A functional genomic model for predicting prognosis in idiopathic pulmonary fibrosis

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Abstract

Background: The course of disease for patients with idiopathic pulmonary fibrosis (IPF) is highly heterogeneous. Prognostic models rely on demographic and clinical characteristics and are not reproducible. Integrating data from genomic analyses may identify novel prognostic models and provide mechanistic insights into IPF.

Methods: Total RNA of peripheral blood mononuclear cells was subjected to microarray profiling in a training (45 IPF individuals) and two independent validation cohorts (21 IPF/10 controls, and 75 IPF individuals, respectively). To identify a gene set predictive of IPF prognosis, we incorporated genomic, clinical, and outcome data from the training cohort. Predictor genes were selected if all the following criteria were met: 1) Present in a gene co-expression module from Weighted Gene Co-expression Network Analysis (WGCNA) that correlated with pulmonary function (p < 0.05); 2) Differentially expressed between observed “good” vs. “poor” prognosis with fold change (FC) >1.5 and false discovery rate (FDR) < 2 %; and 3) Predictive of mortality (p < 0.05) in univariate Cox regression analysis. “Survival risk group prediction” was adopted to construct a functional genomic model that used the IPF prognostic predictor gene set to derive a prognostic index (PI) for each patient into either high or low risk for survival outcomes. Prediction accuracy was assessed with a repeated 10-fold cross-validation algorithm and independently assessed in two validation cohorts through multivariate Cox regression survival analysis.

Results: A set of 118 IPF prognostic predictor genes was used to derive the functional genomic model and PI. In the training cohort, high-risk IPF patients predicted by PI had significantly shorter survival compared to those labeled as low-risk patients (log rank p < 0.001). The prediction accuracy was further validated in two independent cohorts (log rank p < 0.001 and 0.002). Functional pathway analysis revealed that the canonical pathways enriched with the IPF prognostic predictor gene set were involved in T-cell biology, including iCOS, T-cell receptor, and CD28 signaling.

Conclusions: Using supervised and unsupervised analyses, we identified a set of IPF prognostic predictor genes and derived a functional genomic model that predicted high and low-risk IPF patients with high accuracy. This genomic model may complement current prognostic tools to deliver more personalized care for IPF patients.

Keyword: Idiopathic pulmonary fibrosis (IPF), Peripheral blood mononuclear cells (PBMCs), Gene expression profiling, Functional genomic model, Prognosis prediction

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Background

Idiopathic pulmonary fibrosis (IPF) is a fibrotic interstitial lung disease characterized by irreversible scarring of the lung parenchyma that predominantly affects older adults. While older retrospective studies suggested median survival was 2–3 years [1–3], IPF has a highly heterogeneous disease course, making prognostication difficult [4, 5]. While lung transplantation remains the sole intervention to prolong survival in patients with IPF [6], organ scarcity, and ineligibility secondary to comorbid health conditions, make this available to only a few. Pirfenidone [7] and nintedanib [8] have emerged as promising therapies that slow disease progression. Several other medications are currently under investigation. Without the ability to predict disease course, it is difficult to identify which IPF patients are most likely to benefit from these new therapies or from lung transplantation.

Many clinical parameters, including race, gender, age, radiographic and/or histopathologic patterns, and pulmonary function tests have been linked to prognosis in patients with IPF [9, 10]. Lung tissue-based molecular genomic signatures [11, 12] have also been used to predict IPF progression; however, given the resources needed to perform lung biopsy and risk associated with the procedure, the applicability of such genomic signatures is limited. Peripheral blood mononuclear cells (PBMC), comprised of circulating monocytes, T-cells, B-cells, and natural killer cells, have been successfully used as an alternative for exploratory transcriptional profiling studies [13–15]. Advantages of using PBMC over lung biopsy specimens to delineate molecular mechanisms of IPF include easier access, larger quantities, and the ability to dynamically assess disease status through longitudinal sample collection.

Using PBMC gene expression profiling, our group previously identified a genomic signature consisting of 52 genes that predicted survival in patients with IPF [16]. While this investigation drew attention to the potential role of T cell signaling in IPF progression, the contribution of other genes identified in the study were not addressed. Furthermore, the gene set identified from our previous study did not provide a weighted score for the gene expression pattern, which has the potential to be useful in practical application. We therefore aim to construct a functional genomic model to better predict prognosis of IPF patients. To do so, we compiled a set of IPF prognostic predictor genes from previously reported microarray data in the training cohort (accession number GSE28221) [17, 18]. First, we coupled PBMC gene expression profiling to IPF clinical traits using an unbiased “Weighted Gene Co-expression Network Analysis (WGCNA)” approach which is useful for describing the pairwise correlated expression among gene transcripts with co-regulation implications [19–21] and to restrict the search space of genes to those genes in modules associated with pulmonary function. Second, we performed a supervised “Significance Analysis of Microarray (SAM)” approach to identify differentially expressed genes between observed “good” vs. “poor” prognosis IPF patients. Third, we identified genes based on their association with survival. The IPF prognostic predictor gene set satisfying all aforementioned three functional genomic criteria was used to construct a genomic prediction model and derived a prognostic index (PI) score for each patient in the training cohort. We then assessed the prognostic prediction specificity in the training cohort and further validated it in two independent cohorts. This work produced a functional genomic model with a mechanism-anchored IPF prognostication score for each patient, which may better identify those most likely to benefit from IPF-anchored IPF management and provide a tool for personalized IPF management.

Methods

Study populations

Study populations were collected, as previously described, from the University of Chicago Medical Center (UCMC) and University of Pittsburg Medical Center (UPMC) [16]. The training cohort consisted of 45 individuals with IPF collected from November 2007 to July 2009 at UCMC. The University of Chicago validation cohort (UCV) consisted of 21 individuals with IPF along with 10 healthy control subjects without lung disease collected from February 2007 to October 2007. The University of Pittsburg validation cohort (UPV) consisted of 75 individuals with IPF collected from March 2001 to September 2010. While the site source of the samples overlapped between cohorts, all samples were independent of each other. All patients with IPF met American Thoracic Society/European Respiratory Society (ATS/ERS) diagnosis criteria [2]. The local Institutional Review Boards at the University of Chicago and University of Pittsburg Medical Center approved the study and informed consent was provided by all study subjects.

Demographic information, clinical characteristics, and pulmonary function tests were collected from all patients with IPF. Spirometry testing, including forced vital capacity percent predicted (FVC% predicted), diffusion capacity for carbon monoxide percent predicted (DlCO % predicted) as well as lung volumes by plethysmography were obtained per ATS guidelines [19–21]. The composite physiologic index (CPI) was calculated as described by Wells et al. [22]. Survivorship was obtained from medical records, telephone interviews, and the social security death index database. The prognosis of IPF subjects was dichotomously categorized as good or poor based on observed survival over 3 years of follow-up.

PBMC sample collection, RNA isolation, microarray hybridization, and data processing

See details in Additional file 1. Microarray experiments were compliant with MIAME (Minimum Information
Identification of gene co-expression modules correlated with clinical traits in training cohort
Normalized microarray data were filtered to remove redundant genes and genes with minimum variation (i.e. coefficient of variation <0.3 across all samples). Genes that passed filtering criteria were clustered into gene modules, based on their co-expression pattern, using an unsupervised “Weighted gene co-expression network analysis (WGCNA)” package in R 2.13 [23]. Principal Component Analysis (PCA) was used to calculate an eigengene for each gene module. Pearson’s correlation was used to determine the significance of correlation (p < 0.05) between the eigengenes of individual gene modules with each clinical parameter including race, sex, age, FVC % predicted, D_{1}CO % predicted, and CPI.

Identification of differentially expressed genes in the training cohort
Significant Analysis of Microarray (SAM) software [24] was used to identify differentially expressed genes between observed good vs. poor IPF prognosis using criteria of fold change (FC) >1.5 and false discovery rate (FDR) < 2 %.

Survival analysis
Survival analysis was performed using unadjusted log rank testing along with univariate and/or multivariate Cox regression analysis. After checking to ensure that the proportional hazard assumption was met with each Cox model, subdistributional mortality hazards were determined for covariates by treating lung transplantation as a competing event, as previously described by Fine and Gray [25]. Survival time was defined as time from blood draw to death, transplant, loss-to-follow-up or study conclusion. Patients who were lost to follow-up were censored at that time in survival modeling. Survival between groups was plotted using the Kaplan-Meier estimator.

Compilation of the IPF prognostic predictor gene set from the training cohort for a genomic model construction
To construct a functional genomic model predictive of IPF prognosis, genomic, clinical, and outcome data from the training cohort were analyzed to identify a set of genes with individual prognostic significance. Genes were selected for the “IPF prognostic predictor gene set” if they met all of the following criteria: 1) genes in specific gene co-expression modules that correlated with pulmonary function (p < 0.05) in WGCNA, 2) genes differentially expressed (FC> 1.5 and FDR < 2 %) between observed good vs. poor prognosis by SAM, and 3) genes predictive of mortality (p < 0.05) in univariate Cox regression analysis.

Development and validation of the functional genomic model to predict prognosis
The set of IPF prognostic predictor genes identified was used to construct a genomic model using “Survival risk group prediction” implemented in BRB-ArrayTools 4.2 [26] to predict prognosis in IPF patients. The output of the genomic model is a patient-specific “prognostic index (PI)” score. PI of each patient in training cohort was derived from formula, \[ \sum W_i \times X_i + 13.5, \] where \( W_i \) and \( X_i \) represent the weight (computed by supervised PCA) and log-intensity of the \( i \)-th gene in the gene set. To assign a patient to either a high- or low-risk group, each patient’s PI was compared to a predetermined classification threshold. For this study, the threshold was set at the upper tertile in the training cohort according to clinical observation [3]. A “10-fold Cross-Validation (CV)” algorithm was used to assess the classification specificity. Briefly, 10 % of patients were randomly omitted leaving the remaining 90 % of patients to construct the genomic model and derive a PI for each of the omitted samples. The PI of omitted individuals was then ranked relative to the PI of patients included in the CV model. Finally, we determined the predicted risk category based on the percentile ranking, the number of risk groups specified (i.e. \( n = 2 \) in current study), and the empirical risk percentile setting (i.e. low/ high risk = 66.7/33.3). Misclassification rate was determined by the discrepancy between the predicted low or high-risk category with the observed good or poor prognosis according to follow-up. Receiver-Operating-Characteristic (ROC) analysis with area under curve (AUC) calculation was performed to assess how well the PI distinguished IPF patients with low vs. high-risk prognosis. To perform an independent validation of the predictor, we applied the PI weights computed from the training set of 45 IPF samples to the calculation of the PI on the UCV and UPV cohorts. Details can be found in Additional file 1.

Functional pathways enrichment analysis
Significant biological processes in Gene Ontology associated with the set of IPF prognostic predictor genes were identified using R package “GOSim” [27] with the criterion of \( q \)-value (Benjami-Yekutieli adjusted \( p \)-value) <0.01. Significant canonical pathways or gene interaction networks were analyzed using Ingenuity Pathway Analysis (IPA) software (Ingenuity Systems, Redwood City, CA) with the criterion of the right-tailed (referring to the overrepresented pathway) Fisher’s exact test \( q \)-value (Benjamini-Hochberg adjusted \( p \)-value) <0.05.
IPF diagnosis prediction using prognosis index derived from the functional genomic model

Using the generated PI, ROC analysis with AUC calculation was performed in UCV cohort to assess how well the PI distinguishes IPF patients from healthy controls. The true positive rate (sensitivity) is plotted in function of the false positive rate (1-specificity) for different cut-off points. Each point on the ROC curve represents a sensitivity/false alarm pair corresponding to a particular decision threshold.

Statistical analysis

Continuous variables are reported as a mean (± standard deviation) and compared using a one-way analysis of variance. Categorical variables are reported as counts and percentages and compared using a chi-square or Fischer’s exact test, as appropriate. Pearson’s correlation was used to evaluate the correlation of prognostic index (PI) derived from genomic model with clinical parameters. ROC analysis with AUC calculation was performed using R package “caTools”. Other than when indicated above, statistical analysis was conducted using STATA 12 (StataCorp. 2011. College Station, TX).

Results

Demographic and clinical characteristics of patients with IPF

Demographic and clinical characteristics for each IPF cohort are shown in Table 1. Significant differences between the training, UCV and UPV cohorts were observed with respect to male gender (90 % vs. 71.4 % vs. 69.3 %, respectively; \( p = 0.05 \)), white race (82.2 % vs. 81.8 % vs. 97.3 %, respectively; \( p = 0.004 \)), follow-up months (18.8 vs. 43.8 vs. 23.5 months, respectively; \( p < 0.001 \)), months to death (12.7 vs. 26.8 vs. 14.2, respectively; \( p = 0.02 \)) and lung transplantation (2.2 % vs. 9.5 % vs. 20 %, respectively; \( p = 0.009 \)). No differences between cohorts were observed with respect to age, FVC % predicted, DLCO % predicted or CPI.

Identification of gene co-expression modules correlated with clinical traits in training cohort

“WGCNA” package in R was utilized to cluster 2718 genes that passed the filtering criteria into eight gene co-expression modules denoted by different colors (Fig. 1). Optimization of the power for adjacency transition and the parameters for gene clustering dendrogram are depicted in Additional file 2: Figure S1A and S1B, respectively. The eigengene values of individual modules were then computed by PCA and correlated with clinical traits to envision the association between co-expressed gene pattern features with clinical features [23]. The significance of correlation with clinical traits was determined by Pearson’s correlation assay with \( p < 0.05 \). As shown in Fig. 1, three gene modules demonstrated significant correlation (red box) or anti-correlation (green box) with clinical traits: turquoise module with male gender (\( p = 0.009 \)), FVC % predicted (\( p = 0.0002 \)), DLCO % predicted (\( p = 0.005 \)), and CPI (\( p = 0.002 \)); red module with FVC % predicted (\( p = 0.03 \)); black module with FVC % predicted (\( p = 0.002 \)) and CPI (\( p = 0.03 \)). Of note that 1199, 157, and 131 genes consist of 55 % (1487/2718) of total genes were clustered into turquoise, read, and black modules, respectively (Fig. 1).

Compilation of the set of IPF prognostic predictor genes

In an effort to reduce the number of genes in turquoise, red, and black modules associated with pulmonary function to a clinically applicable number and to incorporate predictive prognostic feature to co-expressed gene pattern feature, we applied two additional distinct approaches to the analytic pipeline (Fig. 2a). Using SAM, we identified 155 that were differentially expressed (DE) genes between IPF patients with good and poor prognosis. Using univariate Cox regression analysis, we identified 836 genes which were significantly correlated with survival (\( p < 0.05 \)) (Fig. 2b). Notably, 147 of the 155 DE genes were overlapped with the 1487 genes combined from the turquoise, black, and red gene modules. This integrative functional genomic approach yielded a set of 118 prognostic genes.

Table 1 Demographic and Clinical Characterizations among Study Cohorts

| Characteristic                | Training cohort (n=45) | UCV cohort (n=21) | UPV cohort (n=75) | \( p\)-value |
|------------------------------|-----------------------|-------------------|-------------------|--------------|
| Age, mean (±SD)              | 67.1 (8.2)            | 68.9 (8.2)        | 68.5 (7.8)        | 0.48         |
| Male gender, \( n\) (%)      | 40 (90)               | 15 (71.4)         | 52 (69.3)         | 0.05         |
| White race, \( n\) (%)       | 37 (82.2)             | 18 (81.8)         | 73 (97.3)         | 0.004        |
| Follow-up months, mean (±SD) | 18.8 (11.9)           | 43.8 (29.4)       | 23.5 (12.7)       | <0.001       |
| Months to death, mean (±SD)  | 12.7 (10.9)           | 26.8 (20.1)       | 14.2 (10.6)       | 0.02         |
| FVC % predicted, mean (±SD)  | 60.6 (14.3)           | 64.7 (12.7)       | 65.4 (16.7)       | 0.25         |
| DLCO % predicted, mean (±SD) | 43.4 (17.7)           | 43.2 (15.6)       | 48.9 (18.6)       | 0.19         |
| CPI, mean (±SD)              | 55.6 (13)             | 54.7 (10.7)       | 50.7 (13.7)       | 0.11         |
| Lung transplantation, \( n\) (%) | 1 (2.2)               | 2 (9.5)           | 15 (20)           | 0.009        |
predictor genes (Fig. 2b). A list of 118 prognostic predictor genes attributed to turquoise \((n=110)\), red \((n=5)\), and black \((n=3)\) modules was shown in Table 2.

Pathway and network characterization of the IPF prognostic predictor genes

To assess the pathways and networks of the 118 IPF prognostic predictor genes involved, we carried out a functional enrichment analysis using Ingenuity pathway analysis (IPA) and gene network analysis software. Surprisingly, all significant canonical pathways with \(-\log(q\text{-value}) > 1.3\) were involved in T-cell biology (Fig. 2c). Several genes were involved in multiple T-cell signaling pathways, including CD28 receptor (CD28), inducible T-cell co-stimulator (ICOS), lymphocyte-specific protein tyrosine kinase (LCK), interleukin 7 receptor (IL7R), and major histocompatibility complex, class II, DQ alpha 1 (HLA-DQA1) (Additional file 1: Table S1). Of note that these pathways comprised genes mostly if not all in the turquoise module which represent \(>93\%\) of genes in the IPF prognostic predictor gene set. Ingenuity network modeling based on “Ingenuity Knowledge” database prioritized five significant gene networks with score \(\geq 32\). Seventy-eight of the 118 IPF prognostic predictor genes (66\%) was functionally connected in these five gene networks, supporting a concordance between expression correlations and functional connections of individual genes. The first of the five gene interaction networks is displayed in Additional file 3: Figure S2 showing that five hub genes have significantly higher degree of linkage to other nodes in the network including ras homolog gene family member H (RHOH), G protein-coupled receptor 18 (GPR18), G protein-coupled receptor 171 (GPR171), and G protein-coupled receptor 174 (GPR174), and lymphocyte-specific protein tyrosine kinase (LCK).

Construction and cross-validation of a functional genomic model for prognosis prediction

To determine prognosis prediction power of the gene set, we constructed a novel functional genomic model using “Survival risk group prediction” implemented in BRB-
**A**

**Array Data Processing**
- Affymetrix Exon array normalization
- Summarize exon probes into genes and map to U133 plus 2 array
- Filter probe sets by redundancy, intensity and coefficient of variation

**Prognostic Predictor Gene Set Compilation**
- Gene co-expression modules by WGCNA
- Differentially expressed genes by SAM
- Survival correlated genes by Cox regression

**Functional Genomic Model For Prognosis Prediction**
- Model construction with prognostic predictor genes
- Calculate "Prognostic Index (PI)" score for each patient in training cohort
- Assess prognostic prediction specificity by 10-fold cross validation
- Validate model in two independent cohorts

**B**

- WGCNA (n=1487)
- SAM (n=155)
- Cox regression (n=836)
- 118 IPF prognostic predictor genes

**C**

- PKCθ Signaling in T Lymphocytes
- Nur77 Signaling in T Lymphocytes
- Primary Immunodeficiency Signaling
- CTLA4 Signaling in Cytotoxic T Lymphocytes
- G-Protein Coupled Receptor Signaling
- T Helper Cell Differentiation
- CD28 Signaling in T Helper Cells
- Role of NFAT in Regulation of Immune Response
- T Cell Receptor Signaling
- iCOS-iCOSL Signaling in T Helper Cells

*Fig. 2* (See legend on next page.)
Compilation and functional characterization of IPF prognostic predictor gene set. a A flowchart illustrates the procedures and approaches used for IPF prognostic predictor gene set compilation. Left panel: Array data processing. Affymetrix Exon 1.0 ST Array data was normalized, probe sets mapped to U133 plus 2 Array, and filtered based on redundancy, intensity, and coefficient of variation across all samples. Middle panel: IPF prognostic predictor gene set compilation. Three approaches used to compile IPF prognostic predictor gene set: Co-expressed gene modules correlated with pulmonary function identified by WGCNA; Differentially expressed genes between “good” and “poor” prognosis patients identified by SAM (fold change > 1.5 & FDR < 2.5 %); Survival-correlated genes identified by Cox regression (p < 0.005). Right panel: Genomic model IPF prognosis prediction. IPF prognostic predictor gene set was used to construct a genomic model; Prognostic Index (PI) score was calculated from each patient in training cohort; Prediction specificity was assessed by 10-fold cross validation; Genomic model was validated in two independent cohorts using weights of PI calculated from training cohort. b Venn diagram illustrates the selection criteria for IPF prognostic predictor genes. A total of 118 genes were compiled for downstream data analyses. c Canonical pathways enriched from IPF prognostic predictor genes by Ingenuity Pathway Analysis software. Significant pathways were set with criterion of q-value < 0.05 (i.e. -log (q-value) > 1.3) using one-tailed Fisher’s exact test.

Table 2 List of 118 IPF prognostic predictor genes within the red, black and turquoise gene modules

| Gene     | FC | Gene     | FC | Gene     | FC | Gene     | FC |
|----------|----|----------|----|----------|----|----------|----|
| IL1R2    | 2.0| PPWD1    | −1.7| AS1F1    | −1.6| ABCD2    | −1.5|
| ERAF     | 2.0| CETN3    | −1.6| LM07     | −1.6| GZMK     | −1.5|
| CEACAM8  | 1.8| SH2D1A   | −1.6| GCET2    | −1.6| TRIM52   | −1.5|
| ARG1     | 1.6| SLC39A10 | −1.6| PAQR8    | −1.6| C8orf15  | −1.5|
| FOXO3    | 1.5| SPPR4    | −1.6| BIRC3    | −1.6| ITK      | −1.5|
| TNS1     | 1.5| WDR75    | −1.6| CAMK4    | −1.6| ICOS     | −1.5|
| CYP4F2   | 1.5| C14orf64 | −1.6| ZC3H6    | −1.6| FHT      | −1.5|
| CYP4F3   | 1.5| KPNAS    | −1.6| CD28     | −1.6| TSEPA    | −1.5|
| ARHGAP5  | −1.8| NOP58    | −1.6| GTPBP0   | −1.6| NPCD1    | −1.5|
| ORC3L    | −1.8| PARP5    | −1.6| C5orf51  | −1.6| OXNAD1   | −1.5|
| ZNF100   | −1.8| PRO0471  | −1.6| TRBC1    | −1.6| IL7R     | −1.5|
| UtP5     | −1.8| RCAN3    | −1.6| CAMK2D   | −1.5| HLA-DQA1 | −1.5|
| ANKR36B  | −1.8| C7orf64  | −1.6| PPMP1K   | −1.5| TMEM156  | −1.5|
| LOC399753| −1.8| ANKR36   | −1.6| CCDC76   | −1.5| HLA-DQA1 | −1.5|
| KCNA3    | −1.8| GPR174   | −1.6| CASD1    | −1.5| LOC401397| −1.5|
| RHOH     | −1.8| NDUFA4   | −1.6| pRy10    | −1.5| CDK6     | −1.5|
| LCK      | −1.8| CCDC141  | −1.6| DPP4     | −1.5| GCNT4    | −1.5|
| C16orf52 | −1.7| GPR18    | −1.6| S1PR1    | −1.5| NELL2    | −1.5|
| TC2N     | −1.7| DDX60    | −1.6| ITGA6    | −1.5| FLJ33630 | −1.5|
| HIVEp    | −1.7| TMEM209  | −1.6| GBP4     | −1.5| TRAT1    | −1.5|
| KIF3A    | −1.7| GV1N1    | −1.6| ABCE1    | −1.5| LEF1     | −1.5|
| IFT80    | −1.7| TMEM161B | −1.6| TXK      | −1.5| FCRL3    | −1.5|
| TIA1     | −1.7| USP53    | −1.6| TRAF5    | −1.5| GUSBL2   | −1.5|
| ZNF83    | −1.7| TRA17    | −1.6| SLAMF6   | −1.5| SEPS3    | −1.5|
| SETD82   | −1.7| MRPL1    | −1.6| CD96     | −1.5| BTLA     | −1.5|
| WDR36    | −1.7| SNORD116 | −1.6| PRKACB   | −1.5|         |     |
| ZNF141   | −1.7| GPR171   | −1.6| ALG10B   | −1.5|         |     |
| TRBC1    | −1.7| MGC40069 | −1.6| NBP10    | −1.5|         |     |
| FAM69A   | −1.7| LOC439949| −1.6| MGAT4A   | −1.5|         |     |
| C1GALT1  | −1.7| CCR7     | −1.6| INPP4B   | −1.5|         |     |
| GIMAP5   | −1.7| NUP43    | −1.6| STAT4    | −1.5|         |     |

FC = Fold change; * denotes genes in red module; § denotes genes in black module; the rest of genes are in turquoise module.
ArrayTools. The output of the genomic model is a patient-specific prognostic index (PI). The genomic model was displayed in Fig. 3a, where $W_i$ and $X_i$ represented the weight and log-intensity of the $i$-th gene in the gene set identified from training cohort. Based on previous clinical observation, we empirically set the percentile population of low vs. high risk to the lower tertiles vs. upper tertile, respectively. Patients in the training cohort were categorized as low or high risk based on whether the patient-specific PI fell in the lower tertiles vs. upper tertile, respectively. We obtained PI values ranging from $-2.14$ to $3.45$, which were continuously associated with risk of death.

Of the 45 individuals in the training cohort, 30 (66.7 %) and 15 (33.3 %) were assigned to the low risk or high risk groups, respectively based on PI. Misclassification rate was determined by comparison of the predicted class to clinically observed outcomes in the training cohort, in which 28 patients were with "good" and 17 with "poor" prognosis. Misclassification rate (20 %) was determined by 10-fold CV and computed as $(k + n)/total \ cases$, where $k$ represents the predicted high risk that are observed as good prognosis, and $n$ represents the predicted low risk that are observed as poor prognosis. PI independently predicted survival in univariate competing-risk Cox regression (Sub-hazard ratio (SHR) 2.7; 95 % CI 1.9-3.9; $p < 0.001$) and in multivariate competing-risk Cox regression after adjustment for baseline CPI (SHR 2.3; 95 % CI 1.5-3.4; $p < 0.001$).

IPF genomic model predicts prognosis in two independent validation cohorts

The IPF genomic model shown in Fig. 3a was applied to two independent validation cohorts, UCV and UPV, where two microarray platforms (Affymetrix and Agilent, respectively) were used. The weight of each gene and the constant (13.5) derived from training cohort were carried over for independent validation of the genomic model. After annotating the gene expression data from each cohort with UniGene annotations, 10 of the IPF prognostic predictor genes did not map to Agilent Human 4x44k Whole Genome Expression array in UPV. Therefore, we computed the PI using 108 classifiers for UPV and 118 classifiers for UCV, respectively. Individuals in the UCV and UPV cohorts were classified as low vs. high risk based on whether the patient-specific PI fell in the lower tertiles vs. upper tertile, respectively. In both validation cohorts, patients classified as low risk demonstrated significantly improved survival over those classified as high risk ($p < 0.001$ for UCV and $p = 0.002$ for UPV) (Fig. 4a&b). PI remained a significant predictor of survival in univariate competing-event Cox regression in the UCV (SHR 2.0; 95 % CI 1.2-3.4; $p = 0.005$) and UPV (SHR 1.8; 95 % CI 1.1-2.7; $p = 0.01$).
cohorts. This association remained in the UCV (SHR 1.7; 95% CI 1.04-2.93; p = 0.035) and UPV (SHR 1.9; 95% CI 1.2-3.0; p = 0.005) cohorts after adjusting for baseline CPI in multivariate Cox regression.

To evaluate the biological reproducibility of IPF prognostic predictor genes within the set, fold change of each gene was calculated between predicted low-risk and high-risk prognosis patients in each cohort, and then regressed between training versus each validation cohort. The regression plot revealed a strong concordance of fold changes between training and each validation cohort (Additional file 4: Figure S3). Pearson’s correlation analysis showed significant correlation of the classifier fold changes between training and UCV (p = 2.2 × 10⁻¹⁶) as well as between training and UPV cohort (p = 1.4 × 10⁻⁶).

Multivariate correlation of Prognostic Index with clinical parameters

We investigated the impact of clinical parameters on the genomic model (Additional file 1: Table S2). Pearson’s correlation analysis showed that PI was significantly correlated with CPI in the training (coefficient = 0.36, p = 0.016) and UCV (coefficient = 0.53, p = 0.01) cohorts, but not in UPV cohort (coefficient = −0.05, p = 0.69). While the PI of an individual sample was not associated with age, it was significantly higher in males compared to females in the training and UPV cohorts (p = 0.02 and 0.04, respectively). The impact of ethnic diversity on PI was evaluated and no difference was found between Caucasian and non-Caucasian in the training (p = 0.92) and UCV cohort (p = 0.79) despite each cohort containing 18% non-Caucasians (Additional file 1: Table S2). This result was consistent with the data obtained from WGCNA analysis showing the turquoise gene module was significantly correlated with female gender (Fig. 1, coefficient = −0.39, p = 0.009).

Application of genomic model in IPF diagnosis

To evaluate whether our genomic model is able to discriminate IPF patients from healthy individuals, we conducted the ROC analysis of the PI by plotting the true positive (sensitivity) versus false alarm (1-specificity). PI accurately distinguished IPF subjects from healthy individuals in the UCV cohort with an AUC 0.96. The sensitivity of IPF diagnosis by PI at 10% false alarm was 80% (red line in Additional file 5: Figure S4).

Discussions

In this study, we constructed a functional genomic model that predicted survival in three independent cohorts of IPF patients. In the training cohort, we analyzed genomic data using both unsupervised WGCNA and supervised SAM approaches. By applying WGCNA algorithm, we first associated the pathophysiological alterations in the transcriptome level to the clinic traits of IPF and found 55% of the genes clustered into the turquoise, black and red modules which were significantly correlated with pulmonary function. In a parallel analysis, 95% of the differentially expressed genes between IPF patients with good and poor prognosis identified by SAM were attributed to these three pulmonary function associated gene modules.
This analytical pipeline highlights the potential applicability of an unsupervised correlation network approach, whereby functional characterization of correlated gene modules provides insight into the molecular mechanisms underlying a clinical trait of a complex pulmonary disease. Lastly, we correlated gene expression levels with survival, which contributed another important feature of IPF. We defined genes met all three selection criteria as “IPF prognostic predictor genes”.

Pathway analysis of the IPF prognostic predictor genes revealed several canonical pathways including T-cell receptor signaling pathway in turquoise module (q = 0.0087); hemoglobin metabolic process and oxygen transport in black module (q = 0.0031 and 0.0032, respectively); and defense response to bacterium and neutrophil degranulation in red module (q = 0.000 and 0.0041, respectively) (Additional file 1: Table S3). The enriched T-cell biology, including ICOS signaling in T-helper cells, CD28 signaling, and T-helper cell differentiation is supported by our prior work, with similar analyses of a smaller gene set demonstrating that decreased expression of CD28, ICOS, LCK, and ITK predicted mortality in patients with IPF [16]. Impaired regulatory T-cells from bronchoalveolar lavage fluid have been strongly correlated with pulmonary dysfunction of IPF patients [28]. Down-regulation of CD28 on circulating CD4 T-cells has been associated with poor outcomes in IPF patients [29]. IL-17A, a cytokine produced by CD4* and gamma-delta* T cells, has been shown to play a critical role in inducing fibrosis in a mouse model [30]. Although the role of the immune system in IPF remains unclear, a large multicenter study has shown that IPF patients treated with prednisone and azathioprine had an increased risk of death and hospitalization compared to those receiving placebo [31]. It remains unknown whether a down-regulated immune system is causally involved in IPF pathogenesis, or is the result of primary lung injury. In addition, a down-regulated immune system could result in a reduced T-cell population [32, 33]. Nevertheless, the down-regulated T-cell pathways or reduced T-cell population can both lead to impaired immune function. These studies are congruent with the functional profile of our IPF genomic model suggesting that suppression of the immune system with medications such as prednisone and azathioprine may worsen the clinical course for IPF patients whose immune systems are already down-regulated.

By evaluating the performance of the genomic model in two independent validation cohorts with different microarray platforms performed at different medical centers, we demonstrate the potential applicability of our findings for real-world use. Notably, the prognostic index (PI) derived from the genomic model showed consistent prognostic prediction specificity in each validation cohort and produced similar mortality hazard estimation across all three cohorts. Genomic model constructed using IPF prognostic predictor genes also displayed concordant fold changes between patients with predicted low- and high-risk prognosis in training and validation cohorts. Furthermore, the PI was able to discriminate between IPF and healthy controls with great accuracy, suggesting a future potential screening tool. However, it is unclear whether the PI can distinguish IPF patients from patients with other pulmonary fibrotic diseases such as nonspecific interstitial pneumonia (NSIP), hypersensitivity pneumonitis (HP), and respiratory bronchiolitis-associated interstitial lung disease (RB-ILD) etc. This question can be addressed in future studies.

While the genomic model developed in this study has been successfully validated in two independent (UCV and UPV) cohorts, certain technical issues and potential clinical confounders require further study. First, there were demographic differences in gender and race between the training and UPV cohorts. The training population was strongly biased towards male patients, while the UPV population was more balanced with respect to gender. There was a greater prevalence of Caucasians in the UPV cohort. Interestingly, the PI scores were higher and indicative of poorer outcome in women overall, while being primarily derived from a male cohort. While IPF is more common in men than women [31], women appear to have improved survival [34, 35]. Although the reasons for this clinical observation are unclear, our results indicate that there may be differential gene expressions between male and female patients with IPF that underlie this observation.

In addition, the different microarray platforms used in different cohort studies might affect the prediction specificity of the model. Notably, the correlation of the PI with pulmonary function in UPV cohort is less strong compared to that in training and in UCV cohorts. We speculated that this observation may be partially attributed to the loss of the 10 classifiers when mapped from Affymetrix to Agilent microarray platform. Another potential confounding factor is the higher rate of lung transplant in the UPV cohort (20 %) compared to the UCV cohorts (7 %). We attempted to adjust for this in our survival analysis by treating transplant as a competing event. Validation with a larger prospective cohort would be beneficial. Finally, the PI cannot be standardized across different microarray platforms, because the gene expression levels in microarray assay were measured by arbitrary fluorescent intensities rather than transcript copy numbers. Therefore, an absolute cut-off or carry-over of PI across individual studies is not feasible, and clinical elaboration of the hazard ratio of PI is impractical at this stage. Future approaches with direct assessment of these IPF prognostic predictor genes could overcome this issue.

Conclusions
We identified an IPF genomic model with both diagnostic and prognostic prediction ability. The unsupervised
WGCNA analysis appears to be a promising approach to elucidate the molecular mechanism underlying IPF progression as an extension of its previous use in oncologic studies [36–38]. The genomic model constructed from the IPF prognostic predictor genes demonstrated robust clinical applications. Functional analysis of the IPF prognostic predictor genes strongly supported the involvement of T-cell immune response in IPF progression [29, 39]. These data continue to support and highlight the use of genomic profiles from the peripheral blood for pulmonary disease.

Additional files

Additional file 1: Additional Methods and Tables. (DOCX 103 kb)

Additional file 2: Figure S1. Detection of gene co-expression modules in training cohort. Gene expression intensities obtained from Exon 1.0 ST Array were normalized. Probe sets were mapped to U133 plus 2.0 Array and filtered as described in Additional file 1. A total of 2,718 unique genes were retained and subjected to R package "Weighted Gene Co-expression Network Analysis (WGCNA)" to identify co-expressed gene modules. A. Optimization and selection of power for adjacency transition of gene-gene correlation matrix (power = 7). B. Cluster dendrogram of the gene co-expression modules represented by different colors. Seven gene co-expression modules were detected by hierarchical clustering using dynamic tree cut algorithm integrated in WGCNA with the following parameters: power=7, minModuleSize=120, mergeCutHeight=0.3. Undistributed genes (genes not correlated with other genes) were collected in Grey module. (PPTX 187 kb)

Additional file 3: Figure S2. Co-expression Network Analysis (WGCNA) genes were retained and subjected to R package 'WGCNA' to identify co-expressed gene modules. A. Optimization and selection of power for adjacency transition of gene-gene correlation matrix (power = 7). B. Cluster dendrogram of the gene co-expression modules represented by different colors. Seven gene co-expression modules were detected by hierarchical clustering using dynamic tree cut algorithm integrated in WGCNA with the following parameters: power=7, minModuleSize=120, mergeCutHeight=0.3. Undistributed genes (genes not correlated with other genes) were collected in Grey module. (PPTX 187 kb)

Additional file 4: Figure S3. Detection of gene co-expression modules in training cohort. Gene expression intensities obtained from Exon 1.0 ST Array were normalized. Probe sets were mapped to U133 plus 2.0 Array and filtered as described in Additional file 1. A total of 2,718 unique genes were retained and subjected to R package "Weighted Gene Co-expression Network Analysis (WGCNA)" to identify co-expressed gene modules. A. Optimization and selection of power for adjacency transition of gene-gene correlation matrix (power = 7). B. Cluster dendrogram of the gene co-expression modules represented by different colors. Seven gene co-expression modules were detected by hierarchical clustering using dynamic tree cut algorithm integrated in WGCNA with the following parameters: power=7, minModuleSize=120, mergeCutHeight=0.3. Undistributed genes (genes not correlated with other genes) were collected in Grey module. (PPTX 187 kb)

Additional file 5: Figure S4. ROC analysis of genomic model for diagnosis prediction. ROC curves of UCV cohort consisting of IPF patients and healthy individuals were plotted based on the Prognostic Index (PI) derived from IPF genomic model. AUIC (Area-Under-Curve) is displayed in the graph. The red line denotes 10 % false alarm (1-Specificity). (PPTX 79 kb)

Abbreviations

IPF: idiopathic pulmonary fibrosis; PBMC: peripheral blood mononuclear cells; WGCNA: weighted gene co-expression network analysis; SAM: significance analysis of microarray; SHR: sub-hazard ratio; ROC: Receiver-Operating-Characteristic analysis; AUC: Area-Under-Curve; PI: prognostic index; PCA: principal component analysis; UCMC: University of Chicago Medical Center; UPMC: University of Pittsburgh Medical Center; UCV: University of Pittsburgh validation cohort; UPV: University of Pittsburgh validation cohort; FVC: % predicted: forced vital capacity percent predicted; DLCO % predicted: diffusion capacity of carbon monoxide percent predicted; CPI: composite physiologic index; FC: fold change; FDR: false discovery rate; CV: cross validation; IPA: Ingenuity pathway analysis; COMET: correlating outcomes with biochemical markers to estimate time-progression in IPF; NSIP: nonspecific interstitial pneumonia; HP: hypersensitivity pneumonia; RBILD: respiratory bronchiolitis-associated interstitial lung disease.

Competing interests

The authors declare that they have no competing interests.

Authors’ contributions

All authors contributed to the conception and design of the study; YH and JO performed the statistical analysis; YH drafted the manuscript; SFM, RV, JO, YAL, JGN, IN contributed to the writing of the manuscript; YH, SFM, RV contributed to this work equally; SMB, MSW collected biological samples; SFM supervised the sample collection; DHH, IN, NKS, MES, RV, SW collected phenotypic data which were further evaluated by IN and RV; All authors read and approved the final manuscript.

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References

1. American Thoracic Society. Idiopathic pulmonary fibrosis: diagnosis and treatment. International consensus statement. American Thoracic Society (ATS), and the European Respiratory Society (ERS). Am J Respir Crit Care Med. 2001;161(2 Pt 1):646–64.
2. American Thoracic Society/European Respiratory Society International Multidisciplinary Consensus Classification of the Idiopathic Interstitial Pneumonias. This joint statement of the American Thoracic Society (ATS), and the European Respiratory Society (ERS) was adopted by the ATS board of directors, June 2001 and by the ERS Executive Committee, June 2001. Am J Respir Crit Care Med. 2002;165(2):277–304.
3. Roghu G, Collard HR, Egan JJ, Martinez FJ, Behr J, Brown KK, et al. An official ATS/ERS/JRS/ALAT statement: idiopathic pulmonary fibrosis: evidence-based guidelines for diagnosis and management. Am J Respir Crit Care Med. 2011;183(6):888–824.
4. Fernandez Perez ER, Daniels CE, Schroeder DR, Sr. Sauver J, Hartman TE, Bartholmai BJ, et al. Incidence, prevalence, and clinical course of idiopathic pulmonary fibrosis: a population-based study. Chest. 2010;137(1):129–37.
5. Martinez FJ, Safrin S, Weycker D, Starko KM, Bradford WZ, King Jr TE, et al. The clinical course of patients with idiopathic pulmonary fibrosis. Ann Intern Med. 2005;142(12 Pt 1):963–7.
6. Swigris JJ, Kuschnier WG, Kelsey JL, Gould MR. Idiopathic pulmonary fibrosis: challenges and opportunities for the clinician and investigator. Chest. 2005;127(1):275–83.
7. King Jr TE, Bradford WZ, Castro-Bernardini S, Fagan EA, Glaspole I, Glassberg MK, et al. A phase 3 trial of pirfenidone in patients with idiopathic pulmonary fibrosis. N Engl J Med. 2014;370(22):2083–92.
8. Richeldi L, du Bois RM, Raghu G, Azuma A, Brown KK, Costabel U, et al. Efficacy and safety of nintedanib in idiopathic pulmonary fibrosis. N Engl J Med. 2014;370(22):2071–82.
9. Ley B, Collard HR, King Jr TE. Clinical course and prediction of survival in idiopathic pulmonary fibrosis. Am J Respir Crit Care Med. 2011;183(4):431–40.
10. Thomeer M, Grutters JC, Wuyts WA, Willemse S, Demedts MG. Clinical use of serum biomarkers of survival in pulmonary fibrosis. Respir Res. 2010;11:89.
11. Boon K, Bailey NW, Yang J, Steel MP, Goshong S, Kervitsky D, et al. Molecular phenotypes distinguish patients with relatively stable from progressive idiopathic pulmonary fibrosis (IPF). PLoS One. 2009;4(4):e5134.
12. Selman M, Carrillo G, Estrada A, Mejia M, Becerril C, Cisneros J, et al. Accelerated variant of idiopathic pulmonary fibrosis: clinical behavior and gene expression pattern. PLoS One. 2007;2(5):e682.
13. Aune TM, Maas K, Moore JH, Olsen NJ. Gene expression profiles in human autoimmune disease. Curr Pharm Des. 2003;9(23):1905–17.

14. Baechler EC, Battilavalla FM, Kaypish G, Gaffney PM, Ortmann WA, Espe KJ, et al. Interferon-inducible gene expression signature in peripheral blood cells of patients with severe lupus. Proc Natl Acad Sci U S A. 2003;100(5):2610–5.

15. Maas K, Chan S, Parker J, Slater A, Moore J, Olsen N, et al. Cutting edge: molecular portrait of human autoimmune disease. J Immunol. 2002;169(1):15–9.

16. Herazo-Mayo JD, Noto I, Duncan SR, Kim S, Ma SF, Tseng GC, et al. Peripheral blood mononuclear cell gene expression profiles predict poor outcome in idiopathic pulmonary fibrosis. Sci Transl Med. 2013;5(205):205ra136.

17. Simon R. Roadmap for developing and validating therapeutically relevant genomic classifiers. J Clin Oncol. 2005;23(29):7332–41.

18. Simon R. Development and validation of biomarker classifiers for treatment selection. J Stat Plan Inference. 2008;138(2):308–20.

19. MacIntyre N, Capo RO, Viegi G, Johnson DC, van der Grinten CP, Brasacoso V, et al. Standardisation of the single-breath determination of carbon monoxide uptake in the lung. Eur Respir J. 2005;26(4):720–35.

20. Miller MR, Hankinson J, Brasacoso V, Burgos F, Casaburi R, Coates A, et al. Standardisation of spirometry. Eur Respir J. 2005;26(2):319–38.

21. Wanger J, Clausen JL, Coates A, Pedersen OF, Brasacoso V, Burgos F, et al. Standardisation of the measurement of lung volumes. Eur Respir J. 2005;26(3):511–22.

22. Wells AU, Desai SR, Rubens MB, Goh NS, Cramer D, Nicholson AG, et al. Idiopathic pulmonary fibrosis: a composite physiologic index derived from disease extent observed by computed tomography. Am J Respir Crit Care Med. 2003;167(7):362–9.

23. Langfelder P, Horvath S. WGCNA: an R package for weighted correlation network analysis. BMC Bioinformatics. 2008;9:559.

24. Tusher VG, Tibshirani R, Chu G. Significance analysis of microarrays applied to the ionizing radiation response. Proc Natl Acad Sci U S A. 2001;98(9):5116–21.

25. Jason PF, Gray RJ. A proportional hazards model for the subdistribution of a competing risk. J Am Stat Assoc. 1999;94:496–509.

26. Simon R, Larr A, Li MC, Nian G, Menereses S, Zhao Y. Analysis of gene expression data using BRB-ArrayTools. Cancer Inform. 2007;3:11–7.

27. Alexa A, Rahnenfuhrer J, Lengauer T. Improved scoring of functional groups from gene expression data by decorrelating GO graph structure. Bioinformatics. 2006;22(13):1600–7.

28. Kotsianidis I, Nakou E, Bouchliou I, Tzouvelekis A, Spanoudakis E, Steipoulos P, et al. Global impairment of CD4 + CD25 + FOXP3 + regulatory T cells in idiopathic pulmonary fibrosis. Am J Respir Crit Care Med. 2009;179(12):121–30.

29. Gilani SR, Vuga LJ, Lindell KO, Gibson KF, Xue J, Kaminski N, et al. CD28 down-regulation on circulating CD4 T-cells is associated with poor prognoses of patients with idiopathic pulmonary fibrosis. PLoS One. 2010;5(1):e9959.

30. Wilson MS, Madala SK, Ramalingam TR, Gochuico BR, Rosas IO, Cheever AW, et al. Bleomycin and IL-1beta-mediated pulmonary fibrosis is IL-17A dependent. J Exp Med. 2010;207(3):535–52.

31. MacIntyre N, Crapo RO, Viegi G, Johnson DC, van der Grinten CP, Brusasco V, et al. Standardisation of the measurement of lung volumes. Eur Respir J. 2005;26(3):511–22.

32. Simon R, Larr A, Li MC, Nian G, Menereses S, Zhao Y. Analysis of gene expression data using BRB-ArrayTools. Cancer Inform. 2007;3:11–7.

33. Simon R. Roadmap for developing and validating therapeutically relevant genomic classifiers. J Clin Oncol. 2005;23(29):7332–41.

34. Schwartz DA, Helmers RA, Galvin JR, Van Fossen DS, Frees KL, Dayton CS, et al. Interferon-inducible gene expression signature in peripheral blood cells of patients with severe lupus. Proc Natl Acad Sci U S A. 2003;100(5):2610–5.

35. Maas K, Chan S, Parker J, Slater A, Moore J, Olsen N, et al. Cutting edge: molecular portrait of human autoimmune disease. J Immunol. 2002;169(1):15–9.

36. Kotsianidis I, Nakou E, Bouchliou I, Tzouvelekis A, Spanoudakis E, Steipoulos P, et al. Global impairment of CD4 + CD25 + FOXP3 + regulatory T cells in idiopathic pulmonary fibrosis. Am J Respir Crit Care Med. 2009;179(12):121–30.

37. Clarke C, Madden SF, Doolan P, Aherne ST, Joyce H, O’Driscoll L, et al. Correlating transcriptional networks to breast cancer survival: a large-scale coexpression analysis. Carcinogenesis. 2013;34(10):2300–8.

38. Zhao H, Cai W, Su S, Zhi D, Lu J, Liu S. Screening genes crucial for pediatric plioctylic astrocystoma using weighted gene coexpression network analysis combined with methylation data analysis. Cancer Gene Ther. 2014;21(10):448–55.

39. Sumida A, Hasegawa Y, Okamoto M, Hashimoto N, Imazumi K, Yatsuya H, et al. TH1/TH2 immune response in lung fibroblasts in interstitial lung disease. Arch Med Res. 2008;39(9):503–10.