Research Article

Association between the Platelet-Derived Growth Factor/Platelet-Derived Growth Factor Receptor System and Risk of Rheumatoid Arthritis: A Systematic Review and Meta-Analysis

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This research examines the association between the platelet-derived growth factor/platelet-derived growth factor receptor (PDGF/PDGFR) system and rheumatoid arthritis (RA) susceptibility through a comprehensive search of the PubMed database to study the expression of the PDGF/PDGFR system in RA. Review Manager software version 5.3 was used for statistical analysis. Six eligible studies published in the English language were included, including 108 rheumatoid arthritis cases and 85 controls with the corresponding 126 and 97 tests, respectively, relating the expression of the PDGF/PDGFR system to the risk of RA. The overall results indicated a significant association between the PDGF/PDGFR system expression and RA (OR = 5.25, 95% CI: 3.00-9.18, p < 0.0001), RA patients in Asian countries (OR = 4.13, 95% CI = 2.04-8.39, p < 0.0001) and in Western countries (OR = 9.18, 95% CI = 2.04-83.9, p = 0.03), and only PDGF expression in RA patients (OR = 5.28, 95% CI = 2.73-10.21, p < 0.0001). Thus, only the PDGF expression was insignificantly associated with RA susceptibility (OR = 9.25, 95% CI = 0.63-136.30, p = 0.11). Hence, the PDGF/PDGFR system most likely contributes to susceptibility to RA.

1. Introduction

Rheumatoid arthritis (RA) is a chronic systemic autoimmune disease with unmet treatment needs that disrupts the lining of the synovial joints, resulting in painful joints and damage to other organs, and can lead to progressive disability, early mortality, and socioeconomic burden [1, 2]. The arm and leg joints affected by RA are characterized by synovial inflammation and hyperplasia, destruction of the bones and cartilages, and production of rheumatoid factor and antibody of anticitrullinated protein with clinical manifestations such as pain, swelling, tenderness, and morning stiffness [2, 3]. Other RA-related conditions are systemic disorders of the pulmonary, cardiovascular, nervous, and skeletal systems [2].

RA symptoms and its treatment goals and outcomes vary from person to person. The diagnosis of RA is employed to categorize the diseases into stages as I, II, III, and IV depending on whether the patient’s condition is mild, moderate, severe, and at the end stage, respectively [4, 5]. Treatment options of RA comprise biologic modifiers, receptor-targeted therapies in recognition of the role of cytokines in RA, and medicines for B cell depletion in RA patients [6, 7]. However, therapies for RA treatment are inadequate. As a result, RA patients are faced with a terminal illness that is characterized by progressive destruction of their joints, thereby resulting in pain and poor quality of life [8]. Though reliable biomarkers of RA are few, currently, there are many biomolecules such as anticyclic citrullinated protein antibodies, serum rheumatoid factors, and cytokines...
that are expressed in the serum and synovial tissues at RA patients’ joints and are shown in varying degrees as potential biomarkers of RA [6, 9].

There is a continuing effort to identify candidates of biomarkers in RA, and so far, some biomarkers of RA are likely to be appropriate for target therapy. The platelet-derived growth factor/platelet-derived growth factor receptor (PDGF/PDGFR) system comprises 4 growth factors: PDGFA, PDGFB, PDGFC, and PDGFD, with 2 receptors as PDGFRα and PDGFRβ [10]. The PDGF/PDGFR system is associated with rheumatic diseases, cancers, retinal diseases, fibrotic diseases, and cardiovascular diseases [11–13]. Notably, the PDGF/PDGFR system is expressed in RA-fibroblast-like synoviocyte cells and RA-synovial membranes. It is also involved in proliferation and migration and cell morphological changes and has been shown to stimulate the production of matrix metalloproteinases-1 and proinflammatory cytokines in these cells [11, 14–16].

Though the cohorts’ studies on the expression of the PDGF/PDGFR are few, there are significant studies of the overexpression of the PDGF/PDGFR system in RA without evidence of any marked association between the PDGF/PDGFR system and RA [14, 17–21]. Therefore, a meta-analysis of the PDGF/PDGFR system’s association with RA is needed to develop a fit-for-purpose-targeted treatment of RA. Therefore, this review focuses on assessing the PDGF/PDGFR system’s association with rheumatoid arthritis.

2. Methods

2.1. Identification of Eligible Studies and Data Extraction. A comprehensive search of all English-published literature in June 2021 or earlier, in the PubMed database, to select findings on the expression of platelet-derived growth factors and their receptors in tissues of RA patients and nonarthritic subjects was done for inclusion in this study. The search terms were “platelet-derived growth factor” OR “platelet-derived growth factor receptor” OR “PDGF” or “PDGFR” AND “rheumatoid arthritis.” Two reviewers, LF and WS, independently identified fit-for-purpose studies sequentially by scanning titles/abstracts and reading the full text. All uncertainties and differences were resolved by consensus after rechecking data sources and data conformity to inclusion and exclusion of this research by the third reviewer, WW.

2.2. Inclusion and Exclusion Criteria. A study was placed for the meta-analysis to determine if it met the following criteria: (i) it is a case-control study that measured the expression of the platelet-derived growth factors and/or their receptors in the tissues of nonarthritic and RA patients and (ii) it contained original and sufficient data to calculate odds ratios (ORs). For exclusion criteria of studies, the following was used: (i) studies in which the number of study participants could not be ascertained, (ii) relevant studies in which there is no sufficient data to estimate the number of patient samples that showed significant expression of PDGF and/or PDGFR, (iii) studies using only cell lines and/or animal models or research that is unrelated to the objectives of this study, and (iv) review, expert opinions, editorials, conference abstracts, letters, and case reports involving 1 patient.

2.3. Data Extraction. The first and second reviewers detailed the expression of PDGF and/or PDGFR in the tissues of RA patients and the control, which are nonarthritic subjects from eligible publications using a standard and relevant data extraction form and presented in an orderly manner to facilitate easy comparison and analysis of the extracted data. The following information was collected from the eligible research for the meta-analysis: the surname of the first author, year of publication, country of location of the study subjects, the number RA and control subjects, the cohort of the study, estimation of the number of study RA, and the control study subjects that have shown to express PDGF and/or PDGFR from graphs and tables. All the data were recorded in an Excel spreadsheet for analysis.

2.4. Statistical Analysis. Meta-analysis was done, employing the Review Manager version 5.3 to determine the significance of the association between susceptibility of RA and the expression of the PDGF/PDGFR system in the tissues of RA patients. This was analyzed using the estimated OR of all the eligible studies at a 95% confidence interval (CI). In addition, the pooled effect sizes were obtained from forest plots of the OR of individual studies at 95% CI to evaluate the association between expression and no expression of the PDGF/PDGFR system among combined dichotomous sets of RA and nonarthritic subjects. The forest plots gave the overall effect of growth factors (PDGF) and their receptors (PDGFR) on RA, the effect of only PDGF and only PDGFR on RA, and the effect size of study subjects in Western and Asian studies countries has been analyzed. An OR less than 1, equal to 1, and greater than 1 showed minor association, no association, and a more significant association between the expression of the PDGF/PDGFR system in the tissues of RA patients over nonarthritic subjects.

To estimate OR, a fixed-effect Mantel-Haenszel model was employed in the beginning. Where the I² statistic was greater than 50% or the p value was less than 0.10, it was interpreted as significant heterogeneity across studies; therefore, further analysis was done, using the random-effect model to estimate the pooled effect. Additionally, funnel plots were used to investigate potential biases that might stem from the publications.

3. Results

3.1. Publications Search. Figure 1 details an outline of the data search and the selection of studies in this research. An examination of the PubMed database showed 200 studies, of which 179 were excluded based on the inclusion criteria after scanning the titles and abstracts of 200 articles. A full-text perusal of the remaining 39 studies led to the selection of 6 studies for the meta-analysis. The remaining 15 studies were excluded due to insufficient data to allow data extraction for the analysis. For the 6 studies included in the meta-analysis, 1 study of patients in a Western country had 4 datasets on the PDGFA, PDGFB, PDGFC, and
PDGF [17]. Thus, the 6 studies had 9 datasets drawn from 6 countries, comprising 3 Western countries and 3 Asian countries. The 4 datasets of one of the 6 studies were treated separately by placing “pdgfA,” “pdgfB,” “pdgfC,” and “pdgfD” in parentheses against the surname of the first author of this study.

The association between susceptibility to RA and the PDGF/PDGFR system expression was meta-analyzed using the 6 studies with 9 datasets, as shown in Table 1 and Figure 2. Additionally, subgroup meta-analysis of susceptibility to RA and only PDGFR and only PDGF expression from 3 studies with 3 datasets and 3 studies with 6 datasets, respectively, were undertaken. The 6 studies with 9 datasets had 90 of 126 tests of RA and 27 of 97 tests of nonarthritic subjects had records of PDGFR/PDGFR system expression. For only PDGFR and only PDGF, 32 of 43 tests of RA and 6 of 21 tests of nonarthritic subjects were analyzed for only PDGFR. In contrast, 58 of 83 tests of RA and 21 of 76 tests for nonarthritic subjects had records of expression of only PDGF. Furthermore, comparing the susceptibility to RA due to PDGFR/PDGFR system expression among patients in Western countries and Asian countries was meta-analyzed using 3 studies with 6 datasets and 3 other studies with 3 datasets, respectively. For the expression of the PDGFR/PDGFR system in Asian and Western countries, 51 of 77 tests of RA and 21 of 65 nonarthritic subjects were recorded in Asian countries, while Western countries had 39 of 49 tests of RA patients and 6 of 32 tests of nonarthritic subjects.

3.2. The Association between PDGF/PDGFR Expression and Susceptibility to RA. An overview of the associations between PDGF/PDGFR system expression and RA is provided in Figures 2 and 3.

Six studies with 9 datasets were included in the meta-analysis to evaluate the association between PDGF/PDGFR system expression and RA subjects in Figure 2. The $I^2$ and $p$ value for heterogeneity were 43% and 0.08, respectively. Among the RA and nonarthritic subjects, the PDGF/PDGFR system expression had 5.25 highly significant greater odds of association in RA patients than the nonarthritic subjects. Thus, the meta-analysis showed a more significant association between PDGF/PDGFR system expression and RA patients in the overall population of the study (OR = 5.25, 95% CI = 3.00-9.18, $p < 0.0005$).
Figures 3(a) and 3(b) show the meta-analysis of associations of only PDGFR expression and only PDGF expression on the one hand and PDGF/PDGFR system expression in RA patients examined in Asian countries and Western countries on the other hand in Figures 3(c) and 3(d). To evaluate the association between only PDGFR expression and the RA patients, 3 studies with 3 datasets were included in the meta-analysis. The $I^2$ and $p$ value for heterogeneity were 63% and 0.07, respectively. Therefore, further analysis was performed using the random-effect model. As a result, the PDGFR expression in RA patients has 9.25 significant greater odds of association than that in nonarthritic subjects, as shown in the meta-analysis ($OR = 9.25$, 95% CI = 0.63-136.30, $p = 0.11$). In contrast, studies of PDGF expression in RA and nonarthritic subjects revealed the $I^2$ and the $p$ value for heterogeneity to be 42% and 0.13, respectively. Also, there was a significantly high PDGF expression with a 5.28 greater odds of association with RA patients compared with nonarthritic subjects, as shown in Figure 3(b) of the meta-analysis ($OR = 5.28$, 95% CI = 2.73-10.21, $p < 0.00001$).

For the assessment of the association between PDGF/PDGFR system expression and RA susceptibility of patients in Asian countries, 3 studies with 3 datasets were meta-analyzed. The $I^2$ and the $p$ value for heterogeneity were 0% and 0.65, respectively. The odds of association of the PDGF/PDGFR system expression in RA patients were 4.13, more significant than that in the nonarthritic group in Figure 3(c) of the meta-analysis ($OR = 4.13$, 95% CI = 2.04-8.39, $p < 0.00001$). In contrast, the association between PDGF/PDGFR system expression and RA subjects in

### Table 1: Characteristics of respective studies included in the meta-analysis.

| Author          | Year | Country   | Rheumatoid arthritis (RA) | Control/ nonarthritis (NA) | Sample | Detection | Gene         | PDGF/ PDGFR expression (RA) | PDGF/ PDGFR expression (NA) |
|-----------------|------|-----------|---------------------------|---------------------------|--------|-----------|--------------|----------------------------|-----------------------------|
| Rubin           | 1988 | Sweden    | 15                        | 5                         | 20     | STB       | IHC          | PDGFR                      | 15                          |
| Watanabe        | 2002 | Japan     | 3                         | 3                         | 6      | STB       | RT-PCR       | PDGFRA                     | 2                           |
| Walsh           | 2010 | UK        | 10                        | 11                        | 21     | CSMTMTP   | IHC          | PDGFB                      | 10                          |
| Charbonneau (pdgfA) | 2016 | Canada    | 6                         | 4                         | 10     | STB       | IHC          | PDGFA                      | 2                           |
| Charbonneau (pdgfB) | 2016 | Canada    | 6                         | 4                         | 10     | STB       | IHC          | PDGFB                      | 4                           |
| Charbonneau (pdgfC) | 2016 | Canada    | 6                         | 4                         | 10     | STB       | IHC          | PDGFC                      | 4                           |
| Charbonneau (pdgfD) | 2016 | Canada    | 6                         | 4                         | 10     | STB       | IHC          | PDGFD                      | 4                           |
| Matsumura       | 2019 | Japan     | 25                        | 13                        | 38     | STB       | IHC          | PDGFA/B                    | 15                          |
| Wang            | 2020 | China     | 49                        | 49                        | 98     | VWB       | ELISA        | PDGFB                      | 34                          |

VWB: venous whole blood; STB: synovial tissue biopsy; CSMTMTP: coronal section of middle third of medial tibial plateaux; ELISA: enzyme-linked immunosorbent assay; IHC: immunohistochemistry; RT-PCR: real-time polymerase chain reaction.

**Figure 2:** Odd ratios at 95% CI of individual studies and pooled data for the association between PDGF/PDGFR system expression and RA in all studies.
Western countries involved 3 studies with 6 datasets. Records of the $I^2$ and the $p$ value for heterogeneity were 61% and 0.02, respectively.

Further analysis was done using the random-effect model. As a result, the association between PDGF/PDGFR system expression in RA patients located in Western countries has 9.18 odds of significant association than that in the nonarthritic group as shown in Figure 3(d) of the meta-analysis (OR = 9.18, 95% CI = 2.04-8.39, $p = 0.03$). Thus, the analysis has shown that the odds of association of the PDGF/PDGFR system expression in RA patients are higher among Western countries than in Asian countries.

3.3. Publication Bias. The analyses of PDGF/PDGFR system expression in RA patients and the nonarthritic group demonstrated that all the studies included in the meta-analysis...
develop a development and progression of RA disease are needed to
Invariably, the meta-analyses of cytokines implicated in the
targeted treatments for RA disease in the future [6, 22
Several studies evaluating the role of cytokine protein array
analysis for the 6 studies with 9 datasets was included in the
meta-analysis, as shown in Figure 4. The symmetrical shape
of the funnel plot does not indicate any publication bias
though most of the sample population of eligible studies
for the meta-analysis was small.

4. Discussion
Several studies evaluating the role of cytokine protein array
profile associated with RA have been done with varied out-
comes of these cytokines, including the PDGF/PDGFR
system that calls for developing and integrating cytokine-
targeted treatments for RA disease in the future [6, 22–24].
Invariably, the meta-analyses of cytokines implicated in the
development and progression of RA disease are needed to
develop a fit-for-purpose-targeted treatment for RA disease.
The PDGF/PDGFR system has shown its immense role
in the development and progression of RA disease as follows:
several pieces of research on the PDGF/PDGFR system in
issues of RA patients and in vitro and in vivo studies have
demonstrated the system’s significant role in the occurrence,
development, and progression of RA. The PDGF is known
to play a crucial role in proliferation and migration, cause
an increase in mRNA and protein expressions of proinflam-
atory cytokines such as interleukin-1b and interleukin-8
by synergizing with the tumor necrosis factor in the RA-
fibroblast-like synoviocytes (FLS), increased the amounts of
FLS in the G2/M cell cycle phase, promote morphological
changes of the FLS cells to a dendritic shape, and cause a sig-
ificant increase in cathepsin B secretion that can degrade
some collagen types and proteoglycans in acidic physiological
conditions [14, 16, 20, 21, 25–32]. Furthermore, the mecha-
nism of PDGF’s role in proliferation and migration results
from synergistic action between PDGF and interleukin-1 in
indomethacin [26]. There are also reports of association of
the PDGFR with chronic synovial inflammation and involve-
ment in the development of the destructive phenotype of
synovial cells in RA [17, 19].
These reports call for a meta-analysis to ascertain the
level of significance of the PDGF/PDGFR system expression
with RA susceptibility. The data from published pieces of
research were combined to evaluate the overall association
of PDGF/PDGFR system expression in RA patients and sub-
group analysis of the system’s association with RA based on
the in-country location of the patients, and the association of
only the PDGFs and only PDGFR with RA was evaluated.
We discovered significant associations of the PDGF/PDGFR
system expression with RA patients across the entire popula-
tion of all the study subjects and with RA patients in Asian
countries and Western countries and marked the association
between only PDGF expression with RA patients than with
control subjects. Thus, only PDGFR expression in the study
subject was insignificantly associated with RA susceptibility.
It is worth noting that although there was more tests of the
PDGF/PDGFR system expression in the study subjects of
Asian countries with 142 tests than in the study subjects of
Western countries with 81 tests, the association of PDGF/
PDGFR system expression was more associated with RA
patients in Western countries than in Asian countries.
In addition, the testing for only PDGF and only PDGFR
recorded 64 tests and 159 tests, respectively. Hence, the vast
difference in the numbers might have accounted for the
insignificance of the association between only PDGFR with
RA susceptibility. However, overall, the population of the
study subjects was small. Therefore, we recommend that
more large sample sizes for the study of PDGF/PDGFR sys-

tem expression in RA patients should be done in the future,
with attention to details on the country of origin of RA
patients to allow concrete evaluation of ethnic-origin RA
patients PDGF/PDGFR system expression association with
RA susceptibility.
This meta-analysis is the first of its kind to ascertain the
PDGF/PDGFR system expression with RA susceptibility.
Considering that the PDGF/PDGFR system expression in
human adults is associated with diseases and also has a sig-
nificant association with RA, then targeting this PDGF/
PDGFR has great potential in the design of combination
therapies for RA [10].

5. Conclusion
The overall results indicated a significant association
between the PDGF/PDGFR system expression and RA, with
a higher association among RA patients in Western coun-
tries than in Asian countries. Additionally, only PDGF
expression in RA patients showed a significant association,
while only PDGFR expression in RA patients was not signif-
ically associated with RA susceptibility. In conclusion, the
findings in this meta-analysis suggest that the PDGF/
PDGFR system expression is significantly associated with
RA susceptibility, especially the PDGFs, and more signifi-
cantly associated with RA patients in Western countries.
The limitation of this study is few cohorts. However, these

Figure 4: Funnel plot for assessing publication bias in the meta-
analysis of the 6 studies for PDGF/PDGFR system expression and
susceptibility to the RA disease.
findings present evidence that calls for large cohort studies in PDGF/PDGFR system expression in RA to pave the way for a comprehensive updated investigation in the future.

Data Availability

No data were used to support this study.

Conflicts of Interest

The authors declare no conflict of interest in this research.

Authors’ Contributions

LF and WS did conceptualization in consultation with ZX and XC. The search of relevant publications was done by LF and WS independently and rechecked by WW. WJ, JC, LF, and WS worked on the methodology and data extraction, and LF, WS, and WW meta-analyzed the data in consultation with XH. Finally, ZX and XC reviewed and edited the first draft of the manuscript written by LF and WS. Feiyuan Li and Shaoping Wu have contributed equally to this work and share first authorship. Chao Xie and Xia Zhang are the corresponding authors.

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