High Tissue Factor Microparticle Level in Major Thalassemic Patients with Pulmonary Hypertension

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Abstract

Objective: To analyze the correlation between tissue factor microparticles (TF-MP) levels and pulmonary hypertension (PH) in adult thalassemic patients.

Methods: This study was conducted from September to October 2018, using secondary and primary data. The secondary data consisted of the PH parameter, which was retrieved from a 2017 previous study entitled 'Clinical Characteristic and Complication due to Iron Overload in Thalassaemic Patients' in 2017 while the primary data were the TF-MP which were obtained from the analysis of frozen serum of the same population using ELISA method. The mean pulmonary arterial pressure (mPAP) values were obtained from echocardiography results and PH was defined as mPAP >25 mmHg.

Results: Seven (16.7%) major thalassemic patients experienced PH. The median values of TF-MP levels were higher among major thalassemic patients with PH when compared to the non-PH patients (1569 vs 11.5 pg/dL; p=0.023). No significant difference was observed in the median TF-MP levels between subjects with splenectomy and subjects without splenectomy (11.6 vs 12.3 pg/dL; p=0.44). There was also no difference in mPAP values between subjects with splenectomy and subjects without splenectomy (18.0 vs 17.0 mmHg; p=0.663). When the median TF-MP levels among major thalassemic patients were analyzed in terms of correlation with transfusion level, no statistically significant difference was seen between subjects who received sufficient transfusions (≥180 mL/kg/year) and those who received insufficient transfusions (<180 mL/kg/year) (r=0.138; p=0.390).

Conclusions: There is a positive correlation between the TF-MP levels and PH in adult major thalassemic subjects.

Keywords: Major thalassemia, pulmonary hypertension, tissue factor microparticles

Introduction

Homozygous β-globin carrier defects in thalassemia patients can give rise to several complications due to hemolysis, chronic anemia, and continuous transfusions. In the last few decades, major thalassemia patients can live longer with the availability of blood transfusion and iron chelation. With the longer survival, the previously unrecognized complications associated with hypercoagulability become visible.1,2 Complications associated with hypercoagulability appear to be subclinical, marked only by changes in the hemostasis parameters, until thromboembolic processes such as stroke, transient ischemic attack, deep vein thrombosis (DVT), and pulmonary embolism occur. Pulmonary hypertension (PH) in thalassemia is included in the 5th class
of the PH classification of the 2013 American College of Cardiology and is considered to happen via multifactorial mechanisms. It is considered as the complication that becomes the main factor of morbidity and mortality in these patients. Pulmonary hypertension involves several mechanisms such as endothelial dysfunction, inflammation, and chronic thromboembolism. The mechanism of thromboembolism in thalassemia patients with PH has not been clearly understood.\textsuperscript{3, 4}

The multifactorial mechanism requires microparticles to play distinctive roles in the cellular interactions related to PH pathogenesis. Microparticles are vesicles from the plasma membrane, ranging from 0.1 to 1 \(\mu\)m in size, that are released by cells through activation by their agonists, physical and chemical stresses, and apoptosis. Tissue factor microparticles (TF-MP) or circulating Tissue factor (cTF) is a protein released by cellular activation or apoptosis that plays an important role in producing thrombin in the propagation phase of the cell-based coagulation. Cellular activation in thalassemia is caused by negatively-charged cell membranes and hemolysis process disorders. Previous studies have reported that increased TF-MP levels are found in thalassemia patients, as well as in patients with 1st and 3rd group PH.\textsuperscript{5, 6}

No data available yet on the roles of TF-MP in thalassemia patients with PH. This study aimed to analyze the correlation between TF-MP level and PH in major thalassemia patients in Dr. Hasan Sadikin General Hospital, Bandung, West Java, Indonesia.

**Methods**

This was a cross-sectional quantitative study using numeric-numeric correlative statistical analysis conducted from September to October 2018. Subjects in the study were thalassemia patients treated at the Hematology-Medical Oncology Clinic, Internal Medicine Department, Faculty of Medicine, Universitas Padjadjaran/ Dr. Hasan Sadikin General Hospital Bandung. PH parameters were collected as secondary data from a previous study entitled ‘Clinical Characteristic and Complication due to Iron Overload in Thalassaemic Patients’ which was conducted in 2017 while data on TF-MP were collected by testing the frozen serum of the same population of this 2017 study. The inclusion criteria were thalassemia patients aged ≥14 years who had received treatments and regular transfusions for more than two years after diagnosis, had echocardiographic data, and had their frozen serum stored. The exclusion criteria were intermediate thalassemia patients and patients with left ventricular or left atrial hypertrophy. Pulmonary hypertension (PH) was measured using the GE Vivid S6 echocardiography device. Meanwhile, the severity of PH was determined by estimating the mean Pulmonary Arterial Pressure (mPAP) (≥25 mmHg). The mPAP was measured while the patient was resting through the use of the Pulmonary Velocity Accelerated Time (PVaccT) calculation [mPAP = 80 - (PVaccT/2)]. Data on the MP-TF levels were collected through subject serum analysis using the ELISA method and the results were reported in pg/dL.

Before statistical analyses were performed, data normality testing was performed on numerical data, such as age, period of suffering from thalassemia, transfusion period, total transfusions volume, sufficiency of transfusions, ferritin levels, Hemoglobin (Hb) levels, thrombocyte counts, mPAP values, and TF-MP levels. The Shapiro-Wilk test was selected for this testing because the number of subjects was less than 50. Data on age, ferritin levels, hematocrit percentages, thrombocyte counts, ferritin levels, period of suffering from thalassemia, sufficiency of transfusions, mPAP values, and MP-TF levels were not normally distributed (p>0.05), while data on Hb level were normally distributed (p>0.05).

Normally distributed data were presented in mean and analyzed using Pearson correlation (r) test, while data that were not normally distributed were presented in median and analyzed using non-parametric Spearman correlation test (rs). Point biserial correlation (rpb) analysis was performed to observe the correlation between numerical and categorical data. Results were declared as significant if p<0.05.

**Results**

Subject characteristics, including age, sex, ferritin levels, Hb levels, Ht percentages, thrombocyte counts, sufficiency of transfusions, numbers of subjects with PH or without PH, mPAP values, and TF-MP levels, are presented in Table 1. The median age was 20 years old. More females were included in the study than males (38.1% versus 61.9%). The median period of suffering from thalassemia was 198 months. The sufficiency of transfusion was 65.6 cc/kgBW/year while the mean Hb level, median thrombocyte count, and median ferritin level were 7.0±1.3 g/dL,
The subjects characteristics including the age, sex, ferritin levels, Hb levels, Ht percentages, thrombocyte counts, sufficiency of transfusions, numbers of subjects with PH or without PH, mPAP values, and TF-MP levels can be seen in Table 1 below. The median of age was 20, consisting of males (38.1%) and females (61.9%). The median of period of suffering from thalassemia was 198 months. The sufficiency of transfusions was 65.6 cc/kgbb/year, the mean of Hb level was 7.0±1.3 g/dL, the median of thrombocyte count was 137,000, and the median of ferritin level was 3940 ng/dL.

When the correlation between MP-TF level and mPAP value was assessed using the Spearman correlation test with a 95% confidence interval, it was revealed that the MP-TF level had a statistically significant correlation with the mPAP value (r=0.481; p=0.001).

**Discussion**

This study presented a correlation between...
Fig 1 Scatter Diagram of Correlation between MP-TF Level and mPAP Value

Fig. 2 Boxplot of Median MP-TF Level in PH and Non-PH Groups

Fig. 3 Scatter Diagram of Correlation between Transfusion Sufficiency and TF-MP Level
increased MP-TF levels and high mPAP values (moderate correlation; r=0.481; p<0.001). Based on the point biserial analysis to observe a direct correlation between the MP-TF level and the presence of PH, a significant difference was observed between PH and non-PH subjects (p<0.05; rpb=0.311).

A study conducted by Bakoubula et al. in 2008 that examined the correlation between MP-TF levels and PH by enrolling 20 subjects with 1st and 3rd group of PH discovered a statistically significant correlation between the MP-TF level and PH (p<0.001). Dhiel et al. in 2010 who studied 19 subjects with PH caused by chronic thromboembolism, idiopathic PH, rheumatic-related PH, and chronic pulmonary disease (CPD)-related PH also found a positive correlation between procoagulant MP level and occurrence of PH in study subjects when compared to the normal control.7, 8

Hypercoagulability complications in thalassemia patients are characterized by

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**Fig. 4 Boxplot of TF-MP Level in Splenectomy and Non-splenectomy Groups**

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**Fig. 5 Boxplot of mPAP Levels in Splenectomy and Non-splenectomy Groups**
subclinical defects marked by hemostasis parameter changes until the occurrence of thromboembolic events such as stroke, transient ischemic attack, DVT, and chronic pulmonary embolism. The PH in thalassemia can be included in the 5th group of the 2013 ACC classification. Pulmonary hypertension in thalassemia is caused by various factors and mechanisms, including pulmonary arteriole in situ thrombosis. Interactions among factors that play the roles in the occurrence of PH in major thalassemia patients require a medium such as procoagulant MP.5,9

This study reveals a correlation between MP-TF level and mPAP value in major thalassemia population. This study supports findings of previous studies and adds supporting data of in situ thrombosis process in the pathogenesis of the 5th group PH through the roles of procoagulant microparticles, such as TF-MP.5

The correlation between the TF-MP level and PH discovered in this study is considered moderate (r=0.481). This may relate to other mechanism pathways, such as the endothelial dysfunction pathway, triggered by decreased nitric oxide and inflammation. Endothelial dysfunction causes an imbalance in mediator production, which leads to vasoconstriction, smooth muscle hyperplasia, and pulmonary arterial remodeling. Other factors that can influence the occurrence of PH are not included in this study despite the possibility that they may also influence the significance of the correlation. Hence, this is considered a limitation in this study.5,10

Of all thalassemia patients in this study, 16.7% had PH. This is different from a previous study by Fayed et al. in 2017 which reported that 36% of major thalassemia children have PH. The presence of PH in their study was determined through echocardiographic examination by calculating mPAP using a different formula.11

Based on the 2015 ESC, mPAP is not used for echocardiographic diagnosis of PH. Instead, the 2015 ESC uses the tricuspid regurgitant velocity (TRV) and other imaging techniques for assessing complications of PH in the right side of the heart. This method is more specific and widely used. The results are categorized as low, moderate, or high probability for PH. This study uses secondary data from a previous study. However, the reference study did not use TRV calculation in the determination of PH, which creates another limitation of this study.12

The median TP-MF level of the subjects of this study was 12.1 pg/dL, which is much higher than the TP-MF level in normal people, which is less than 0.21 pg/dL. The increase in the TF-MP level in major thalassemia patients in comparison to the normal population is also demonstrated in some previous studies. A higher level of TF-MP can also be found in several other diseases such as HIV, diabetes mellitus, and sickle cell disease.13,14

The determinant factors of defect in thalassemia are chronic hemolysis and ineffective erythropoiesis with iron overload which leads to complications in thalassemia, including hypercoagulability. Hemichrome precipitation due to thalassemic erythrocyte globin chain defect can cause heme disintegration, enhance intracellular labile iron, and produce reactive oxygen species, which lead to endothelial activation and dysfunction.10

The majority of the subjects in this study (95.2%) had a pre-transfusion Hb level of <9 g/dL, which was caused by insufficient transfusions. The median of transfusion sufficiency in this study was 65.6 cc/kg BW/year, which is below the standard of ≥180 cc/kg BW/year. No significant correlation was observed between the insufficiency of transfusions and either TF-MP level and the presence of PH. A previous study by Hastuti in Dr. Hasan Sadikin General Hospital, Bandung, Indonesia in 2007 on major thalassemia patients discovered a correlation between the sufficiency of transfusions and the hypercoagulability index when examined by TEG. The insufficiency of transfusions can cause a higher level of thalassemic erythrocyte cells.15

The results of this study were different from the results reported by Agouti et al. in 2015 on 37 major thalassemia patients. Their study described higher TF-MP in subjects who received insufficient transfusions, which was assessed using the flow cytometry test. Flow cytometry test is considered to be more sensitive and has the ability to identify the cell origin of TF-MP. Literature stated that complications due to chronic hemolysis and ineffective hematopoiesis in relation to the thromboembolic processes are more frequently seen in intermediate thalassemia than in major thalassemia. Agouti’s study does not include intermediate thalassemia patients so complications caused by insufficiency of transfusions do not influence the higher TF-MP level and PH.16

Splenectomy in thalassemia patients triggers an increase in thrombocyte count, activation, and aggregation. These conditions may lead to a higher level of TF-MP derived
from thrombocytes. Splenectomy is also associated with an increase in mPAP values in thalassemia patients and previous studies have discovered a higher PH incidence in patients with splenectomy when compared to those without splenectomy.11

The result of this study showed no significant correlation between splenectomy and increased TF-MP level and mPAP value. A previous study stated that the incidence of PH is higher in thalassemia patients who had undergone splenectomy. The difference in result is probably caused by the number of the subjects in the study which is lower (36 subjects) than in this study, along with the higher frequency (47.2%) of PH in their study. The mean thrombocyte count in their study is 548.200±242.500/mm$^3$, which is higher than in this study (137.000/mm$^3$).11

Several limitations are noted in this current study, including PH examination using PvaccT calculation, which is less specific than other methods such as the TRV from echocardiography. This study does not analyze the origin of cells that activate and release TF-MP, thus unable to describe the influence of sufficiency of transfusions on the TF-MP level. This study also does not analyze major factors that can influence the pathogenesis of PH from the endothelial dysfunction and inflammation pathways; hence, major determinants in the significance of the correlation between TF-MP level and PH in major thalassemia patients cannot be discovered.17,18

In summary, this study proves that the TF-MP level influences in situ thrombosis in the mechanism of PH in major thalassemia patients. Further studies with cohort design and multiple regression analysis of other determinant factors in the endothelial dysfunction and inflammation pathways that might contribute to the mechanism of PH should be conducted. In major thalassemic patients with sufficient transfusions, it is not crucial to discover the origin of cells that release TF-MP using flow cytometry test.

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