Integrated analysis of immunocyte infiltration and differential gene expression in tricuspid aortic valve-associated thoracic aortic aneurysms

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Background: Progressive dilatation is responsible for significant mortality and morbidity in patients with thoracic aortic aneurysms (TAAs). Studies have shown that the development and progression of TAAs are closely related to immune regulatory pathways and genes. Therefore, it is important to understand the immune regulatory mechanisms and biomarkers of TAA dilatation.

Methods: Systematic bioinformatics analysis was applied, including linear models for microarray data (LIMMA) differential expression analyses, principal component analysis (PCA), immunocyte identification, and genetic function enrichment analysis.

Results: Our results showed that both aortic intima-media (AMed) and outer aortic adventitia (AAdv) tissues were closely associated with T cell activation during the process of tricuspid aortic valve (TAV)-associated TAA dilatation. Additionally, the degree of infiltration of resting memory CD4+ T cells was linked to both AAdv and AMed vascular dilation. The core regulators PPTRC, IL1B, CD4, CD3G, and IL2RA were also identified and are closely related to resting memory CD4+ T cell infiltration in this pathological process.

Conclusions: The candidate genes PPTRC, IL1B, CD4, CD3G, and IL2RA were involved in the regulation of resting memory CD4+ T cell tissue infiltration, which is closely related to the process of AAdv and AMed vascular dilation in TAV patients.

Keywords: Thoracic aortic aneurysms (TAAs); pathway enrichment; immunocyte infiltration; integrated bioinformatic analysis

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Introduction

Thoracic aortic aneurysms (TAAs) are a class of vascular diseases with rapid progression and very high mortality and morbidity. Epidemiological studies have suggested that the overall incidence of TAAs is greater than 7.6/100,000 and that TAAs are identified in only 78% of patients before death (1,2). Even with surgical treatment, 16% of patients died within 30 days after surgery, and the 1-, 5-, and 10-year survival rates were 92%, 77%, and 57%, respectively, while the reoperation rate within 10 years was 7.8% (2,3). With the development of 3D printing technology and hybrid surgery, the treatment and prognosis of TAAs have greatly improved (4). However, the mechanisms of TAA occurrence and progression are still unclear.

In recent years, the vigorous development of high-throughput sequencing technology has provided a new opportunity for studying the mechanism of TAAs. Studies have shown that the development and progression of TAAs are closely related to immune regulatory pathways and genes. Kim et al. investigated gene expression profile differences between the thoracic aortas of TAA patients and normal thoracic aortas in organ transplant patients. They found that the differentially expressed genes (DEGs) associated with TAAs were mainly associated with ion transport, cell signal transduction, and immune inflammatory responses (5). Tang et al. analyzed the pathological process of vascular remodeling (changes in the vascular outer diameter) and intima dilation in ascending aorta specimens from TAAs. They found that the transmural inflammatory state of the aorta and the production of interferon-gamma (IFN-γ) in TAAs were closely related to increases in the outer diameter of the aneurysm, thickening of the intima, maintenance of the density of vascular smooth muscle cells, and decreases in matrix proteins (6). Similarly, Sprague et al. believed that the aneurysm dilation process was closely related to the inflammatory state of blood vessels. Vascular injury can stimulate the expression of endothelial cell adhesion molecules and promote the recruitment of inflammatory cells, growth factors, and cytokines, thus affecting the functions of vascular smooth muscle cells and endothelial cells. In addition, these cytokines can induce the production or activation of vasodilation mediators, such as nitric oxide, prostacyclin, endothelial-derived hyperpolarizing factors, and bradykinin, and vasoconstrictors, such as endothelin and angiotensin II, thereby regulating the pathological process of vascular dilation (7). However, aneurysm formation is not always associated with immune inflammation. By comparing the vascular tissue differential gene expression profiles and pathological mechanisms of bicuspid aortic valve (BAV) and tricuspid aortic valve (TAV)-associated aortic aneurysm expansion, Folkersen et al. found that immune mediators were activated only in TAV tissues, whereas BAV tissues did not exhibit a significant immune process. However, the specific mechanism has not yet been elucidated (8). Based on this result, this study aimed to conduct an in-depth investigation of the pathological molecular mechanism of TAV-associated vascular dilation by analyzing DEGs from the whole gene expression profile, immune cell infiltration, and related enrichment pathways in the dilated and nondilated aortic intima-media (AMed) and outer aortic adventitia (AAdv) of TAV patients. This study will provide a new diagnostic or therapeutic target for this disease.

Methods

Data screening and acquisition

We downloaded the GSE26155 dataset from the GEO (https://www.ncbi.nlm.nih.gov/geo/) database for subsequent analysis (9). These data were acquired from microarray expression profiling of AAdv and AMed tissues of TAAs and were published in the Advanced Study of Aortic Pathology (ASAP). A total of 83 vascular tissue specimens were selected, including 46 AMed tissue samples (17 dilated and 29 nondilated samples and 6 boundaries that needed to be excluded) and 37 AAdv tissue samples (12 dilated and 21 nondilated samples and 4 boundaries that needed to be excluded) (8). The corresponding microarray platform was the GPL570 (HG U133 Plus_2) Affymetrix Human Genome U133 Plus 2.0 Array platform (Affymetrix, Santa Clara, CA, USA). In addition, the clinical information corresponding to each sample was downloaded for further analysis.

Data analysis

The data processing flow was as follows: (I) detection of the CEL fluorescence intensity; (II) quality control; (III) background processing using the robust multiarray average (RMA) method; (IV) processing of missing probe values by log2 transformation and the k-nearest neighbor (kNN) algorithm; (V) gene annotation using probe names; (VI) differential expression analysis of expression profiles using linear models for microarray data (LIMMA) (10); and (VII) examination of the data structure by principal component analysis (PCA). Cross-checking was applied to identify DEGs, and then the Benjamini-Hochberg method was
used to adjust the statistical P values of the false discovery rate (FDR) to calculate the expression fold change (FC) (a log2FC >1.0 and a corrected P<0.05 represented DEGs) (10). All of the data were obtained from the GEO database, and a research ethics application was not needed for this study.

**Gene Ontology (GO) and pathway enrichment analysis of gene sets**

The GO of the AMed- and AAdv-associated DEGs was obtained based on analysis using the clusterProfiler algorithm (11). The results for co-DEG-related GO and Kyoto Encyclopedia of Genes and Genomes (KEGG)-enriched pathways were obtained based on an analysis of the MetaScape gene annotation and retrieval platform (http://metascape.org/gp/index.html) (12).

**Analysis of immune infiltration**

CIBERSORT (https://cibersort.stanford.edu/) is an immune cell subtype infiltration calculation algorithm that was developed based on linear support vector regression (13). Users can comprehensively estimate the infiltration level of each cell subtype from chip expression profile and RNA-seq expression data. The parameters applied in this study were as follows: (I) gene expression values corrected by the RMA algorithm; (II) 1,000 deconvolutions (Perm); and (III) P<0.05 for differential subtypes.

**Association analysis between core genes and differentially infiltrating immune cell subtypes**

In addition, by constructing the co-DEG-related KEGG pathway network, pathway-rich genes of interest were selected for protein–protein interaction (PPI) network analysis. The node correlation degree in the network was calculated using the STRING database (https://string-db.org/) (14) and CytoScape software (15) to identify candidate regulatory factors. To further clarify the associations between core genes and immune genes, we performed a Pearson correlation clustering analysis of candidate genes and differential cell subtype infiltration values.

**Results**

**Data acquisition and pretreatment**

The results of the difference analysis suggested that (I) in the AAdv tissues, compared with the nondilated group, a total of 1,190 differential mRNAs (3 downregulated and 1,187 upregulated) were present in the dilated group; and (II) in the AMed tissues, 173 DEGs were present between the dilated group and nondilated group (7 downregulated and 166 upregulated). The distributions of DEGs in AMed and AAdv tissue samples are shown in Figure 1A and http://cdn.amegroups.cn/static/application/d1026d3979cb63f6749fde5efbe1d50d/atm.2020.03.05-1.pdf.

**Functional enrichment analysis of DEGs**

In the enrichment analysis, we found that GO:0042110–T cell activation (P=1.24E-19, n=75), GO:0002694–regulation of leukocyte activation (P=1.25E-18, n=78), and GO:0001819–positive regulation of cytokine production (P=1.38E-16, n=70) were closely related to the biological process (BP) of DEGs associated with AAdv dilation, whereas GO:0031012–extracellular matrix (P=9.49E-11, n=60), GO:0009897–external side of the plasma membrane (P=2.05E-10, n=35), and GO:0062023–collagen-containing extracellular matrix (P=3.32E-10, n=53) were closely related to the cellular component (CC) of DEGs associated with AAdv dilation. GO:0005201–extracellular matrix structural constituent (P=1.60E-10, n=32), GO:0005539–glycosaminoglycan binding (P=4.68E-05, n=26), and GO:0008201–heparin binding (P=5.88E-04, n=19) were closely related to the molecular function (MF) of DEGs associated with AAdv dilation (Figure 1B and http://cdn.amegroups.cn/static/application/573636b8df0cab7b2c648958a2ea3928/atm.2020.03.05-2.pdf). Similarly, GO:0042110–T cell activation (P=6.61E-17, n=27), GO:0001819–positive regulation of cytokine production (P=7.35E-15, n=25), and GO:0045785–positive regulation of cell adhesion (P=3.03E-14, n=23) were closely related to the BP of DEGs associated with AMed dilation, while GO:0009897–external side of the plasma membrane (P=4.76E-11, n=15), GO:0030667–secretory granule membrane (P=1.01E-08, n=15), and GO:0043235–receptor complex (P=7.26E-08, n=14) were closely related to the CC of DEGs associated with AMed dilation. GO:0042287–major histocompatibility (MHC) protein binding (P=5.78E-07, n=6), GO:0019955–cytokine binding (P=1.55E-06, n=8), and GO:0004896–cytokine receptor activity (P=4.12E-05, n=6) were closely related to the MF of DEGs associated with AMed dilation (Figure 1C and http://cdn.amegroups.cn/static/application/573636b8df0cab7b2c648958a2ea3928/atm.2020.03.05-2.pdf).
Figure 1 The differential expression and genetic function enrichment analysis with regard to both aortic intima-media (AMed) and aortic adventitia (AAdv) dilation. The volcano plot in Figure 1A presents the differentially expressed genes (DEGs) for the comparison of dilated and nondilated AMed or AAdv samples. Figure 1B,C presents the Gene Ontology (GO) and Kyoto Encyclopedia of Genes and Genomes (KEGG) pathway enrichment analyses were performed based on the clusterProfiler and MetaScape databases, respectively. The sizes of the dots represent the counts of enriched DEGs, and the colors of the dots represent the adjusted P value for the GO term enrichment, while the dot size represents the negative Log(P value) for KEGG maps.
Analysis of the MetaScape database showed that AAdv-associated DEGs were mainly associated with type 1 T helper (Th1) and Th2 cell differentiation (enrichment score =5.31, \(P=2.51\times 10^{-11}\), n=25), the PI3K-Akt signaling pathway (enrichment score =2.77, \(P=4.00\times 10^{-10}\), n=46), the tumor necrosis factor (TNF) signaling pathway (enrichment score =4.0, \(P=4.62\times 10^{-8}\), n=21), and other pathways (Figure 1B and http://cdn.americanets.cn/static/application/4196621e963294ffbd07d0dc69f03162/atm.2020.03.05-3.pdf). AMed-associated DEGs were mainly associated with Th17 cell differentiation (enrichment score =22.50, \(P=1.80\times 10^{-16}\), n=15), cell adhesion molecules (CAMs) (enrichment score =14.14, \(P=7.23\times 10^{-15}\), n=17), leishmaniasis (enrichment score =23.92, \(P=9.49\times 10^{-14}\), n=12), and other pathways (Figure 1C and http://cdn.americanets.cn/static/application/4196621e963294ffbd07d0dc69f03162/atm.2020.03.05-3.pdf).

**Identification and functional enrichment of co-expressed DEGs in the dilation process of AMed and AAdv tissues**

By evaluating the intersection set (Figure 2), 107 DEGs shared by both AMed and AAdv tissues were obtained (Figure 2A). PCA showed that co-DEGs could significantly distinguish whether AAdv and AMed tissues contained dilated blood vessels (Figure 2C). Based on cluster analysis, the 107 DEGs were divided into five categories. Functional enrichment analysis showed that GO:0030334~regulation of cell migration, GO:0045785~positive regulation of cell adhesion, GO:0002322~B cell proliferation involved in the immune response, GO:0045321~leukocyte activation, and GO:0002253~activation of the immune response were closely related to the functional enrichment of each gene set (Figure 2B and http://cdn.americanets.cn/static/application/bb4fe75016658b7f263c56ab52348e529/atm.2020.03.05-4.pdf).

**Core gene identification**

Based on the KEGG pathway enrichment network, we found that the co-DEGs of AAdv and AMed tissues were closely associated with T cell-associated immune pathways (Figure 3A). We further constructed a PPI network for genes enriched in T cell-associated immune pathways and found that protein tyrosine phosphatase receptor type C (PTPRC) (degree =11), interleukin-1B (IL1B) (degree =7), CD4 (degree =7), CD3G (degree =7), and IL-2 receptor alpha chain (IL2RA) (degree =11) were closely related to the progression of aortic dilation (Figure 3B). In addition, compared with the nondilated group, these 5 core genes were highly expressed in both the AAdv tissues and the AMed tissues in the dilated group (all \(P<0.05\); Figure 3C, D).

**Analysis of immune infiltration**

The overall immune infiltration profiles of AAdv and AMed vascular tissues are shown in Figure 4A, B. Resting memory CD4 T cells (AAdv \(P=4.36\times 10^{-4}\), AMed \(P=1.79\times 10^{-3}\)), regulatory T cells (Tregs) (AAdv \(P=0.027\), AMed \(P=1.20\times 10^{-2}\)), naïve B cells (AAdv \(P=0.044\), AMed \(P=1.90\times 10^{-2}\)), and monocytes (AAdv \(P=0.051\), AMed \(P=2.10\times 10^{-2}\)) demonstrated significant differential infiltration in both dilated and nondilated AAdv and AMed tissues (Figure 4C, D, http://cdn.americanets.cn/static/application/a0d90651ac6c292730150444766be6bb/atm.2020.03.05-5.pdf).

Through correlation analysis, we also found that the core regulatory genes (including PTFRC, IL1B, CD4, CD3G, and IL2RA) had strong correlations with the degree of infiltration of resting memory CD4 T cells (AAdv: PTPRC coefficient =0.71, IL1B coefficient =0.52, CD4 coefficient =0.59, CD3G coefficient =0.78, IL2RA coefficient =0.60; AMed: PTPRC coefficient =0.82, IL1B coefficient =0.62, CD4 coefficient =0.78, CD3G coefficient =0.73, IL2RA coefficient =0.64) in AAdv and AMed tissues (Figure 5A, B, http://cdn.americanets.cn/static/application/5c02a2b7023c218db1d6026daee4f0d1/atm.2020.03.05-6.pdf).

**Discussion**

The association between the pathogenesis of TAAs and the immune inflammatory response has always been a popular research topic. Our study found that both AAdv and AMed tissues were closely associated with T cell activation during the process of vascular dilation. By further building a network of disease mechanisms, our results also suggested that PTPRC, IL1B, CD4, CD3G, and IL2RA may be the core regulatory genes of vascular dilation; these genes were closely related to the degree of infiltration of resting memory CD4 T cells in AAdv and AMed vascular tissues, indicating that these genes may be important regulatory mediators in TAA pathogenesis.

An increasing number of researchers believe that TAAs are immune inflammatory diseases, and the risk of disease increases with increasing age (16). Compared with other aneurysms, the pathological changes in the outer adventitia and media of the aortic wall are more closely associated with the immune inflammatory response, especially those in TAV-
associated TAA vascular tissues (16,17). Among these changes, T cell activation is the most important molecular mechanism. Itani et al. found that angiotensin II can promote the infiltration of leukocytes [CD45 (+)], memory T cells [CD3 (+)/CD45 Ro (+)] and T lymphocytes (CD3 (+) and CD4 (+)) in thoracic aortic tissues, increase activated CD4+ and CD8+ T cells in the circulation, and increase the production of IL17a and IFN-γ, suggesting that functional activation of T cells and their subpopulations is associated with hypertension-induced vascular remodeling and dilation (18). Similarly, Ju et al. found that the cytokine IL6 can induce Th17 lymphocytes to aggregate in dilated vascular tissues through the transcription-3 signaling pathway in an angiotensin II perfusion-induced vascular dissection model. At the same time, these lymphocytes promoted macrophage recruitment and mediated the development of vascular dissection through the transcription-3 signaling pathway (19). Ye et al. found that CD4 T cell infiltration was closely related to aortic root inflammation and the degree of root dilation in TAA patients (20).
The comparison between dilated and nondilated vascular walls revealed that resting memory CD4+ T cells were significantly infiltrated in the dilated AAdv and AMed tissues, especially the AAdv tissues. Crotty believes that memory CD4 T cells have a degree of plasticity and can differentiate into other subtypes of T cells; however, no experimental data have confirmed the plasticity or differentiation ability of resting memory CD4 T cells (21). McKinstry et al. suggested that memory CD4 T cells can not only differentiate into subcells such as secretory T cells (Th) but also secrete a large amount of cytokines to recruit immune cells and enhance the immune response, facilitating the immune response of CD8+ T cells and B cells (22). Sbrana et al. found that the degree of CD4+ T lymphocyte infiltration was increased and the production of IFN-γ, IL-17a, and IL-21 was increased in the vascular tissues of patients with ascending aortic dilation (23). Jones et al. found that compared with other immune cells, effector memory and central memory CD4+ T cells showed higher levels of glycolysis and oxidative phosphorylation and a higher metabolic capacity, and regulation of this metabolic capacity and cell recruitment were closely related to early activation of naïve CD4+ T cells (24).

In addition, our study suggested that IL1B, CD3G, CD4, IL2RA, and PTPRC may be the core regulatory genes of disease progression in TAV-associated TAs and that these genes are positively correlated with the degree of infiltration of resting memory CD4+ T cells. Cochain et al. performed single-cell sequencing on mouse aortic arch CD45+ macrophages in a low-fat diet group and a high-fat diet group and found that IL1B was closely associated with the inflammatory status of aortic arch endothelial cells (25). Yang et al. found that miR-30c could participate in the process of vascular dilation of abdominal aortic aneurysms (AAAs) by targeting IL1B, phosphatidylinositol-4,5-bisphosphate 3-kinase catalytic subunit delta (PIK3CD), and Ras-related C3 botulinum toxin substrate 2 (RAC2) (26).
CD3G is an important regulatory factor during the process of T cell development and differentiation (27) and is highly expressed in the torn vascular tissues of patients with acute aortic dissection (D13). Goudy et al. showed that IL2RA mutations can significantly affect the function of regulatory and effector T cells, which are associated with lymphocyte proliferation and T cell activation and are the key mediators of immune function and homeostasis in the body (28).

Figure 4 Detection of immunocyte infiltration and the significant immunocyte subtypes. The hierarchical clustering map of Figure 4A,B shows the immunocyte infiltration difference between dilated and nondilated AMed or AAdv samples, respectively. The boxplots of Figure 4C,D presenting the significantly infiltrated immunocyte subtypes involved in AMed or AAdv dilatation.
Figure 5 Interaction analysis of candidate genes and significantly infiltrated immunocyte subtypes. Figure 5A,B shows the relationship between immunocytes and hub genes was presented by a clustering heatmap and circus plot with regard to AMed or AAdv dilatation, respectively.
Similarly, a study by Belot et al. also found that methylation of the IL2RA promoter region can affect IL2RA expression and T cell activation (29). PTPRC (CD45) is an important marker of macrophage and leukocyte activation. Gallo et al. found that high CD45 expression in TAA patients was positively correlated with increased monocyte infiltration in the vascular walls and increased IFN-γ, IFN-inducible protein 10, and IFN-induced T cell α chemokine levels in the circulation (30).

**Conclusions**

In summary, our study found that during the process of AAdv and AMed vascular dilation in TAV-associated TAAs, PPTRC, IL1B, CD4, CD3G, and IL2RA were involved in the regulation of resting memory CD4 T cell tissue infiltration, which was closely related to the process of vascular dilation. Most of those candidate regulators were verified in previous studies. However, several limitations remain. First, although the correlations among these candidate markers and immunocyte infiltration in TAV-associated TAAs were identified, further experimental evidence concerning the mechanism is still needed. Second, the mechanism of TAV-associated vascular dilation is complicated; thus, immunocyte infiltration may be important but not essential.

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