**Introduction**

Idiopathic pulmonary fibrosis (IPF) is a form of chronic progressive fibrosing interstitial lung disease characterized by the histopathologic pattern of usual interstitial pneumonia. Patients with IPF usually present with progressive respiratory symptoms with periods of acute exacerbations, resulting in significant morbidity and mortality. Prognosis of patients with IPF is poor with median survival of only two to five years (1). The exact etiology is still unknown despite extensive research effort. There is no medication with proven efficacy for IPF and current standard treatment focuses mainly on symptom relief although lung transplantation could be an option for selected patients (2).

Venous thromboembolism (VTE), which consists of deep venous thrombosis (DVT) and pulmonary embolism (PE), is one of the common medical problems with approximately 900,000 new and recurrent cases diagnosed every year in the United States (3). Traditional risk factors of VTE include cancer, trauma, surgery, immobilization hospitalization and use of certain medications (4, 5). More recently, chronic inflammation has been recognized as an independent risk factor for VTE as increased in-

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**Risk of venous thromboembolism in patients with idiopathic pulmonary fibrosis: a systematic review and meta-analysis**

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**Abstract.** Background: Recent studies have suggested that patients with idiopathic pulmonary fibrosis (IPF) may have a higher risk of venous thromboembolism (VTE) compared to general population even though the results were inconsistent. Objective: To investigate the risk of VTE among patients with IPF. Methods: Comprehensive literature review using MEDLINE and EMBASE database were performed to identify studies that compared the risk of VTE among patients with IPF to general population. Effect estimates from each study were combined together using random effect model, generic inverse variance method of DerSimonian and Laird. Results: Out of 510 retrieved articles, 5 studies met the inclusion criteria and were included in the meta-analysis. A significant risk of VTE in patients with IPF was observed with the pooled risk ratio of 2.11 (95% confidence interval, 1.28-3.48). The heterogeneity was moderate with I² of 64%. Conclusion: An approximately 2-fold increased risk of VTE among patients with IPF was observed in this meta-analysis. (Sarcoidosis Vasc Diffuse Lung Dis 2018; 35: 109-114)

**Key words:** idiopathic pulmonary fibrosis, venous thromboembolism, thrombosis, deep vein thrombosis, pulmonary embolism
cidence of VTE has been observed in several chronic inflammatory conditions, such as rheumatoid arthritis, psoriasis, systemic vasculitis and inflammatory myositis (6-9).

Patients with IPF may be at an elevated risk of VTE as well due to their increased systemic inflammatory burden. In addition, patients with advanced respiratory symptoms are also likely to have limited mobility, resulting in venous stasis that can predispose to thromboembolism. In fact, several epidemiologic studies have suggested an association between IPF and VTE although the results are inconsistent (10-14). To further investigate this possible association, we performed a systematic review and meta-analysis of studies that compared the VTE risk in patients with IPF to subjects without IPF.

**Method**

**Search strategy**

Both investigators independently searched published articles in MEDLINE and EMBASE database from inception to February 2017 using the search terms for idiopathic pulmonary fibrosis and venous thromboembolism as described in online supplementary data without any language restriction. References of selected articles were also manually searched for additional studies.

**Inclusion criteria**

Studies were eligible for this meta-analysis if they met these inclusion criteria: 1) Cohort (either prospective or retrospective), case-control study or cross sectional study published as original study to evaluate the association between IPF and VTE, 2) odds ratios (OR), relative risk (RR), hazard ratio (HR), and standardized incidence ratio (SIR) with 95% confidence intervals (CI) or sufficient raw data to calculate these ratios were provided, and 3) subjects without IPF were used as comparators in cohort and cross-sectional study while subjects without VTE were used as controls in case-control study.

Study eligibility was independently evaluated by the two investigators. Any disagreement was resolved by mutual consensus. The quality of each study was appraised using the Newcastle-Ottawa quality scale (15). This scale assesses each study in three domains including 1) the representativeness of the subjects, 2) the comparability between the study groups, and 3) ascertainment of the exposure of interest for case-control study and the outcome of interest for cohort study. The modified version of Newcastle-Ottawa scale as described by Herzog et al. was used for cross-sectional study (16).

**Review process and data extraction**

The two study investigators independently reviewed the titles and abstracts of all retrieved articles. Articles that clearly did not fulfill the inclusion criteria were excluded. Only potentially relevant articles underwent full-text review to determine the eligibility. A standardized data collection form was used to extract the following information from the included studies: first author’s name, year of publication, year of study, country where the study was conducted, study design, source of population, number of subjects, baseline characteristics of the subjects, methods used to identify IPF and VTE, and effect estimates. This data extraction process was also performed by both investigators to ensure the accuracy.

**Statistical analysis**

All statistical analyses were performed using Review Manager 5.3 software from the Cochrane Collaboration (London, UK). The pooled RR of VTE in IPF patients in comparison to subjects without IPF was calculated using generic inverse method of DerSimonian and Laird (17). Random effect model was used given the high likelihood of between-study variance due to the difference in underlying population and methodology. As the outcome of interest was relatively uncommon, the ORs of cross-sectional study and case-control study were used as an estimate to pool with the RR from cohort study. Cochran’s Q-test, which is supplemented by I² statistic, was used to evaluate the statistical heterogeneity. This I² statistic quantifies the proportion of the total variation across studies, that is, due to true heterogeneity rather than chance. A value of I² of 0% to 25% represents insignificant heterogeneity, more than 25% but ≤50% represents low heterogeneity, more than 50% but ≤75% represents moderate heterogeneity, and more than 75% represents high heterogeneity(18).
**Results**

The initial search yielded 510 articles, all of which underwent title and abstract review. The majority of them were excluded at this step as they were case report, letter to editor, review article or interventional study which clearly did not fulfill our inclusion criteria. A total of 15 studies underwent full-length article review and 10 of them were excluded at they did not include patients with IPF or did not report the outcome of interest. Therefore, a total of five studies met our inclusion criteria (four cohort studies and one cross-sectional study (10-14)) and were included in the meta-analysis. Baseline characteristics of the included studies are summarized in table 1.

| Country         | Study design          | Year of publication | Participants                                                                 | Diagnosis of IPF                        | Diagnosis of VTE                        | Number of participants | Mean age (years) | Percentage of male | Confounder adjusted | Quality assessment (Newcastle-Ottawa scale) | Quality assessment (Newcastle-Ottawa scale) |
|-----------------|-----------------------|---------------------|------------------------------------------------------------------------------|-----------------------------------------|----------------------------------------|------------------------|------------------|-------------------|---------------------|------------------------------------------|------------------------------------------|
| Hubbard et al. [10] | UK Retrospective cohort | 2008                | Cases were identified from The Health Improvement Network database from general practitioners across the UK from 1991 to 2003. Comparators were sex and age-matched randomly selected from the same database. | Diagnostic code from the registry       | Diagnostic code from the registry     | Case 920 Comparator 3,593 | Case 71          | Case 62            | Age, sex, smoking habit and medications | Selection: 4 stars, Comparability: 2 stars, Outcome: 2 stars | Selection: 4 stars, Comparability: 2 stars, Outcome: 2 stars |
| Collard et al. [11] | USA Retrospective cohort | 2012                | Cases were identified from two US insurance databases from January 1, 2001 to September 30, 2008. Comparators were sex and age-matched randomly selected from the same database. | Diagnostic code from the registry       | Diagnostic code from the registry     | Case 9,286 Comparator 9,286 | Case 74          | Case 62            | None                                            | Selection: 4 stars, Comparability: 1 star, Outcome: 2 stars | Selection: 4 stars, Comparability: 1 star, Outcome: 2 stars |
| Sprunger et al. [12] | USA Cross-sectional   | 2012                | Cases were identified from the US cause-of-death mortality database of the National Centre for Health Statistics from 1988 to 2007. Comparators were the rest of subjects in the database. | Diagnostic code from the registry       | Diagnostic code from the registry     | Case 218,991 Comparator 46,450,489 | N/A              | N/A               | Age, sex, and year of death                        | Selection: 4 stars, Comparability: 1 star, Outcome: 2 stars | Selection: 4 stars, Comparability: 1 star, Outcome: 2 stars |
| Dalleywater et al. [13] | UK Retrospective cohort | 2014                | Cases were identified from The Health Improvement Network database from general practitioners across the UK from 2000 to 2013. Comparators were sex and age-matched randomly selected from the same database. | Diagnostic code from the registry       | Diagnostic code from the registry     | Case 3,211 Comparator 12,307 | Case 76          | Case 75            | Age and sex                                | Selection: 4 stars, Comparability: 2 stars, Outcome: 2 stars | Selection: 4 stars, Comparability: 2 stars, Outcome: 2 stars |
| Navaratnam et al [14] | UK Prospective cohort  | 2014                | Cases were recruited from five teaching hospitals and eight district general hospitals in the Greater Trent region and Wales between January 2010 and February 2012. Comparators were sex and age-matched recruited from the same hospitals. | Diagnostic code from the registry       | Diagnostic code from the registry     | Case 211 Comparator 256 | Case 74          | Case 75            | Selection: 4 stars, Comparability: 2 stars, Outcome: 2 stars | Selection: 4 stars, Comparability: 2 stars, Outcome: 2 stars | Selection: 4 stars, Comparability: 2 stars, Outcome: 2 stars |

*USA indicates United States of America; UK, United Kingdom; N/A, Not available; IPF, idiopathic pulmonary fibrosis.*
Our meta-analysis revealed a significantly increased risk of VTE among patients with IPF with the pooled RR of 2.11 (95% CI, 1.28-3.48). The heterogeneity was moderate with $I^2$ of 64%. Figure 1 demonstrates the forest plot of this study.

**Evaluation for publication bias**

The funnel plot is shown in figure 2. It is symmetric and does not suggest the presence of publication bias in favor of positive study.

**Sensitivity analysis**

Since the statistical heterogeneity was not low in this meta-analysis, a sensitivity analysis was performed by excluding one study at a time to investigate the effect of each study on the overall heterogeneity. Interestingly, exclusion of the study by Sprunger et al. (12), the only cross-sectional study, dramatically reduced $I^2$ to 0%. The pooled effect estimate from this sensitivity analysis remained essentially unchanged (RR 2.58; 95% CI, 1.66-4.02).

**Discussion**

This is the first meta-analysis to demonstrate a significantly increased risk of VTE among patients with IPF. The risk is increased by approximately 2-fold. The exact mechanism behind the increased risk is not known but several possible explanations have been proposed.

First, the higher inflammatory burden in patients with IPF could be responsible for the increased tendency for blood clot. The underlying mechanisms of inflammation-induced thrombosis include up-regulation of coagulation factors and down-regulation of anticoagulants/fibrinolysis by inflammatory cytokines as well as injury to endothelial cells by free radicals and oxidative stress. Further evidence to support that inflammation plays an important role in the development of VTE is that VTE is observed more frequently when the inflammatory disease is active (19, 20).

Second, VTE and IPF may share a common origin. Thrombin, a key enzyme in coagulation cascade, could be the link as it is also an inducer of fibrogenic cytokines (21) and has been found in increased concentration in bronchoalveolar lavage from patients with fibrotic lung disease (22). In fact, a recent study has suggested that recombinant human thrombomodulin, which could form a reversible complex with thrombin to inactivate coagulation cascade, is effective for treatment of acute exacerbation of IPF (23).
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Third, it is also possible that the increased risk is simply due to immobility as most patients with IPF have respiratory symptoms and reduced exercise capacity (24).

Fourth, the increased risk could be due to exposure to glucocorticoids, the medication often used to treat acute exacerbation of IPF. In vitro studies have demonstrated that glucocorticoids increase levels of coagulation factors and fibrinogen (25, 26). In addition, a dose-response relationship between exposure to glucocorticoids and incidence of VTE has been demonstrated by a recent population-based study (27).

Although the literature review process was rigorous and the included studies were of high quality, this meta-analysis has some limitations. Therefore, the interpretation of the results needs to be performed with caution. First, most of the included studies were medical registry-based studies, with the exception for the study by Navaratnam et al. (14), which were inherently at risk of inaccurate coding for both IPF and VTE. As a result, the completeness of case/event identification and the accuracy of diagnosis were limited. Second, statistical heterogeneity was not low in this study. Interestingly, the I² dropped dramatically to 0% with the sensitivity analysis that excluded the study by Sprunger et al. (12). We suspect that the difference in study design was responsible for the between-study heterogeneity as the study by Sprunger et al. was the only cross-sectional study. Third, this is a meta-analysis of observational studies that can only demonstrate an association but cannot confirm causality. It is possible that confounders that were not adjusted in the primary studies, rather than IPF itself, are accountable for the increased risk of VTE. Last, surveillance bias may also play a role. It is possible that patients with IPF may have more medical examinations because of their chronic illness. Also, they may have more imaging studies of the thorax due to their respiratory symptoms.

Conclusion

An approximately 2-fold increased risk of VTE among patients with IPF was observed in this meta-analysis.

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Search Strategy

Database: Ovid MEDLINE
1. interstitial lung disease.mp. or exp Lung Diseases, Interstitial/
2. pulmonary fibrosis.mp. or exp Pulmonary Fibrosis/
3. or/1-2
4. exp Thromboembolism/
5. Thromboembolism.mp.
6. exp Venous Thrombosis/
7. venous thrombosis.mp.
8. exp Pulmonary Embolism/
9. pulmonary embolism.mp.
10. or/4-9
11. 3 and 10

Database: EMBASE
1. idiopathic pulmonary fibrosis.mp. or exp fibrosing alveolitis/
2. fibrosing alveolitis.mp.
3. pulmonary embolism.mp. or exp lung embolism/
4. deep vein thrombosis.mp. or exp deep vein thrombosis/
5. venous thromboembolism.mp. or exp venous thromboembolism/
6. exp thromboembolism/ or thromboembolism.mp.
7. or/3-6
8. or/1-2
9. 7 and 8