Bioinformatic identification and analysis of immune-related chromatin regulatory genes as potential biomarkers in idiopathic pulmonary fibrosis

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Background: Idiopathic pulmonary fibrosis (IPF) is a disease with a very poor prognosis. The search for new IPF biomarkers is particularly urgent due to the uncertainty of the mechanisms and treatment. Studies have shown that chromatin regulators (CRs) are involved in the development of IPF and are associated with tumor immunity. However, there are no studies on immune-related CRs in IPF. Therefore, we conducted a systematic study to analyze the expression levels and immune correlation of CRs in IPF tissues and normal tissues and to explore their potential as diagnostic biomarkers.

Methods: GSE53845, GSE179781 and GSE24206 datasets from Gene Expression Omnibus (GEO) database were merged into an integrated dataset as the training set; GSE70866 was used as the validation dataset. The cr-related differentially expressed genes (DEGs) between normal and IPF tissues were identified using the “Limma” software package. Weighted gene co-expression network analysis (WGCNA) was performed using the “WGCNA” package to screen eigengenes, which were intersected with DEGs to identify hub genes. The “ggcorrplot” package was used to analyze the correlation between hub genes and immunity, and immune-related hub genes were defined as immHub. A logistic regression model was constructed using immHub as the independent variable and whether the diagnosis was IPF as the dependent variable.

Results: One hundred and sixty-nine DEGs were identified between IPF and normal tissues. wGCNA identified 3 key modules in brown, green and yellow genes that were present in all 3 modules and met module membership (MM) >0.8 and gene significance (GS) >0.5 were called signature genes (n=390). Four intersecting genes were obtained by intersecting DEGs with signature genes (PADI4, IGFBP7, GADD45A, and SETBP1) all associated with immunity were defined as immHub genes Logistic regression models were constructed based on immHub genes. The area under the curve (AUC) of the ROC curve is used to evaluate the diagnostic accuracy of the logistic regression model for IPF. The AUC in the ROC analysis was 0.771 for the training dataset, and 0.759 for the validation dataset.

Conclusions: PADI4, IGFBP7 and GADD45A may be biomarkers for IPF, which will provide assistance in the diagnosis, treatment and prognostic assessment of IPF patients, and provide an important basis for future studies on the relationship between CRs genes and IPF.

Keywords: Chromatin regulators (CRs); idiopathic pulmonary fibrosis (IPF); immunity

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Introduction

Idiopathic pulmonary fibrosis (IPF) is a chronic, progressive, and fibrotic interstitial lung disease with an extremely poor prognosis, having a median survival of only 2–3 months after diagnosis (1). Only two drugs have been approved by the US Food and Drug Administration (FDA) for treatment of IPF (pirfenidone and nintedanib), but neither can prevent or reverse the disease progression, only slow down the deterioration in lung function. Their common side effects include nausea, fatigue, and diarrhea, which are often difficult to tolerate. In selected IPF patients, lung transplantation may be the only treatment option (2-4). At present, the pathogenesis of IPF is still unclear but has been well elucidated from the perspective of epigenetics (5). Chromatin regulators (CRs) are indispensable upstream regulatory factors of epigenetics and can modulate the underlying genes of cancer and other diseases (6). According to their modulation mode, CRs can be divided into three categories: DNA methylators, histone modifiers, and chromatin remodelers (7). Current studies have reported that CRs can affect the progress of IPF by modulating the proliferation and differentiation of alveolar myofibroblasts and regulating apoptosis (8-10).

DNA methylation is a chemical modification of the DNA structure. With S-adenosyl methionine as the methyl donor, it is mediated by DNA methyltransferases to complete the covalent bonding with specific bases in the DNA sequence. In eukaryotes, DNA methylation mainly exists in CpG islands in gene promoter regions and exons, primarily involving cytosine nucleotides, and the expression and silencing of its methylation can regulate the expressions of related genes (11). DNA methylation is associated with the occurrence of IPF, and DNA methylation in IPF may affect the expressions of specific genes in the lungs of IPF patients (12,13). In addition, it can regulate IPF-related transforming growth factor-beta (TGF-β) and Wnt signaling pathways (14). Through methylation array data analysis, Evans et al. (14) found that the transcriptional regulator gene C8orf4 was downregulated in fibrotic fibroblasts, and its three CpG islands in the promoter region were hypermethylated. The knockdown of C8orf4 in lung fibroblasts by small interfering RNA revealed that the levels of cyclooxygenase-2 (COX-2) and prostaglandin E2 (PGE2) were downregulated. Therefore, it is believed that downregulation of COX-2 and PGE2 levels in lung fibroblasts is related to the hypermethylation of C8orf4, which leads to fiber growth in the lungs. DNA methylation can also lead to hyperproliferation of fibroblasts and an altered differentiation phenotype of airway macrophages (15).

Histones are a family of basic proteins that associate with DNA in the nucleus of eukaryotic cells. There are five histone types, including H1, H2A, H2B, H3, and H4. Histones assemble to form a nucleosome together with DNA, thereby packing it tight inside the nucleus. Covalent modifications (phosphorylation, methylation, acetylation, and ubiquitination) at the amino terminus of core histones in nucleosomes play an important role in gene regulation, and most of the histone modifications associated with IPF are acetylation, methylation, and ubiquitination (16-18).

Chromatin remodeling refers to the weakening of nucleosome-DNA binding and aggregation between adjacent nucleosomes through physical ATP-dependent chromatin modifications or chemical covalent modifications. Different chromatin remodeling factors can affect epithelial-mesenchymal transition (EMT) processes in multiple organs (19). Obrien et al. exposed non-Brg1 (Brahma-related gene 1) mutant wild-type cell line H441 and Brg1-mutant cell line A549 to TGF-β and found that both cell lines showed a myofibroblast-like phenotype; however, H441 had more obvious upregulation of epithelial cell markers (e.g., E-cadherin) and downregulation of fibroblast markers (e.g., vimentin). In addition, the upregulation of TGF-β target genes (such as PAI-1 and JUN) was more pronounced in H441 than in A549. Therefore, the chromatin remodeling protein Brg1 mediates the EMT process in IPF by affecting the TGF-β signaling pathway (20).

Thus, CRs are closely related to the occurrence of IPF. The recent decades have witnessed dramatic advances in research on the immune system. It has been found that immune dysregulation is a driver of IPF (21). M2 macrophages, Th17 cells, CD8+ T cells, and possibly regulatory T-cells (Tregs) promote fibrosis, while Th1 and TRM CD4+ T cells appear to be protective. Targeted immunomodulators show promise as IPF treatment (22).

Epigenetic dysregulation is an important feature of immune escape in tumorigenesis, and CRs have cell-intrinsic effects on the immune sensitivity of cancer cells (23). Because the immune response is closely related to CRs, there are no functional studies of immune-related CRs in IPF. The aim of this study is to screen immune-related CRs as biomarkers of IPF for diagnosis, treatment and prognostic assessment of IPF. It also provides a basis for further studies on the function of CRs in IPF. We present the following article in accordance with the TRIPOD reporting checklist (available at https://atm.amegroups.com/article/view/10.21037/atm-22-3700/rc).
Methods

Data sources

All the data were downloaded from the Gene Expression Omnibus (GEO) database (https://www.ncbi.nlm.nih.gov/geo/). Data on normal and IPF tissues were collected from four datasets (GSE53845, GSE17978, GSE24206, and GSE70866) on May 1, 2022. Three (GSE53845, GSE17978, and GSE24206) were combined into an integrated dataset to be used as the training dataset, and GSE70866 was used as a validation dataset. Data from the three datasets were merged and normalized using the “SVA” R package. The study was conducted in accordance with the Declaration of Helsinki (as revised in 2013).

Identification of CRs and screening of differentially expressed genes (DEGs)

We identified 870 CR-related genes from the appendix of a literature review (24). We used the “Limma” R package to screen DEGs (n=169), with |log2 Fold Change| >1 and adjusted P value <0.05 as the cut-off values. Volcano plots and heat maps of the DEGs were drawn. Gene Ontology (GO) and Kyoto Encyclopedia of Genes and Genomes (KEGG) pathway enrichment analyses of the DEGs were performed with the “clusterProfiler” package. A protein-protein interaction (PPI) network of DEGs was constructed and visualized by the STRING database (with a minimum required interaction score of 0.9) and Cytoscape 3.8.2 software, respectively.

Identification of hub genes on WGCNA

Weighted gene coexpression network analysis (WGCNA) of all genes in IPF and normal tissues was performed using the “WGCNA” R package. Low-expression genes were filtered, and samples were clustered to remove outliers as abnormal values. An adjacency matrix was constructed to map the association between the graph nodes. The pickSoftThreshold function was used to select the optimal value for the weighting parameters of adjacent functions, with a soft threshold β=3 (scale-free R2=0.9). The modules of related genes were constructed based on hierarchical clustering of the dissimilarity measure (1-TOM) of the topological overlap matrix (TOM). The minimum number of genes in modules was set to 60 and the threshold for merging similar modules was set to 0.25. The coexpression module is a collection of genes with high topological overlap similarity, and genes in the same module often have a higher degree of coexpression. Module eigengene is defined as the first principal component of the expression matrix of the corresponding module and is used to describe the expression pattern of the module in each sample. The key modules (yellow, green, and brown) were screened for analysis. The inclusive criterion for identifying eigengenes (n=390) in the key modules was as follows: module membership (MM) >0.8 and gene significance (GS) >0.5. MM refers to the correlation coefficient between genes and module eigengenes, which is used to describe the reliability of a gene belonging to a module. GS values reflect the correlation between gene expression levels.

Construction of a diagnostic model based on CRs

Four genes (PADI4, IGFBP7, GADD45A, and SETBP1) were obtained by overlapping and intersecting the DEGs and eigengenes. The “ggstatsplot” package was used to perform Spearman correlation analysis on diagnostic markers and infiltrating immune cells and the “ggplot2” package was used to visualize the results, which showed that PADI4, IGFBP7, and GADD45A correlated with immunity. A logistic regression model was constructed with PADI4, IGFBP7, and GADD45A as joint predictors. The sample information of normal and IPF tissues in the GSE53845, GSE17978, and GSE24206 datasets was extracted as the training sets, and the receiver operating characteristic (ROC) curves were drawn using the “ROC” R package. Area under the ROC curve (AUC) was used to assess the accuracy of the regression model for differential diagnosis. GSE70866 was used as the validation dataset. Nomograms were drawn for various outcomes with the “nomogram()” function in R.

Analysis of immune activity

Single-sample gene set enrichment analysis (ssGSEA) was performed using the “gsva” package to calculate the score of infiltrating immune cells and to assess immune function. Heatmaps of the expressions of immune cells and immune function in each sample were drawn using the “heatmap” package. Heatmaps reflecting the correlations between immune cells and immune function were drawn using the “corrplot” package. The “vioplot” package was used to visualize and compare the levels of immune cells in IPF and normal tissues.
Statistical analysis

R software (version 4.2.0) was used for statistical analysis. The R packages applied in each step were downloaded from the Bioconductor website (https://bioconductor.org/). All reported P values are two-tailed. P<0.05 was set as the threshold for statistical significance.

Results

Identification of CR DEGs between normal and IPF tissues

Among the three GEO datasets (GSE53845, GSE17978, and GSE24206) selected, GSE53845 contained 8 normal lung tissue samples and 40 IPF tissue samples, GSE17978 contained 20 normal tissue samples and 38 IPF tissue samples, and GSE24206 contained 6 normal lung tissue samples and 17 IPF tissue samples. By comparing the expression levels of CR-related genes in the 34 normal tissues and 95 IPF tissues, we screened 169 CR DEGs (P<0.05, |log2FC| ≥ 0.2), of which 88 genes were upregulated and 81 were downregulated. Volcano plots and heat maps of these DEGs were drawn (Figures 1,2). To further analyze the interactions among the CR DEGs, a PPI network with 169 nodes and 194 edges was constructed using the highest stringent minimum required interaction score of 0.9 by STRING (Figure 3). KEGG enrichment analysis (Figure 4) showed that DEGs were mainly enriched in the cell cycle, p53 signaling pathway, viral life cycle - HIV-1, Notch signaling pathway, transcription coregulator activity, and other signaling pathways. GO enrichment analysis, which depicts three complementary biological concepts including biological process (BP), molecular function (MF) and cellular component (CC), showed the top 5 enriched terms. BP enrichment showed that DEGs were mainly involved in histone modification, chromatin organization, chromatin remodeling, peptidyl-lysine modification, and peptidyl-lysine acetylation; GO:0018393 internal peptidyl-lysine acetylation; the most enriched CC included SWI/SNF superfamily-type complex, ATPase complex, histone deacetylase complex, histone acetyltransferase complex, and histone methyltransferase complex; and the top 5 enriched MF terms included histone binding, transcriptional coregulator activity, nucleosome binding, modification-dependent protein binding, and transcriptional coactivator activity (Figure 5).

Gene coexpression network and identification of key modules and hub genes

After data preprocessing, a gene expression matrix (14,725 genes) was generated after the three databases (GSE53845, GSE17978, and GSE24206) were merged. The missing values of the samples were checked, and cluster analysis was performed to remove outliers and abnormal samples. The modules of related genes in modules were constructed based on hierarchical clustering of the dissimilarity measure.
Figure 3 Protein-protein interaction network based on DEGs between normal and idiopathic pulmonary fibrosis lung tissues. DEGs, differentially expressed genes.

Figure 4 Bar chart (left) and bubble chart (right) of KEGG analysis. Longer bars (left) or larger bubbles (right) represents more gene enrichment and deeper red color represents more significant difference. KEGG, Kyoto Encyclopedia of Genes and Genomes.

(1-TOM) of the TOM. The minimum number of genes in modules was set to 60 and the threshold for merging similar modules was set to 0.25. A total of 9 modules were yielded. Genes in the same module had high connectivity and similar functions (Figure 6). Each module was assigned a unique color identifier and an expression profile summarized by module eigengenes (MEs) (Figure 7). The ME of the brown modules negatively correlated with IPF (r=-0.52, P=2^{-10}), and the MEs of the green modules (r=0.41, P=2^{-6}) and yellow modules (r=0.75, P=4^{-24}) positively correlated with IPF. The brown, green, and yellow modules were selected as key modules for further analyses. Phenotypic data, and correlation graphs were used to show the correlation between the MM and GS of IPF-related modules (Figures 8-10). Genes with MM >0.8 and GS >0.5 in the three modules were selected, and the eligible genes in the
three key modules were merged into the eigengenes of IPF (n=390).

**Identification and validation of immune-related CR genes**

Four hub genes (PADI4, IGFBP7, GADD45A, and SETBP1) were obtained by overlapping and intersecting the DEGs and eigengenes screened by WGCNA (Figure 11). The relationships of these four hub genes with immune cells and immune function were analyzed using the “ggcorrplot” R package, which showed that PADI4, IGFBP7, and GADD45A correlated with immunity (Figure 12). Analysis of the potential correlation between relevant biomarkers and infiltrating immune cells showed that: GADD45A positively correlated with APC_co_stimulation, CCR, parainflammation, and Treg; IGFBP7 negatively correlated with APC_co_inhibition and positively correlated with Mast_cells; and PADI4 negatively correlated with HLA, Macrophages, and Mast_cells and positively correlated with Neutrophils. A logistic regression model was constructed based on the three immHub genes (PADI4, IGFBP7 and GADD45A), with whether the tissue was IPF as the dependent variable and the levels of PADI4, IGFBP7, and GADD45A as the dependent variables. PADI4, IGFBP7, and GADD45A were divided into high- and low-expression groups according to their median values. The levels of PADI4 [odds ratio (OR) =0.2498; 95% confidence interval (CI): 0.092–0.629], IGFBP7 (OR =2.7784; 95% CI:
and $GADD45A$ (OR $=3.0641$; 95% CI: 1.18–8.435) correlated with the occurrence of IPF (all $P<0.05$). According to the prediction formula $[P=1/(1+\exp(-Z))]$ of the regression model, the prediction formula of our current study was: $P=1/(1 + \exp[-(1.022 \times IGFBP7 - 1.387 \times PADI4 + 1.120 \times GADD45A)])$. For the training dataset, the AUC was used as the assessment parameter, which was 0.771. Nomograms were drawn for various outcomes (Figures 13,14).

Validating model performance in the validation dataset. In the validation dataset, the AUC was 0.759. Nomograms are shown in Figures 15,16.

The ROC is used to determine the differential diagnostic accuracy of the i.e. logistic regression model, with the closer
the AUC value to 1 the more effective the model is. We believe that models with AUC between 0.5 and 0.6 have low diagnostic accuracy, models with AUC between 0.6 and 0.7 have some diagnostic accuracy, models with AUC between 0.7 and 0.8 have high accuracy, and models with AUC >0.9 have excellent diagnostic performance.

The AUCs of the models ranged from 0.7 to 0.8 in the training and validation datasets, indicating that the model had good predictive ability.

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Immune activities of IPF and normal tissues

Heatmaps were constructed to display the expressions of immune cells and immune function in each sample (Figure 17). Analyses of the enrichment scores of 16 immune cells in IPF and normal tissues and the activities of 13 immune function showed that, among intestinal mucosal mast cells (IMMCs), B_cells, CD8+ T_cells, DCs, iDCs, Macrophages, Mast_cells, T_helper_cells Tfh, and Th2_ cells were highly expressed in IPF whereas NK_cells and Neutrophils were lowly expressed in IPF. In IMMF, APC_co-stimulation, Check-point, HLA, Inflammation-promoting, MHC_class_I, Parainflammation, T_cell_co-inhibition, and T_cell_co-stimulation were all highly expressed in IPF tissues (Figure 18). Analysis for the potential correlations between immune cells and immune function showed that immune cell Tumor Infiltrating lymphocytes positively correlated with B_cells (r=0.75) and Neutrophils negatively correlated with Mast_cells (r=–0.38). Analysis for the potential correlations among immune function showed that Check-point positively correlated with T_cell_co-stimulation (r=0.88) and Type_II_IFN_Response negatively correlated with HLA (r=–0.06) (Figure 19).

Prediction of model-related gene therapy drugs

Through the Enrichr website (https://maayanlab.cloud/Enrichr/), some small-molecule compounds that may be used in the treatment of IPF were obtained, and the top 10 drugs are listed (Table 1).

Discussion

CRs are a class of enzymes with special functional structures that can recognize, form, and maintain epigenetic states in a cell-environment-dependent manner. CRs are indispensable upstream regulators of epigenetics, and their abnormal expressions are closely related to various BPs. Research has shown that immune dysregulation is the main driver of IPF pathogenesis, and CRs can help tumor immune escape. We constructed a logistic regression model based on three immune-associated CR genes (PADI4, IGFBP7, and GADD45A). The AUC was 0.771 in the training dataset and 0.759 in the validation dataset, indicating our model had good predictive power. The three genes are potential biomarkers in IPF.

PADI4 is a member of the peptidyl arginine deiminase (PAD) family. Depending on PAD enzyme activity, PADI4 can modify the guanidine group of arginine into the ureido group of citrulline. Although PADI4 plays an important role in human innate immunity, NETosis is a special pathway of neutrophil death. When an extracellular pathogen invades, neutrophils release a network of linear fibers called neutrophil elastase traps (NETs) into the extracellular environment to trap pathogens. The PADI4 enzyme predominates in neutrophils and initiates the process of NETs by unfolding and decondensing nuclear chromatin via citrullination of histones. Subsequently, the PADI4 co-expanded NETs enter the extracellular high-calcium space to complete the bactericidal process by citrullinating extracellular proteins such as fibrin (25,26).

PADI4 was first reported by the genome-wide association study as a disease susceptibility gene for non-MHC rheumatoid arthritis (RA). PADI4 is currently considered to be an important in the pathogenesis of RA (27). Studies have shown that PADI4 interferes with fibrosis in several
Figure 12 Correlations of hub genes with immune cells and immune function.

Figure 13 Receiver operating characteristic curve of immHub genes in the training dataset. immHub genes, hub genes correlating with immunity. AUC, area under the curve.

Figure 14 Nomogram of immHub genes in the training dataset. immHub genes, hub genes correlating with immunity; IPF, idiopathic pulmonary fibrosis.

Figure 15 Receiver operating characteristic curve of immHub genes in the validation dataset. immHub genes, hub genes correlating with immunity. AUC, area under the curve.

Figure 16 Nomogram of immHub genes in the validation dataset. immHub genes, hub genes correlating with immunity; IPF, idiopathic pulmonary fibrosis.
Figure 17 Heat map of the enrichment of immune cells and immune function in IPF and normal lung tissues. IPF, idiopathic pulmonary fibrosis.

Figure 18 Comparison of enrichment scores for 16 immune cells and 13 immune function in IPF and normal lung tissues. P values are shown as: ns: not significant, *P<0.05, **P<0.01, ***P<0.001. IPF, idiopathic pulmonary fibrosis.

organs, including the lung, by affecting the release of NETs. Suzuki et al. found that the expression of NETs in PADI4 knockout mice was suppressed, and the level of tumor necrosis factor-α showed a downward trend. PADI4 deficiency prevented a reduction in the number of alveolar epithelial cells and pulmonary vascular endothelial
cells and an increase in ACTA2-positive mesenchymal cells and S100A4-positive fibroblasts in the lungs. Thus, PADI4 deficiency can ameliorate bleomycin-induced NET formation and pulmonary fibrosis (28). In this study, PADI4 gene expression was negatively regulated and suppressed in IPF patients. Recent studies have shown that nintedanib suppressed PADI4 expression in mice. Nintedanib was administered to mice models of RA-related interstitial lung disease (RA-ILD). Changes in PADI4 expression after nintedanib administration were suppressed in the RA-ILD lung. However, similar results have been shown after treatment with nintedanib for galectin-3, which is expected to be the next anti-fibrotic target (29,30).

Insulin-like growth factor-binding protein 7 (IGFBP7) is a glycoprotein with a similar structure to insulin. It localizes to the cell surface and interacts with proteoglycans to promote cell adhesion to fibronectin and cohesin (31,32). A study has shown that IGFBP7 can intervene in the immune system through the IGF pathway, and deletion of IGFBP7 leads to downregulation of a set of interferon (IFN)-γ-mediated immune responses. This suggests that IGFBP7 may be a potential target for the treatment of chronic inflammatory diseases.

### Table 1

| Rank | Name                        | P value     | Genes               |
|------|-----------------------------|-------------|---------------------|
| 1    | Dimethyl sulfoxide CTD 00005842 | 3.18E-04    | GADD45A; PADI4      |
| 2    | Dronabinol CTD 00006853      | 0.001192927 | GADD45A; IGFBP7     |
| 3    | Primaquine PC3 UP            | 0.001389158 | IGFBP7; PADI4       |
| 4    | Scopolamine PC3 UP           | 0.001519268 | IGFBP7; PADI4       |
| 5    | Pregnenolone PC3 UP          | 0.001532594 | IGFBP7; PADI4       |
| 6    | Etoposide CTD 00005948       | 0.00156616  | GADD45A; IGFBP7     |
| 7    | PTAQUILOSIDE CTD 00001981    | 0.001649143 | GADD45A             |
| 8    | Lead dinitrate CTD 00000898  | 0.001649143 | GADD45A             |
| 9    | Streptotigin CTD 00006785    | 0.001649143 | PADI4               |
| 10   | PRODIONE CTD 00001594        | 0.001649143 | GADD45A             |

Figure 19 Correlations among 16 immune cells and 13 immune function. DCs, dendritic cells; NK, natural killer; TIL, tumor infiltrating lymphocyte; APC, antigen-presenting cell; CCR, cytogenetic complete remission; HLA, human leucocyte antigen; MHC, major histocompatibility complex; IFN, interferon.
regulated genes that regulate antigen presentation, which manifests as inhibition of antigen presentation and the decreased infiltration of CD8+ and CD4+ T cells and natural killer cells in tumor tissue (33).

IGFBP7 can promote various BPs related to fibrotic diseases in different organs. For example, it is involved in the occurrence and development of kidney and liver fibrosis (34-35). TGF-β enhances the expression of IGFBP7 by stimulating renal tubular epithelial cells, and its silencing reduces the number of TGF-β-induced α-SMA-positive cells (35). Likewise, IGFBP7 has been shown to promote hepatic stellate cell activation, favor the development of a myofibroblast phenotype, and promote the production of extracellular matrix proteins (34). Extracellular vesicles (EVs) are important players in intercellular communication. Velázquez-Enríquez et al. revealed key proteins in the EV cargo associated with IPF in vitro through proteomic analysis and found that the IGFBP7 level in EVs was higher in IPF tissue than in normal tissue (36), which was consistent with the findings of Hsu et al. (37).

Growth arrest and DNA-damage-inducible alpha (GADD45A) is a member of the GADD family and highly conserved. The GADD45A protein is widely expressed in normal tissues (38). R-loops are DNA-RNA hybrids enriched at CpG islands (CGIs) that can regulate chromatin states. GADD45A binds directly to the R-loops and mediates local DNA demethylation by recruiting TET1. GADD45A is considered to be an epigenetic R-loop reader (39). As an immune cell, T cells play an important role in the pathogenesis of IPF. There is increasing evidence that Th1/Th2 imbalance is associated with the inflammatory phase in pulmonary fibrosis. IFN-γ and interleukin (IL)-12 secreted by Th1 cells have anti-fibrotic effects, while IL-4, IL-5, and IL-13 secreted by Th2 cells are involved in pulmonary fibrosis. It has been proposed that Th1 response is protective whereas Th2 response causes damage (40). In addition, Tregs are also involved in the pathogenesis of IPF. Boveda-Ruiz et al. found that Tregs can increase TGF-β1 release and collagen deposition in the early stage of pulmonary fibrosis, proving that Tregs have a pro-fibrotic effect in this stage (41); however, Tregs in the late stage have anti-fibrotic effects. Xiong et al. showed that Treg depletion attenuated pulmonary fibrosis by inducing a Th17 response and shifting the Th1/Th2 cytokine balance into Th1 dominance (42). Therefore, immune cells play an auxiliary role in the pathogenesis of IPF, and some immune cells can regulate IPF in a bidirectional manner (43).

GADD45A plays essential roles in controlling differentiation and activating T cells in the immune system; in particular, GADD45A can inhibit p38 kinase activation downstream of T-cell receptors. Salvador et al. found that GADD45A was a negative regulator of T cell proliferation induced by T-cell antigen-receptor-mediated activation and that lack of GADD45A led to the development of a lupus-like syndrome; in addition, the counts of leukocytes and lymphocytes in the peripheral blood of GADD45A-deficient mice decreased (44,45).

Although the pathogenic mechanisms of IPF are still unclear, more studies have shown that immune activation is linked to its pathogenesis. In our current study, comparisons of the enrichment scores of 16 immune cells in IPF and normal tissues and the activities of 13 immune function from the ssGSEA showed that, among IMMCs, B_cells, CD8+ T_cells, DCs, iDCs, Macrophages, Mast_cells, T_helper_cells Tfh, and Th2_cells were highly expressed in IPF whereas NK_cells and Neutrophils were lowly expressed in IPF. The expression of PADI4 was downregulated in IPF tissues and showed a positive correlation with the level of neutrophils. Because PADI4 can prevent tissue fibrosis caused by excessive release of NETS, it is believed that PADI4 plays a negative regulatory role in the occurrence of IPF. IGFBP7 was highly expressed in IPF and negatively correlated with APC_co_inhibition and positively correlated with Mast_cells. It has been demonstrated that IGFBP7 is involved in BPs (e.g., cell adhesion), the pathogenesis of IPF, and regulation of the antigen presentation system, which is consistent with the results of our current study.

To sum up, our study showed that IPF is closely related to CRs and immune dysregulation. Analysis based on models constructed from three immune-related CR genes showed that high expression of PADI4 negatively correlated with the occurrence of IPF, and high expressions of IGFBP7 and GADD45A positively associated with the pathogenesis of IPF. The AUC was 0.771 in the training dataset and 0.759 in the validation dataset, suggesting that...
PADI4, IGFBP7, and GADD45A are potential biomarkers of IPF for predicting risk. This study will help to clarify the pathogenic mechanism of IPF and provide a reference for future targeted therapies.

Our study still had some limitations, and the results of this study have yet to be validated by relevant in vivo and in vitro experiments. Our future studies will continue to focus on the roles of genes such as PADI4, IGFBP7, and GADD45A in IPF.

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Footnote
Reporting Checklist: The authors have completed the TRIPOD reporting checklist. Available at https://atm.amegroups.com/article/view/10.21037/atm-22-3700/rc

Conflicts of Interest: All authors have completed the ICMJE uniform disclosure form (available at https://atm.amegroups.com/article/view/10.21037/atm-22-3700/coif). The authors have no conflicts of interest to declare.

Ethical Statement: The authors are accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved. The study was conducted in accordance with the Declaration of Helsinki (as revised in 2013).

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