sputum eosinophils (8.2 vs 0.6%, $p=0.017$). In contrast, Group 3 participants possessed the highest levels of sputum eosinophils (15.8 vs. 0.6%, $p=0.014$), high BMI (33.1 vs. 28.0, $p=0.010$) and a trend toward intermediate-high FeNO (35.5 vs. 19.0 ppb, $p=0.058$).

Levels of gene expression of IL-33 and TSLP but not IL-25 were relatively increased in Cluster A compared to Cluster non-A and in Group 1 compared to Groups 3 and 4 (for IL-33) and to Group 4 (for TSLP) in bronchial biopsies but not in epithelial brushings (Fig S4).

Inference of phenotype from non-invasive clinical measurements

The clinical features and non-invasive biomarkers from the patients classified into the 4 groups were used to build an inference tree framework. FeNO >55 ppb (Fig 6 Node 7, 58% for Group 1), sputum eosinophil >4.5% (Fig 6 Node 6, 42% for Group 1) or oral corticosteroid dependency (Fig 6 Node 4, 36% for Group 1) predicted subjects as the highest eosinophilic inflammation and steroid insensitivity subtype (Group 1), with an 84% sensitivity and 72% specificity. Patients with FeNO ≤55 ppb, sputum eosinophil ≤4.5% and not dependent on oral corticosteroids had an 86% probability of having non-eosinophilic inflammation (Fig 6 Node 5, 22% for Group 2 and 64% for Group 4). Clustering analysis based on the 3 top gene sets in the 9-GSVA signature associated with these clinical traits confirmed the molecular association of Group 1 subjects with high Th2 activation and corticosteroid insensitivity in bronchial biopsy and epithelial brushing samples (Fig 7).

Discussion

We clustered gene expression data from bronchial biopsies and epithelial brushings obtained from patients with moderate-to-severe asthma in order to define clinical phenotypes of asthma. Direct comparison of the transcriptome obtained from severe asthma to moderately-severe asthma subjects led to the identification of few differentially-expressed genes in bronchial biopsies and epithelial brushings using false discovery rate. We then examined their gene expression profiles based on important disease drivers as described by specific gene-sets and derived phenotypes from the gene-set clusters. A collection of 9 gene-set signatures applied to genes expressed in bronchial biopsies and airway epithelial brushings led to two subtypes of patients with eosinophilic inflammation and relative corticosteroid insensitivity. Using machine-learning tools, we found that currently-available inflammatory biomarkers such as sputum eosinophilia and exhaled nitric oxide levels could be used to predict the subtypes of asthmatics described by gene expression profiling. This approach could ultimately define patients who may benefit the most from specific molecular agents that target Th2-mediated inflammation and/or relative corticosteroid insensitivity. It also defines the importance of the site of expression of these 9 gene-set signatures in either biopsies or brushings or in both in determining the phenotype.

It is now generally accepted that asthma is a heterogeneous disease with distinct phenotypes. Our analysis confirmed the presence of a previously-described Th2-high cluster derived from an analysis of Th2-gene expression in epithelial brushings which is indicative of Th2 activation (4, 41) and which was defined by two of the 42 GSVA gene-sets we used. This signature has been linked to other features of asthma, particularly corticosteroid responsiveness. The remaining signatures we used reflected the heterogeneous nature of asthma and represent many of the mechanisms that have been proposed to drive the asthmatic disease process in addition to the Th2 pathway.

Interestingly, GSVA gene-sets associated with steroid responsiveness accounted for the second and sixth most important signatures that were significantly up-regulated, when they

are expected to be down-regulated by corticosteroids in non-severe asthmatics. In addition, a moderate correlation was noted between Th2 and corticosteroid insensitivity signatures (*Supplementary Figure S1*) reflecting the clinical problem of clinical insensitivity to corticosteroid treatment in a subgroup of severe asthmatics. Further analysis of the genes that were associated in the corticosteroid insensitivity and Th2 signatures in the biopsies and brushings has shown an association between *CD44* and the Th2-associated genes *CCL26*, *IL1R2* and *CST2*, indicating potential underlying mechanisms. The patients in Group 1 where the corticosteroid insensitivity signatures were most highly expressed in the airway epithelium and bronchial biopsy were on the highest doses of oral corticosteroids. Therapeutic strategies other than corticosteroid dose escalation should be considered for these patients.

Relative corticosteroid insensitivity was defined at the molecular level by detecting the signature of genes usually down-regulated by corticosteroids as being up-regulated. However, such a definition assumes that the patient is adherent to corticosteroid therapy, particularly oral therapy. The patients in UBIOPRED have been followed-up for at least for 6 months in a specialist severe asthma clinic during which time their degree of compliance has been deemed to be satisfactory. Our observation that Group 1 where the expression of the relative corticosteroid insensitivity signature together with the Th2 high and oxidative stress signatures were found in both bronchial biopsies and brushings was associated with a greater level of oral steroid dependency would support the notion that relative corticosteroid insensitivity may be associated with corticosteroid dependency. A potential limitation of the analysis of relative corticosteroid insensitivity is the use of the gene signatures obtained from patients with COPD, where the gene response to corticosteroids may be different from that seen in asthma.

Two gene-sets used were related to ozone stimulation as a marker of elevated oxidative stress and other important GSVAs signatures that were critical in differentiating asthma included those from monocytes relating to autoimmune diseases and to Th1 immunity as

reflected by the increase in the numbers of submucosal CD3⁺ and CD8⁺ T cells. Amongst them, we noted that the oxidative stress signature revealed a moderate to high correlation with Th1 signature and with the steroid insensitivity signature (*Supplementary Figure S2 & S3*). While ROS has been reported to play a role in promoting asthmatic inflammation (14, 15) and reducing steroid responsiveness (16), the mechanism remains largely controversial. Therefore, the implication of oxidative stress in our finding may shed light on the impact of ROS in asthma. Overall, we found that there were non-Th2 pathways that were associated with severe asthma in the analysis of both bronchial biopsies and epithelial brushings despite the presence of eosinophilic inflammation.

One potential concern regarding our approach in this study is the appropriateness of gene sets used for clustering. We endeavored to be as inclusive as possible by using gene sets reflecting the diversity of processes previously reported to be associated with various aspects of asthma pathobiology in addition to key mechanisms involved in chronic obstructive airway disease, baseline immune cell signatures and those reported in a number of systemic immune disorders. In keeping with this deployment of gene-sets, we found that Group 1 subjects were distinguished by diverse immune activation across airway compartments, whereas nearly half of Group 4 participants did not show pronounced activation of the immune cells selected. Interestingly, the enrichment of GSVA signatures representative of multiple immune pathways including Th2, Th1, Th17, neutrophil, dendritic cells and influenza response were largely lower in Group 4 compared to Group 1 (*Supplementary Figure S4*), suggesting the lack of a dominant disease driving immune pathway and may reflect a pauci-immune phenotype of asthma. Further unbiased bioinformatic analyses such as the use of weighted gene co-expression network analysis (WGCNA) will be necessary to determine other potential disease-driving mechanisms. In a preliminary WGCNA analysis of epithelial cell gene expression, we found several modules related to FeNO, sputum eosinophilia and oral corticosteroid use with distinct eigengenes associated with these clinical parameters.

Although the 9 GSVA gene-set signature was initially derived from gene expression in

bronchial biopsies, its expression in epithelial brushings was associated with similar clinical characteristics. The strength of this signature was greater when it was highly expressed in epithelial cells than when it was expressed solely in bronchial biopsies particularly when linked to clinical parameters such as FeNO, blood eosinophils, sputum eosinophils and subtypes of submucosal T cells. This finding echoes previous studies highlighting the key role of the bronchial epithelium in dictating Th2 asthma phenotype and emphasizes the importance of the bronchial epithelium in driving asthma pathobiology (42). However, cross-talk between these compartments must exist as the most severe asthma subtype we describe here featuring eosinophilic inflammation and corticosteroid insensitivity (Group 1) occurred in subjects where the 9 GSVA signatures were concomitantly highly expressed in both bronchial biopsies and epithelial brushings. Interestingly, the expression of the 9 GSVA gene-set signatures in epithelial cells was mainly associated with sputum eosinophilia (Group 3 participants).

This observation may reflect priming of the asthmatic airway upon which Group 1 subtype was dependent when the signature was also found in the bronchial biopsies. This priming is further supported by the observation that subjects lacking this 9 GSVA signature expression in epithelial cells do not show eosinophilic inflammation irrespective of their signature expression in bronchial biopsies (Groups 2 and 4 participants). This highlights the need to further clarify the role played by airway epithelial cells in driving the development of airway inflammation in asthma. Indeed, recent work indicates that the airway epithelium is an important upstream cell that controls the regulation of Th2 cytokines through the production of cytokines such as thymic stromal lymphopoietin (TSLP), IL-25 and IL-33, which can be stimulated by exposure of epithelial cells to external stimuli such as pollutants, viruses and allergens (42, 43). The epithelium in severe asthma is also reported to be thicker than in mild-to-moderate asthma with increased proliferation, apoptosis and release of pro-inflammatory factors (44). However, our results indicate that IL-33 and TSLP gene expression is increased in the biopsies not in the brushings in asthmatic patients in Cluster A

and in Group 1 who also show the highest Th2 gene signature with bronchial eosinophilia.

The subtypes of asthma reported here, based on transcriptomic rather than clinical data, share some overlapping clinical and biomarker features with phenotypes driven by clinical parameters from the Leicester (23) and Severe Asthma Research Program (SARP) (22) clustering studies. The small Group 3 phenotype (10% of patients) reported here is very similar to Cluster 3 in SARP (8%) in terms of higher BMI and higher sputum eosinophils but lower biopsy eosinophils, relative to the remaining asthmatic groups. In addition, our Group 1 phenotype (23%) has some resemblance to Cluster 5 in SARP (16%) regarding the lower FEV₁, whereas they were discordant with respect to its neutrophilic profile. Indeed, we found no difference in the level of either blood or sputum neutrophilia in the 4 groups. These findings provide a bridge between phenotypes derived from clustering of clinical parameters and our approach of using gene expression profiles within key airway structural cells along with tissue histopathology. We have provided a path to exploring the mechanisms underlying different subsets of patients where discordant inflammation is found.

A key question is whether there is any clinical utility or application of the phenotypes described here particularly as they were determined using transcriptomic analysis of samples obtained using bronchoscopy. The clinical utility is addressed by the tree-based recursive partitioning algorithm selecting significant non-invasive parameters or clinical traits based on their conditional distribution statistics against phenotypes using an unbiased permutation method (45). This approach successfully translated our phenotypes originally defined by gene signatures and by histopathological results to a clinically inferable scheme. Following a stepwise binary value split of FeNO (>55ppb), sputum eosinophils (>4.5%) or oral corticosteroid dependency, this scheme identified Group 1 subjects with an 84 % sensitivity and 72% specificity. This is clinically important as this group bears the Th2 and steroid insensitivity signatures across airway structural cells that justifies the need for molecular agents that target Th2-specific pathobiology (20). However, other groups such as Group 2 and 3 may also benefit from these agents with the possibility that different targets within the

Th2 phenotype may be more appropriate for each of the 3 groups. Thus, epithelial-dependent release of different eosinophilic cytokines/chemokines might be appropriate for Groups 1 and 3, while in Group 2, a non-eosinophilic Th2 or T2 targets might be more useful.

In conclusion, transcriptomic profiling in bronchial biopsies and epithelial brushing cells showed heterogeneity of underlying mechanisms with respect to Th2 and relative steroid insensitivity signatures which allowed the definition of different asthma phenotypes. Routinely-used clinical biomarkers successfully translated these findings, and used in this way may help in taking therapeutic decisions for patients with severe eosinophilic inflammation and relative corticosteroid insensitivity.

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Figure legends:

Figure 1: Differential expression analysis comparing moderate versus severe asthma from bronchial biopsies and epithelial brushing transcriptomic profiles on the basis of individual genes or Gene set variation analysis (GSVA). Given the FDR correction (broken line), no differentially expressed genes (A) or gene sets (B) in epithelial brushing was noted. Two genes were marginally significant (C) and three gene sets (D) showed a trend towards significance in bronchial biopsies. ES: enrichment score.

Figure 2: Topological network analysis of gene-set variation analysis using 42 gene sets of relevance to the pathogenesis of asthma in bronchial biopsies revealing one large signature-driven cluster (Cluster A) and 4 smaller distinct clusters. Cluster A was distinguished by higher submucosal eosinophil counts, higher levels of nitric oxide in exhaled breath and higher oral corticosteroid use. The network is coloured on the basis of the submucosal eosinophil count with low cell count in blue and high in red. The colour key histogram (bottom left) showed the number of subjects with respect to the given cell count (x-axis) from low to high. Subjects of high submucosal eosinophil count were noted to be highly enriched in Cluster A as compared to the rest.

Figure 3: Heat map of differential gene set expression from bronchial biopsies between Cluster A (blue bar) and Cluster non-A (grey bar). High expression is denoted by red and low expression by green. Twenty-six of 42 gene sets used were significantly differentially-expressed in Cluster A with 23 over-expressed and 3 under-expressed.

Figure 4: Hierarchical clustering showing 91 bronchial biopsies on 42 GSVA signatures and the color bar revealed the clustering result from TDA on the same dataset. The Cluster A (red bar) identified by TDA method overlapped substantially with the subgroup of patients identified by hierarchical clustering that shaded in grey. The agreement on two clustering

methods measured by Rand and adjusted Rand Index were 0.7 and 0.4, respectively.

Figure 5: Machine-learning algorithm using shrunken centroid method on 26 differentially-expressed gene sets in bronchial biopsies. Panel A: Training of classifiers for Cluster A was evaluated for classification error using 10-fold cross-validation. A threshold of 1.82 (red broken line) was selected which reduced classifiers to 9 gene-sets at an 82% cross-validated accuracy. Panel B: Centroid profile of the 9-gene-set signatures. Length of the centroid denoted the relative amount the expression was away from the overall mean expression for each given gene-set signature. From top down, the centroids of Cluster A and Cluster non-A were ranked in decreasing amount. Panel C: Discrimination performance of the 9 gene-set signatures (AUC: 0.866; 95% CI:0.796-0.927, $p=1.3e^{-09}$) based on the estimation of the Cluster A probability from Gaussian density distribution. The probability cut-off at 0.5 gave a sensitivity and specificity of 84.6% and 90.4% respectively, with the confidence intervals shown.

Figure 6: Tree-based inference scheme for the probability of Group 1 to Group 4 using two non-invasive measurements (FeNO: $p=0.014$, sputum eosinophil counts: $p=0.012$) and one clinical parameter (oral corticosteroid dependency: $p=0.030$). Group 1 (red bar) represented a phenotype of high eosinophilic inflammation and steroid insensitivity. Group 3 (pink bar) represented a phenotype of medium eosinophilic inflammation and Group 2 & 4 (black bar) low eosinophilic inflammation.

Figure 7: Three signatures related to Th2 and steroid insensitivity in epithelial (above the broken line) and biopsy cells (below the broken line) across 83 asthmatics were used for clustering. Subject's membership as Group 1 to 4 were shown as color bar and mapped underneath the dendrogram. Group 1 (magenta) patients with the highest and most extensive expression of these signatures were mainly clustered to the right. Group 3 (purple) subjects

with the signatures solely in the epithelium also clustered closely. Group 2 (light blue) and 4 (grey) subjects interlaced with each other and clustered to the left. These 3 out of the 9 signatures were the major representative features of Group 1 and 3 asthma subtypes.

Table 1. Demographic and clinical characteristics of 107 asthmatic subjects

Variables[†]	
Age (years)	45.9 (13.9)
Female	59 (55.1%)
BMI	28.7 (5.8)
Severe asthma category	67 (62.6%)
Oral corticosteroid use (number)	29 (27.1%)
Exacerbation numbers (per year)	2 (1-4)
Atopy	76 (71.0%)
Allergic rhinitis	53 (49.5%)
Eczema	42 (39.3%)
Nasal polyp	27 (25.2%)
FEV ₁ (% predicted)	74.9 (62.3-95.1)
Total serum IgE (IU/ml)	115.6 (43.0-316.0)
Blood eosinophil (%)	3.1 (1.7-4.4)
Sputum eosinophil (%)	1.5 (0.4-7.5)
Sputum neutrophil (%)	52.5 (36.7-70.1)
FeNO (ppb)	31.5 (19.0-53.8)
Bronchial biopsy available	91 (85.0%)
Epithelial brushing available	99 (92.5%)

†: Data presented as N (%) and mean (SD) or median (IQR).BMI: Body mass index, FEV₁: Forced expiratory volume in 1 second, FeNO: Fractional exhaled nitric oxide

Table 2. Clinical characteristics of Cluster A versus non-A

Variables[†]	Cluster A	Cluster non-A	p-value
No (%)	N= 39	N=52	
Age (year)	48.4±13.6	45.8±13.1	0.355
Female	18 (46.2%)	31 (59.6%)	0.288
BMI	28.1±4.9	28.6±5.8	0.661
Severe asthma	26 (66.7%)	30 (57.7%)	0.258
Oral corticosteroid use	16 (41.0%)	8 (15.4%)	0.006
Exacerbation number (time/year)	2.5 (2.0-3.75)	2.0 (1.0-3.0)	0.139
Atopy	25 (64.1%)	40 (76.9%)	0.356
Allergic rhinitis	17 (43.6%)	26 (50.0%)	0.793
Eczema	15 (38.5%)	21 (40.4%)	1.000
Nasal polyp	14 (35.9%)	11 (21.2%)	0.092
FEV ₁ (%)	74.2 (59.8-91.6)	74.7 (62.3-94.6)	0.630
Total serum IgE (IU/ml)	125.0 (41.0-231.5)	104.0 (44.0-364.5)	0.813
Blood eosinophil (%)	3.2 (1.6-4.8)	2.7 (1.7-3.8)	0.523
Sputum eosinophil (%)	3.9 (0.6-12.1)	1.1 (0.2-4.7)	0.156
Sputum neutrophil (%)	53.2 (42.7-76.9)	52.4 (34.6-63.4)	0.421
FeNO (ppb)	34.0 (20.0-61.0)	21.0 (14.5-40.0)	0.028
<i>Submucosal cells (mm⁻²)</i>			
Eosinophils	5.6 (2.7-11.4)	1.2 (0-3.3)	6.6 x 10 ⁻⁵
Neutrophils	11.0 (6.6-18.1)	12.1 (6.8-18.4)	0.833
CD3 T cells	36.3 (22.6-50.0)	35.4 (16.6-49.5)	0.471
CD4 T cells	11.2 (5.9-18.9)	9.5 (4.8-15.8)	0.545
CD8 T cells	17.8 (9.5-30.4)	14.2 (4.9-27.7)	0.531

†: Data presented as N (%) and mean (SD) or median (IQR).

BMI: Body mass index, FEV₁: Forced expiratory volume in 1 second, FeNO: Fractional exhaled nitric oxide

Table 3. Nine GSVA signatures differentiating Cluster A from the rest

GSVA signature	ES difference	p-value	FDR
Lung.brushings.Th2high.asthma.HS.IVV.UP	0.578	6.0×10^{-10}	2.5×10^{-8}
Lung.biopsy.COPD.FLU.SAL.HS.IVV.DOWN	0.308	5.3×10^{-6}	7.4×10^{-5}
Lung.biopsy.HDM.Rhesus.IVV.UP	0.275	7.8×10^{-7}	1.6×10^{-5}
PBMC.Systemic immune disorders.HS.IVV.UP	0.203	0.001	0.005
Th1.activated.HS.IVS.UP.2	0.199	7.2×10^{-6}	7.6×10^{-5}
PBMC.asthma.GC.HS.IVS	0.234	8.4×10^{-5}	7.1×10^{-4}
Tcell.activated.HS.IVS	0.192	1.5×10^{-4}	0.001
Ozone.Air.MM.IVV.Up	0.157	8.9×10^{-4}	0.004
Ozone.MM.IVV.Down	-0.159	2.1×10^{-4}	0.001

ES: enrichment score, FDR: false discovery rate, HS: Homo sapiens, MM: Mus musculus, IVS: in vitro soil-less, IVV: in vivo, COPD: chronic obstructive pulmonary disease, FLU: fluticasone, SAL: salmeterol, HDM: house dust mite, PBMC: peripheral blood mononuclear cell

Table 4. Clinical characteristics of the 9-gene-set signature expressed in epithelial brushings

Variables[†]	Expressed	Not expressed	p-value
No (%)	N= 31	N=68	
Age (year)	48.0±14.8	44.4±14.0	0.253
Female	12 (38.7%)	40 (58.8%)	0.101
BMI	28.8±5.4	29.0±6.3	0.895
Severe asthma	23 (74.2%)	40 (58.8%)	0.211
Oral corticosteroid use	11 (40.7%)	10 (17.9%)	0.032
Exacerbation number (time/year)	3.0 (2.0-4.0)	2.0 (1.0-3.0)	0.025
Atopy	17 (63.0%)	40 (75.0%)	0.419
Allergic rhinitis	19 (61.3%)	30 (44.1%)	0.171
Eczema	10 (32.3%)	32 (47.1%)	0.245
Nasal polyp	10 (32.3%)	15 (22.1%)	0.404
FEV ₁ (%)	74.2 (57.3-95.8)	87.9 (68.5-95.5)	0.264
Total serum IgE (IU/ml)	184.0 (44.0-373.0)	87.0 (39.6-268.5)	0.067
Blood eosinophil (%)	3.5 (2.7-7.4)	2.3 (1.5-3.8)	0.008
Sputum eosinophil (%)	12.0 (2.6-26.2)	0.6 (0-1.7)	5.2x10 ⁻⁴
Sputum neutrophil (%)	53.2 (42.7-76.9)	52.4 (34.6-63.4)	0.786
FeNO (ppb)	51.5 (20.8-75.8)	21.0 (13.3-37.8)	4.2x10 ⁻⁴
<i>Submucosal cells (mm⁻²)</i>			
Eosinophils	5.8 (3.0-12.5)	1.5 (0-4.8)	1.3x10 ⁻⁵
Neutrophils	11.5 (4.6-17.1)	12.1 (6.8-19.2)	0.780
CD3 T cells	44.7 (32.2-61.7)	24.4 (14.8-46.8)	0.003
CD4 T cells	13.8 (7.9-23.7)	8.3 (3.2-14.6)	0.016
CD8 T cells	24.9 (15.7-37.6)	10.8 (4.2-20.2)	2.0x10 ⁻⁴

†: Data presented as N (%) and mean (SD) or median (IQR).

BMI: Body mass index, FEV₁: Forced expiratory volume in 1 second, FeNO: Fractional exhaled nitric oxide

Table 5. Expression status of the 9 gene-set signature of Cluster A in 83vbronchial biopsy and epithelial brushing

Group	Bronchial biopsy	Epithelial brushing	N (%)
Group 1	Expressed	Expressed	19 (23%)
Group 2	Expressed	Not expressed	17 (20%)
Group 3	Not expressed	Expressed	8 (10%)
Group 4	Not expressed	Not expressed	39 (47%)

Table 6. Clinical features of the expression of 9 gene-set signatures in bronchial biopsy and/or epithelial brush

Variables [†]	Group 1	Group 2	Group 3	Group 4	1 vs 4	3 vs 4	1 vs 3
					<i>p</i> value		
n (%)	19 (23%)	17 (20%)	8 (10%)	39 (47%)			
Age (year)	48.0±13.6	47.4±14.1	50.6±13.4	44.0±13.5	0.284	0.187	0.644
Age of asthma onset	24.7±13.4	22.2±14.4	32.0±11.6	22.1±13.7	0.485	0.039	0.176
Female	7 (36.8%)	8 (47.1%)	3 (37.5%)	24 (61.5%)	0.136	0.390	1.000
BMI	27.8±4.8	28.6±4.9	33.1±3.8	28.0±5.9	0.986	0.010	0.007
Nasal polyps	8 (42.1%)	6 (35.3%)	2 (25.0%)	9 (23.1%)	0.102	0.898	0.357
Rhinitis	8 (42.1%)	8 (47.1%)	6 (75.0%)	16 (41.0%)	0.996	0.100	0.150
Eczema	6 (31.6%)	9 (52.9%)	2 (25.0%)	17 (43.6%)	0.604	0.525	0.900
Severe asthma	14 (73.7%)	10 (58.8%)	6 (75.0%)	20 (51.3%)	0.179	0.402	1.000
Oral corticosteroid use	9 (47.4%)	5 (29.4%)	2 (25.0%)	5 (12.8%)	0.010	0.737	0.515
Atopy	11 (57.9%)	12 (70.6)	6 (75.0%)	30 (76.9%)	0.312	0.476	0.466
Exacerbations/year	3.0 (2.0-5.0)	2.0 (1.3-2.8)	3.0 (2.0-3.0)	1.5 (1.0-3.0)	0.022	0.238	0.563
FEV ₁ (%)	68.1(54.9-78.3)	90.4 (70.9-98.0)	80.3 (65.9-101.6)	79.7(65.5-94.8)	0.133	0.708	0.260
Total IgE (IU/ml)	163 (42–231.5)	86.6 (24.0-196.0)	93.9 (41.3-879.8)	96 (44.3-278.5)	0.659	0.839	0.815
Blood eosinophil (%)	3.5 (2.4-8.0)	1.7 (1.4-3.6)	3.4 (3.2-4.2)	2.4 (1.4-3.9)	0.070	0.100	0.915
Blood neutrophil (%)	58.8 (50.4-64.2)	61.9 (56.6-68.3)	65.5 (61.9-69.5)	58.6 (53.2-68.9)	0.596	0.380	0.217
Sputum eosinophil (%)	8.2 (1.8-27.0)	0.6 (0.6-1.0)	15.8(12.4-16.2)	0.6 (0-1.6)	0.017	0.014	0.863
Sputum neutrophil (%)	52.3 (33.4-77.4)	57.0 (54.2-76.7)	45.3 (41.4-50.8)	52.7 (34.3-65.4)	0.835	0.763	0.727
FeNO (ppb)	56.5 (34.1-74.6)	22.0 (14.5-32.5)	35.5 (20.0-50.4)	19 (12.5-36.1)	0.001	0.058	0.312
<i>Submucosal cells (mm⁻²)</i>							
Eosinophil	6.1 (4.3-13.6)	2.2 (0-5.9)	2.7 (1.5-4.6)	1.0 (0-3.5)	7.6E-06	0.109	0.019
Neutrophil	11.0 (4.6-18.2)	9.8 (7.1-18.7)	12.1 (6.5-13.5)	12.1 (6.8-19.2)	0.853	0.500	0.727
CD4 T cell	15.4 (9.2-24.7)	5.8 (1.5-11.1)	9.2 (6.2-16.3)	10.2 (4.6-15.5)	0.070	0.839	0.187
CD3 T cell	45.3 (32.7-77.2)	21.8 (16.0-37.0)	41.9 (36.9-70.9)	33.3(16.4-49.9)	0.048	0.243	0.874
CD8 T cell	26.4 (15.7-37.0)	9.4 (4.6-14.2)	27.1 (18.1-39.8)	13.9 (4.7-22.1)	0.020	0.150	0.852

†: Data presented as N (%) and mean (SD) or median (IQR). BMI: Body mass index, FEV₁: Forced expiratory volume in 1 second, FeNO: fractional exhaled nitric oxide

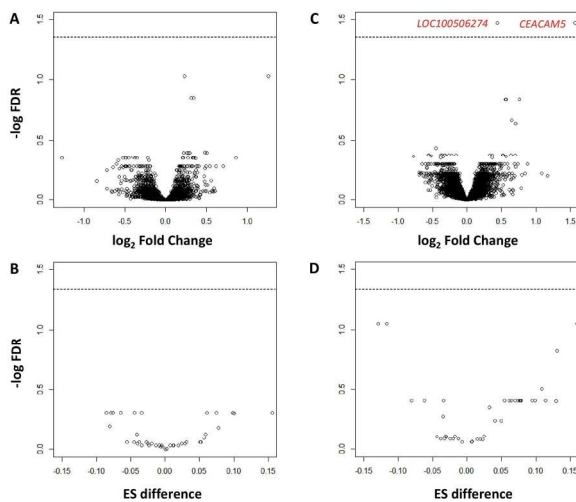


Figure 1

190x107mm (300 x 300 DPI)

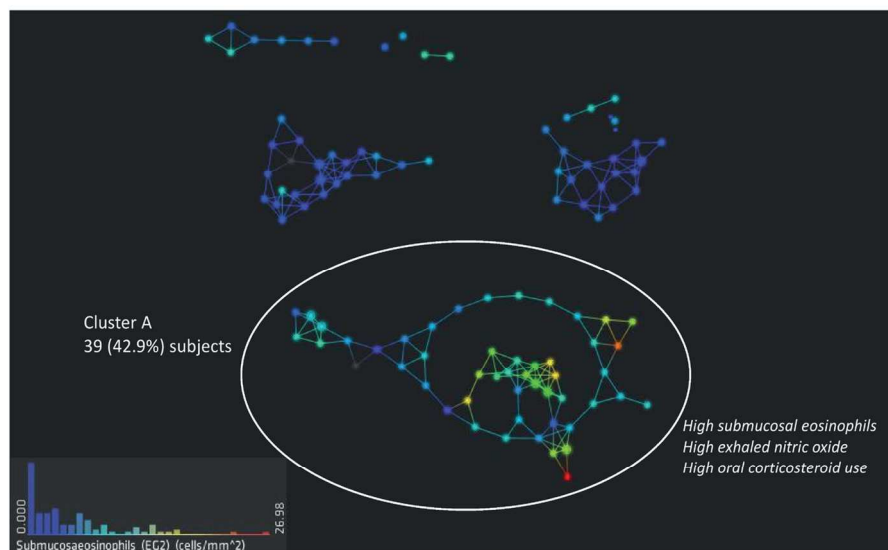


Figure 2

128x76mm (300 x 300 DPI)

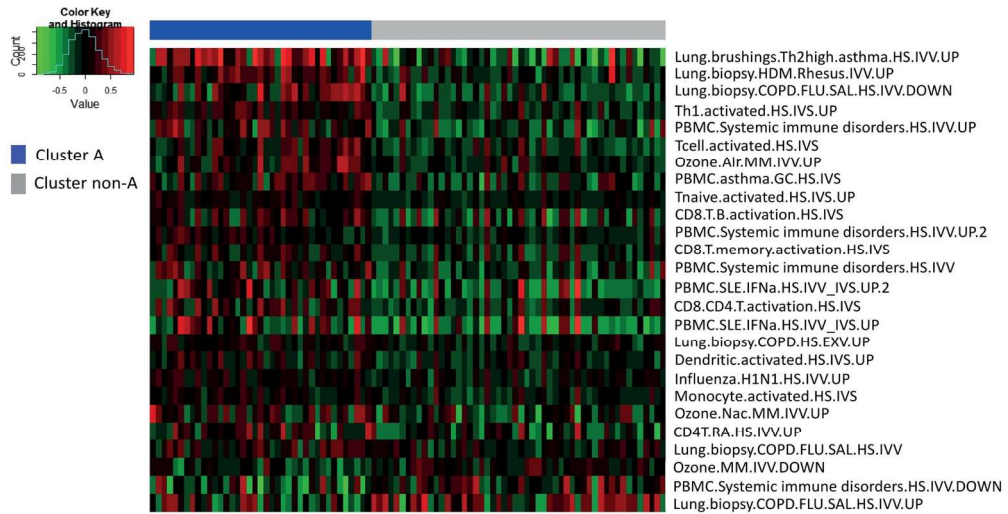


Figure 3

157x85mm (300 x 300 DPI)

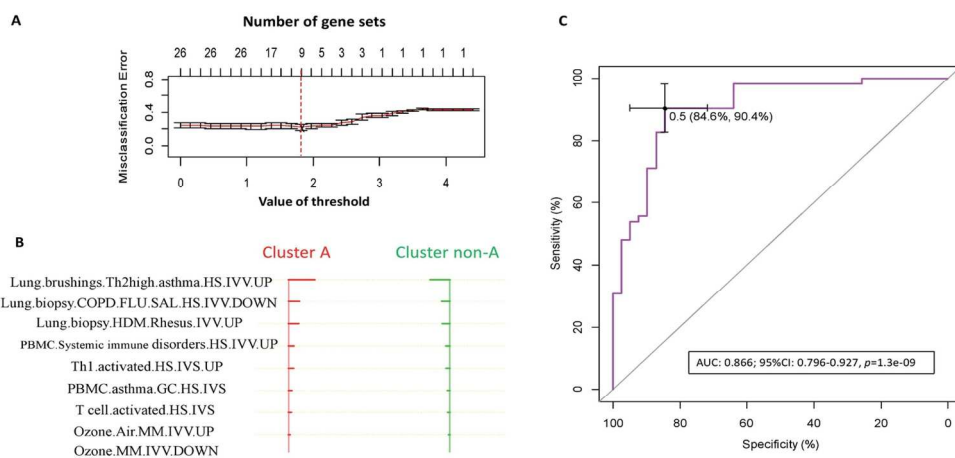


Figure 4

134x70mm (300 x 300 DPI)

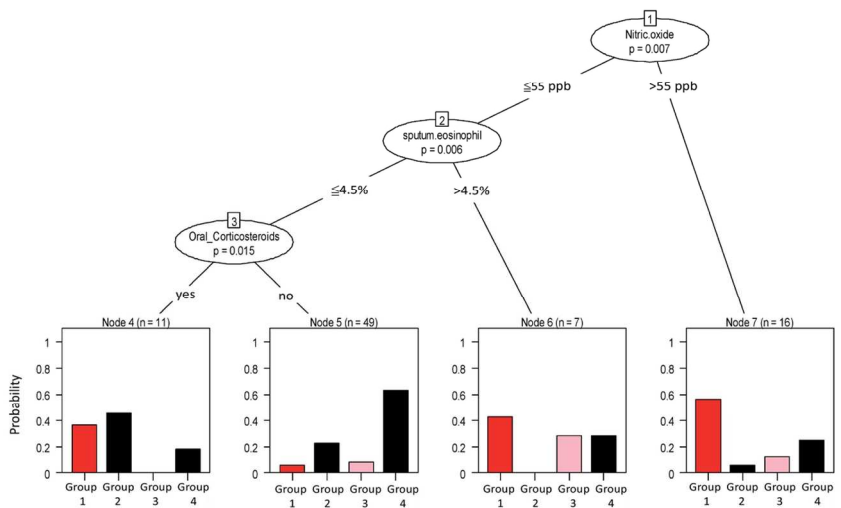


Figure 5

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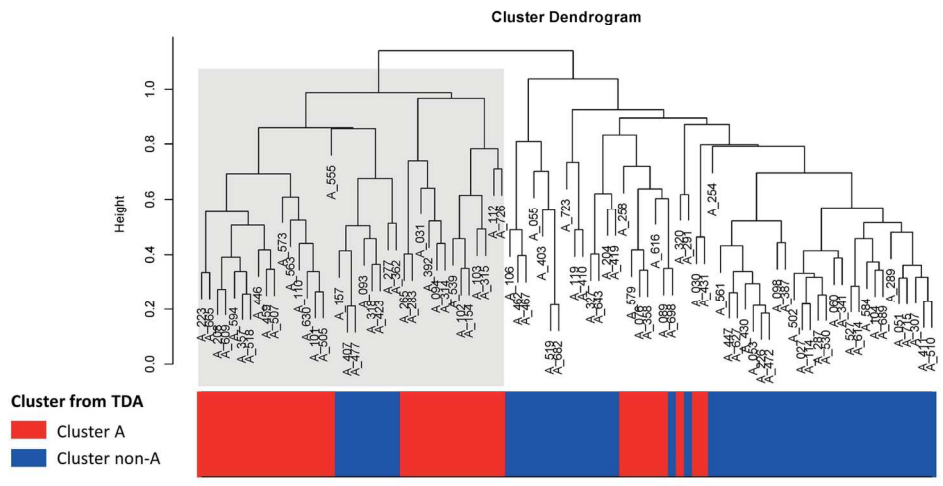


Figure 6

138x75mm (300 x 300 DPI)

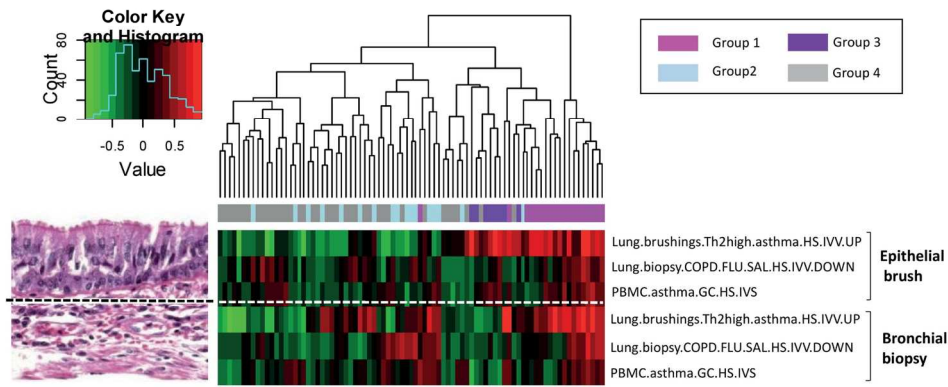


Figure 7

153x85mm (300 x 300 DPI)

Supplementary data

A transcriptomic-driven analysis of epithelial brushings and bronchial biopsies to define asthma phenotypes in UBIOPRED

Chih-Hsi Scott Kuo, Stelios Pavlidis, Matthew Loza, Fred Baribaud, Anthony Rowe, Ioannis Pandis, Uruj Hoda, Christos Rossios, Ana Sousa, Susan J. Wilson, Peter Howarth, Barbro Dahlen, Sven-Eric Dahlen, Pascal Chanez, Dominic Shaw, Norbert Krug, Thomas Sandström, Bertrand De Muelder, Charles Auffray, Peter J. Sterk, Ratko Djukanovic, Yike Guo, Ian M. Adcock & Kian Fan Chung on behalf of the U-BIOPRED Project Team

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Methods for Cluster Analysis and Machine Learning Algorithm

Topology Data Analysis

Software used for TDA was provided by Ayasdi Core™ (Ayasdi, Inc. California, USA). TDA method takes three clustering parameters as input, namely *dissimilarity (distance) metric*, *filter function* and *resolution parameters* (resolution and overlap). A user-defined distance function computes the distance between any pairs of the data points and thus determines a metric space of the data set that serves as the starting point of TDA. The distance metric using correlation distance was a consensus setting in the UBIOPRED project across each individual analysis when applying the TDA approach for clustering. The filter function in TDA serves to generate a real valued quantity associated with each data point that reflects a unique view of the data set. There are various sort of filter functions in TDA that include functions that depend only on the distance metric itself, such as a density estimator or a measure of centrality; functions that produce linear projections on a data matrix, such as principle component analysis or multidimensional scaling; or functions that retrieved from input parameters by researchers, such as clinical metadata. In UBIOPRED, the filter function was an open option across each analysis. In this analysis, filter function of linear projection was chosen as the 1st and 2nd principle component of the 42 GSVA signatures in order to unbiasedly focus on the major variation within the data matrix.

The value of each data point computed by the filter function which spanned from low to high was segmented into overlapping intervals (bins). User-defined resolution parameters in TDA are used to control the size of the bins, a high resolution means fine-grained partitions and a low resolution suggests coarse-grained ones. A network in high resolution will preserve the strongest connections between data points and data points that are more weakly associated tend to break apart as the bins are split. An overlap parameter is used to define the extent to which the data will be

oversampled. This parameter allows users to set the number of bins that are shared by most data points thus a higher overlap parameter is often useful to preserve the relationship of continuity within a data set. In this analysis, we set a moderate-high resolution (Resolution=30) and overlap parameter (Gain=3x) in order to reveal the detailed structure of the network while not compromising the inherent continuity of the data. Using these input set up, data points having their filter function value sat in the same bin were combined when clustering was performed based on the correlation dissimilarity. By clustering similar data points into nodes and joining nodes that had shared data points with edges, TDA produced a coordinate-free network with 3D graphic visualization to enable cluster exploration. Subsequent analysis of these graphs is used to identify specific network features such as flares or separated clusters that represent subgroups within the data set. Ayasdi CoreTM also provides a histogram-based coloring scheme for the constructed network from all available filter functions or metadata which was input by users. This flexible coloring scheme enables a quick overview of the characteristic of interest for all identified clusters or subgroups within the network [1, 2].

Nearest Shrunken Centroid learning algorithm

The nearest shrunken centroid method was used as a machine learning algorithm to further define the optimal number of GSVAs signatures that best differentiated between identified clusters. The centroids (average expression of each GSVAs signature) for each cluster as well as the overall samples were calculated. Standardization of the centroids of each cluster was performed through dividing the difference of the cluster centroids and overall centroids by the within-cluster standard deviation of each signature. This standardized value was treated as an absolute value which was later shrunken by an amount (Δ , threshold value). If the value of a given standardized

centroid was shrunken to zero by Δ for all clusters, then this signature did not contribute to the nearest-centroid classification. Otherwise, a non-zero value of a standardized centroid after shrinkage was retained as a centroid classifier for the given cluster. The inherent property of this method is that many signatures will be eliminated from cluster prediction as Δ increases. Given that each amount of Δ shrunken will result in a set of surviving centroids for each cluster, the amount of shrinkage is chosen by iterative cross-validation on the performance that each set of surviving centroids correctly predicts the cluster classification of each sample. This algorithm is available as a free R software as prediction analysis for microarrays in The Comprehensive R Archive Networks (CRAN-pamr package) [3].

Table S1. 42 signatures used for gene set variation analysis

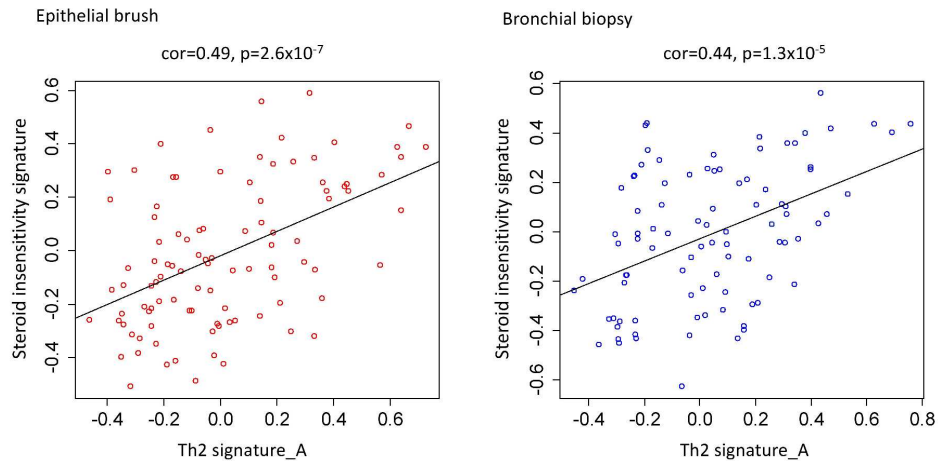
Name of gene-sets	Disease/Treatment	Species	Cell Type	Study Type	Reference
<i>Omnibus immune cells activation</i>					
Tcell.activated.HS.IVS	activated	HS	Tcell	IVS	[4]
Bcell.activated.HS.IVS	activated	HS	Bcell	IVS	[4]
Monocyte.activated.HS.IVS	activated	HS	Monocyte	IVS	[4]
NKcell.activated.HS.IVS	activated	HS	NKcell	IVS	[4]
Dendritic.activated.HS.IVS	activated	HS	Dendritic	IVS	[4]
Neutrophil.activated.HS.IVS	activated	HS	Neutrophil	IVS	[4]
<i>Th1 and CD8 T cell immunity</i>					
CD8.T.memory.activation.HS.IVS	activated	HS	peripheral blood	IVS	[5]
CD8.T.B.activation.HS.IVS	activated	HS	peripheral blood	IVS	[5]
CD8.CD4.T.activation.HS.IVS	activated	HS	peripheral blood	IVS	[5]
Th1.activated.HS.IVS.UP	activated	HS	peripheral blood	IVS	[6]
Th1.activated.HS.IVS.UP.2	activated	HS	peripheral blood	IVS	[7]
Influenza.H1N1.HS.IVV.UP	H1N1trivalent influenza vaccine	HS	peripheral blood	IVV	[8]
<i>Th2, Th17 and T-reg cell immunity</i>					
Th17.activated.HS.IVS.UP	activated	HS	peripheral blood	IVS	[6]
Th17.activated.HS.IVS.UP.2	activated	HS	peripheral blood	IVS	[7]
Th2.activated.HS.IVS.UP	activated	HS	peripheral blood	IVS	[7]
Treg.activated.HS.IVS.UP	activated	HS	peripheral blood	IVS	[7]
Tnaive.activated.HS.IVS.UP	activated	HS	peripheral blood	IVS	[7]
<i>Oxidative stress</i>					
Ozone.MM.IVV.Down	Ozone	MM	lung biopsy	IVV	#
Ozone.Air.MM.IVV.UP	Ozone followed by air	MM	lung biopsy	IVV	#
Ozone.Air.MM.IVV.DOWN	Ozone followed by air	MM	lung biopsy	IVV	#
Ozone.Nac.MM.IVV.UP	Ozone followed by N-acetylcysteine	MM	lung biopsy	IVV	#
Smoking.signature.Mouse	smoke	MM	lung biopsy	IVV	#
Lung.biopsy.COPD.HS.EXV.UP	COPD	HS	lung biopsy	EXVIVO	[9]
<i>Systemic immune disorders</i>					
PBMC.Systemic immune disorders.HS.IVV	4 autoimmune diseases	HS	peripheral blood	IVV	[10]
PBMC.Systemic immune	4 autoimmune	HS	peripheral blood	IVV	[10]

disorders.HS.IVV.UP	diseases				
PBMC.Systemic immune disorders.HS.IVV.DOWN	4 autoimmune diseases	HS	peripheral blood	IVV	[10]
PBMC.Systemic immune disorders.HS.IVV.2	3 autoimmune diseases	HS	peripheral blood	IVV	[10]
PBMC.SLE.IFNa.HS.IVV_IVS.UP	SLE (Bonferroni)	HS	peripheral blood	IVV_IVS	[11]
PBMC.SLE.IFNa.HS.IVV_IVS.UP.2	SLE (FDR)	HS	peripheral blood	IVV_IVS	[11]
CD4T.RA.HS.IVV.UP	RA	HS	peripheral blood	IVV	[12]
Asthma specific T cell immunity					
CD4.severe.asthma.HS.IVV.UP	Severe Asthma	HS	peripheral blood	IVV	[13]
CD4.severe.asthma.HS.IVV.UP	Severe Asthma	HS	peripheral blood	IVV	[13]
CD8.severe.asthma.HS.IVV.UP	Severe Asthma	HS	peripheral blood	IVV	[13]
CD8.severe.asthma.HS.IVV.DOWN	Severe Asthma	HS	peripheral blood	IVV	[13]
Lung.biopsy.HDM.Rhesus.IVV.UP	HDM	Rhesus	lung biopsy	IVV	[14]
Lung.biopsy.Th2high.asthma.HS.IVV.U P	Th2 high/low asthma	HS	lung biopsy	IVV	[15]
Lung.brushings.Th2high.asthma.HS.IVV .UP	Th2 high/low asthma	HS	lung brushings	IVV	[16]
Glucocorticoid response					
Lung.biopsy.COPD.FLU.SAL.HS.IVV	COPD fluticasone & salmeterol	HS	lung biopsy	IVV	[17]
Lung.biopsy.COPD.FLU.SAL.HS.IVV. UP	COPD fluticasone & salmeterol	HS	lung biopsy	IVV	[17]
Lung.biopsy.COPD.FLU.SAL.HS.IVV. DOWN	COPD.fluticasone & salmeterol	HS	lung biopsy	IVV	[17]
Lung.ASM.asthma.Prednisolone.HS.IV V.UP	Asthma, Prednisolone	HS	lung biopsy	IVV	[18]
PBMC.asthma.GC.HS.IVS	Asthma,IL-1 β , TNF- α , with or without GC	HS	peripheral blood	IVS	[19]

HS: Homo sapiens, MM: Mus musculus, IVS: in vitro soil-less, IVV: in vivo, INF: interferon, COPD: chronic obstructive pulmonary disease, FLU: fluticasone, SAL: salmeterol, HDM: house dust mite, PBMC: peripheral blood mononuclear cell, SLE: systemic lupus erythematosus, RA: rheumatoid arthritis, GC: glucocorticoid, #: unpublished data from Philip Hansbro et al.

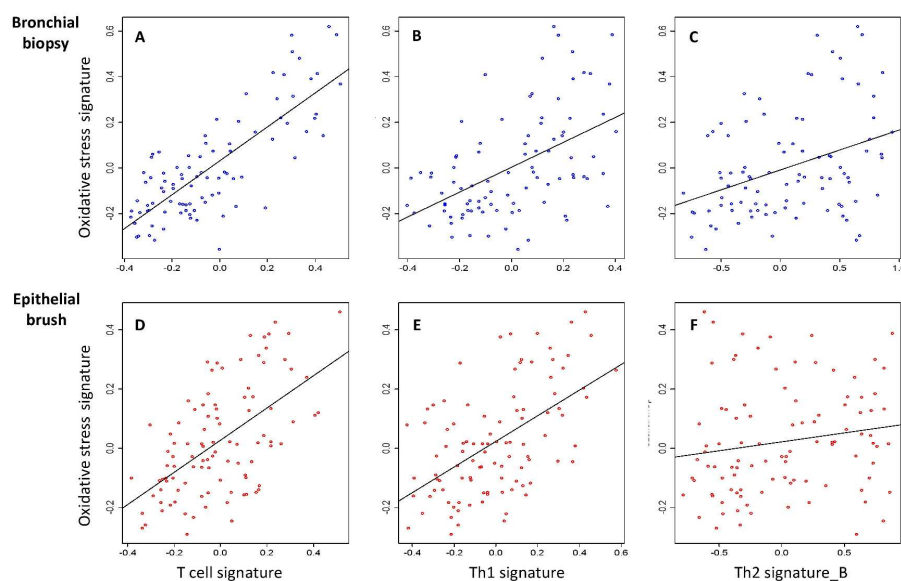
Supplementary Figures

Figure S1



Relationship between steroid insensitivity (PBMC.asthma.GC.HS.IVS) and Th2 signature-A (Lung.biopsy.HDM.Rhesus.IVV.UP) showing a moderate correlation in epithelial brushings (Pearson's r : 0.49, $p=2.6 \times 10^{-7}$) and bronchial biopsies (Pearson's r : 0.44, $p=1.3 \times 10^{-5}$). There was also a significant correlation between the steroid insensitivity signature and the Th2 signature-B (Lung.brushings.Th2high.asthma.HS.IVV.UP) in epithelial brushings and bronchial biopsies (data not shown).

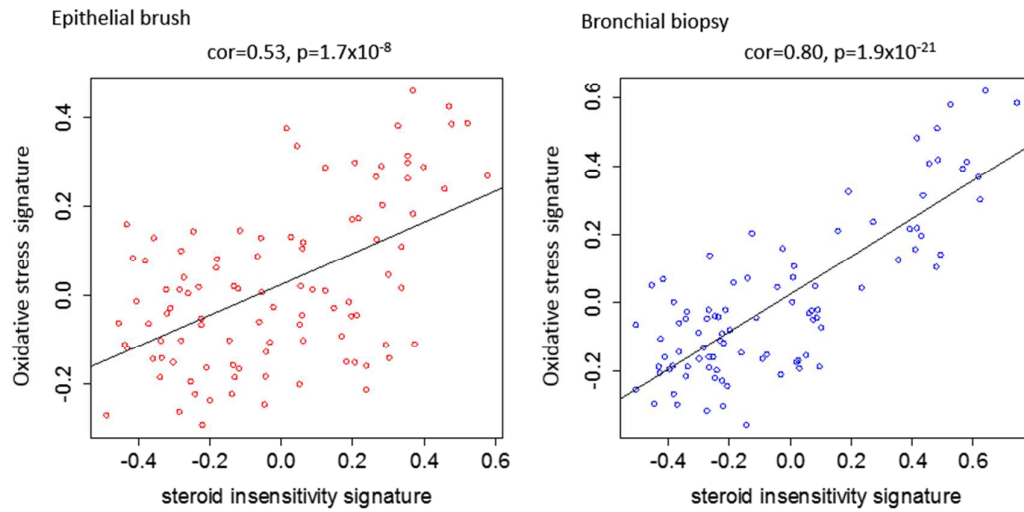
Figure S2



Relationship between oxidative stress (Ozone.Air.MM.IVV.UP) and adaptive immunity signatures. In bronchial biopsy, oxidative stress showed high correlation with (A) T cell signature (Tcell.activated.HS.IVS; Pearson's r : 0.79, $p=1.3 \times 10^{-20}$), moderate correlation with (B) Th1 signature (Th1.activated.HS.IVS.UP; Pearson's r : 0.50, $p=4.5 \times 10^{-7}$) and mild correlation with (C) Th2 signature_B (Lung.brushings.Th2high.asthma.HS.IVV.UP; Pearson's r : 0.36, $p=4.6 \times 10^{-7}$). In epithelial brushing, oxidative stress showed moderate correlation with (D) T cell signature (Tcell.activated.HS.IVS; Pearson's r : 0.58, $p=3.1 \times 10^{-10}$), moderate correlation with (E) Th1 signature (Th1.activated.HS.IVS.UP; Pearson's r : 0.55, $p=3.7 \times 10^{-9}$) and weak to negligible correlation with (F) Th2 signature_B (Lung.brushings.Th2high.asthma.HS.IVV.UP; Pearson's r : 0.16, $p=0.110$). There was also a significant correlation between the oxidative stress signature and the Th2

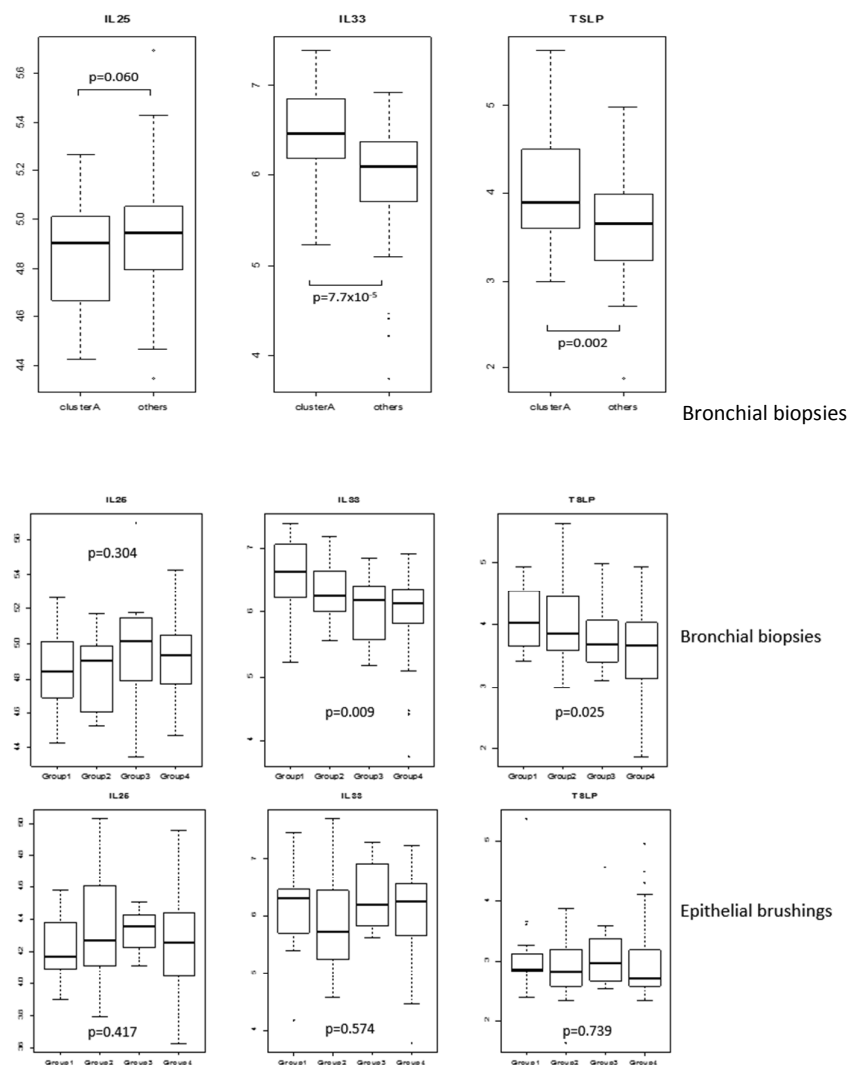
signature-A (Lung.biopsy.HDM.Rhesus.IVV.UP) in bronchial biopsies and epithelial brushings (data not shown).

Figure S3



Relationship between steroid insensitivity (Lung.biopsy.COPD.FLU.SAL.HS.IVV.DOWN) and oxidative stress (Ozone.Air.MM.IVV.UP) signature showing a moderate correlation in epithelial brushings (Pearson's r : 0.53, $p=1.7 \times 10^{-8}$) and high correlation in bronchial biopsies (Pearson's r : 0.80, $p=1.9 \times 10^{-21}$).

Figure S4



Gene expression of IL-25, IL-33 and TSLP in bronchial biopsies according to cluster A and non-A (upper panel), and in bronchial biopsies and epithelial brushings according to the 4 Groups (lower panels). For bronchial biopsies, there was a significant difference in the level of gene expression for IL-33 and TSLP between Group1 and Group4 ($p=0.003$ for both) and between Group2 and Group4 ($p=0.014$ and $p=0.051$, respectively). There was a difference in the level of gene expression of IL-33 between Group1 and Group3 ($p=0.053$).

Reference:

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