Interaction of Panax quinquefolius Saponin and Dual Antiplatelets on Vascular Endothelial Function in Rats with Acute Myocardial Infarction

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1. Introduction

Cardiovascular diseases remain the predominant cause of morbidity and mortality all over the world. Immediately percutaneous coronary intervention (PCI) is one of the most frequently therapies used in clinical practice for acute myocardial infarction (AMI), which remarkably increased the successful rate of revascularization and reduced the mortality and morbidity of AMI patients [1, 2]. Despite the fast development of interventional cardiology, drug therapy has always been a cornerstone in the treatment of cardiovascular diseases. Based on the integrative medicine of Eastern and Western worlds, the application of herbal medicine (HM) has valuable significance in reducing the risk of cardiovascular event [3]. In Asia, HM is used for prevention and treatment of coronary heart disease (CHD) in many countries for thousands of years. Many studies showed that the combination therapy of HM and conventional western medicine significantly improved life quality and prognosis of patients after PCI [4, 5]. Clinical application of dual antiplatelet drugs such as aspirin and clopidogrel is important in the prevention and therapy of CHD, but prolonged treatment with dual or triple antiplatelet drugs revealed a diversity of platelet reactions, which limits the clinical application widely.

Panax quinquefolius saponin (PQS) is an active part extracted from the stems and leaves of Panax quinquefolius (American ginseng), which has a beneficial effect on the treatment of CHD with increasing energy storage in ischemic myocardium, promoting angiogenesis, inhibiting oxidative stress injury and ventricular remodeling, and so forth.
just as indicated in our previous studies [6–8]. However, whether PQS has some additional beneficial effects with dual antiplatelets (aspirin and clopidogrel) in thrombosis, antiplatelet activity and improving endothelial function for CHD patients remain unclear. This study was designed to examine the interaction of PQS and dual antiplatelets (aspirin and clopidogrel) on antiplatelet activity and vascular endothelial function in rats with AMI.

2. Methods

2.1. Animals Grouping and Treatment. Male Sprague-Dawley rats (weight 200–220 g) were purchased from Beijing University Laboratory Animal Center (the animal certificate number: SCXK (Jing) 2011-0004). Rats were housed in humidity-controlled (60 ± 10)% rooms at (24 ± 1)°C with a 12 h on/12 h off light cycle. The animals were maintained with free access to standard diet and tap water.

After one week of adaptive feeding, AMI model was created in rats by ligating the left anterior descending coronary artery (LAD) as described before [9]. Twelve rats without AMI were assigned to sham group, and rats with successful operation of coronary artery ligation were divided randomly into 3 groups as follows, model group; dual antiplatelets group (9 mg/kg/d of aspirin and 6.75 mg/kg/d of clopidogrel); PQS + dual antiplatelets group (Xinyue capsules (162 mg/kg/d) combined with dual antiplatelets), with 12 rats in each group. All drugs were diluted with distilled water, and the dosages were evaluated with body surface coefficient conversion between human and rat. Rat weights were measured every week. Rats in sham and model groups were administered with the same volume of distilled water. All rats were intragastrically administrated for 28 days. The Animal Care and Use Committee of Xi’An Jiao Tong University of Science and Technology approved the experimental protocol.

2.2. Drugs. Xinyue capsules (50 mg PQS/capsule) were purchased from Yisheng Pharmaceutical Co., Ltd., Jilin Province of China (SFDA Approval number Z20030073); clopidogrel hydrogen sulfate tablets (75 mg/pill) were purchased from Sanofi Winthrop Industry, France (SFDA Approval number J20080090); aspirin enteric-coated tablets (50 mg/pill) were purchased from Peking Shuguang Pharmaceutical Co., LLC (SFDA Approval number H11020827); and penicillin sodium injection (80 IU/bottle) was purchased from North China Pharmaceutical Co., Ltd. (SFDA Approval number XI105313). All the drugs were dissolved before use.

2.3. Thrombosis Time. Six rats were chosen randomly in every group to test thrombosis time. After the last intragastric administration, rats were anesthetized with intramuscular 4% chloral hydrate (0.9 mL/100 g body weight) solution and disinfected for operation. The skin was sheared at the right side of cervical part. The length of right carotid artery exposed was about 2 cm with padded cellophane under the vessel to protect the surrounding tissues. Stimulating electrode and temperature reporter were placed under the vessel separated to stimulate vascular by current of 80 µA for 7 min, resulting in thrombosis. The sudden temperature decreases at the distal vascular detected by temperature reporter meant thrombosis. The thrombosis time, also known as occlusion time, was measured from the beginning of simulation to thrombosis.

2.4. Thrombus. As described before [10], the length, wet weight, and dry weight of thrombus were tested in the other 6 rats in vitro. Blood sample was drawn out from abdominal aortic and then injected into silicone tube of 2 mm diameter and 25 cm length quickly to reach about 1/2 length of the tube. The silicone tubes were put into Thrombosis and Platelet Adhesion Instrument (XSN-RII, Wuxi 2nd Electrics and Electronics Factory, Jiangsu province) and uniformly rotated for 15 min. Thrombus was picked out of the tubes to measure the length and wet weight after absorbing liquid gently with filter paper. The dry weight of thrombus was tested after heating in dryer at 37°C for 20 min.

2.5. Detection of Platelet Aggregation. About 2 mL of blood sample was drawn out from abdominal aortic and mixed with 3.8% sodium citrate (v/v = 9:1). Platelet-rich plasma was obtained by centrifuging at 800 r/min for 10 min, while platelet-poor plasma was obtained by centrifuging at 3000 r/min for 10 min. Concentration of platelet was adjusted in PRP to 600–700 × 10^9/L. According to Born’s method [11], platelet aggregation was induced by adenosine diphosphate (ADP), and the maximum platelet aggregation rate was measured by Automatic Four-channel Platelet Aggregation Instrument (LBY-NJ2, Shanghai Precil Instrument, Inc.).

2.6. Concentration of Thromboxane B_2 (TXB_2) and 6 Ketone Prostaglandin F_1α (6-Keto-PGF_1α). The concentrations of TXB_2 and 6-keto-PGF_1α in plasma were detected with radioimmunoassay (r-911, Industrial Corporation of China University of Science and Technology) according to the procedures of kits specification provided by Beijing Huaying Biotechnology Research Institute.

2.7. Concentration of Endothelin-1 (ET-1) and Nitric Oxide (NO). ET-1 concentration in plasma was measured by radioimmunoassay according to the procedures of kits specification (British Enzo Co.). NO level in serum was detected with nitrate reductase method using automatic microplate reader (MK3, Finland Labsystems Co.) referring to as the kit instruction (British Enzo Co.).

2.8. Ratio of Ventricular Cavity Area and Cardiac Transverse Area. After blood sample was drawn out from abdominal aortic, heart was removed quickly. Cardiac cavity was irrigated by saline and liquid on surface was absorbed. Pathological section was cut into 5 pieces (5 mm in each piece) down from the ligation and parallel with coronary ditch. Sections were stained with haematoxylin-eosin (HE), and ventricular cavity area and cardiac transverse area were measured and the ratios between them were calculated by color pathological image analysis system (DpxView Pro, Denmark DeltaPix Co.). Pathological changes of rat myocardial tissues were observed by optical microscopy (BN-2, Japan Olympus Co.).
2.9. Coagulation Related Markers. Semiautomatic blood coagulation instrument (produced by Teco Medical Instruments Production, Germany; supplied by Beijing Chuangxin technology Co., LTD.) was used for measuring APTT (kit number A04-080929-N05), PT (kit number V28417), TT (kit number 011-I10397), and FIB (kit number S210028) following the introduction of each kit.

2.10. Statistical Analysis. Statistical analysis was performed using SPSS 17.0 software. The mean ± standard deviation (SD) was determined for each group. Statistical analysis was performed with ANOVA F test and the nonparametric Kruskal-Wallis test. Differences were considered statistically significant when P < 0.05, and the exact P values were shown unless P < 0.001.

3. Results

3.1. The Ratio of Ventricular Cavity Area and Cardiac Transverse Area. Rats in model group showed thinned ventricular wall in infarction area, thickened ventricular wall in non-infarctional area, ventricular cavity dilatation, deformation and ventricular cavity area and heart transverse section ratio significantly increased compared with sham group (P < 0.001). Ventricular wall changes had been improved to different degree in all intervention groups, and the ratio of the ventricular cavity area and cardiac transverse area was decreased significantly in PQS + dual antiplatelets group (P = 0.001) compared with model group. The ventricular cavity area and cardiac transverse area ratio in PQS + dual antiplatelets group showed a decrease tendency, but there was no statistical difference (P = 0.058) compared with dual antiplatelets group (see Figures 1 and 2).

3.2. Concentrations of Plasma ET-1 and Serum NO. Compared with sham group, plasma ET-1 concentration increased remarkably (P = 0.009), whereas serum NO concentration decreased significantly in model group (P = 0.009). Plasma ET-1 concentration showed a significant decrease in PQS + dual antiplatelets group (P < 0.001), while serum NO concentration showed a significant increase in PQS + dual antiplatelets group (P = 0.001) compared with model group. The ventricular cavity area and cardiac transverse area ratio in PQS + dual antiplatelets group showed a decrease tendency, but there was no statistical difference (P = 0.058) compared with dual antiplatelets group (see Figures 1 and 2).

3.3. Platelet Aggregation Rate (PAgT(%)). PAgT(%) decreased significantly in both dual antiplatelets group and PQS + dual antiplatelets group (P = 0.039, P = 0.013) compared with model group. PQS + dual antiplatelets group showed no statistical difference in PAgT(%) (P = 0.515 > 0.05) compared with dual antiplatelets group (see Figure 3).

3.4. Thrombosis. Thrombosis time in vivo shortened significantly (P = 0.031), and thrombus length, wet and dry weight in vitro increased significantly in model group (P = 0.017, P = 0.031, and P = 0.038) compared with sham group. Compared with model group, thrombosis time in vivo prolonged significantly and thrombus length, wet and dry weight in vitro decreased significantly in dual antiplatelets group (P = 0.013, P = 0.010, P = 0.002, and P = 0.003) and PQS + dual antiplatelets group (P = 0.003, P = 0.005, P = 0.001, and P = 0.002). Rats in PQS + dual antiplatelets group showed a decreased tendency in thrombosis time in vivo (P = 0.405), thrombus length (P = 0.817), and wet and dry weight (P = 0.725 and P = 0.861, resp.) in vitro as well, but no significant difference was observed compared with dual antiplatelets group (see Figures 5 and 6).

3.5. Concentration of Plasma TXB<sub>2</sub> and 6-Keto-PGF<sub>1α</sub>. A significant decrease of 6-keto-PGF<sub>1α</sub> concentration has been shown in model group (P < 0.001) compared with sham group. TXB<sub>2</sub> concentration significantly decreased in dual antiplatelets and PQS + dual antiplatelets groups (P < 0.001), whereas 6-keto-PGF<sub>1α</sub> concentration significantly increased in PQS + dual antiplatelets group (P = 0.022) compared with model group. A decreased tendency of TXB<sub>2</sub> concentration (P = 0.274) and an increased tendency of 6-keto-PGF<sub>1α</sub> concentration showed (P = 0.539), respectively, in PQS + dual antiplatelets group compared with dual antiplatelets group (see Figure 7).

3.6. Coagulation Related Markers. Activated partial thromboplastin time (APTT) shortened significantly but FIB increases significantly in model group compared with sham group (P = 0.002, P = 0.005). Compared with model group, APTT and PT prolonged significantly in dual antiplatelets group (P < 0.001, P = 0.006) and PQS + dual antiplatelets group (P < 0.001, P = 0.004), whereas higher level of FIB has been observed in two groups (P < 0.001, P < 0.001). However, there was no significant difference between antiplatelets group and PQS + dual antiplatelets group (see Figure 8).

4. Discussion

The Xinyue capsule, mainly composed of PQS, could significantly improve clinical symptoms of CHD patients, improving left ventricular ejection fraction (LVEF) and life quality in combination with conventional western medicine treatment [12]. Previous experiments have showed that PQS increased the levels of vascular endothelial cell growth factor (VEGF) and basic fibroblast growth factor (bFGF), thus promoting angiogenesis in ischemic myocardial area [13] and improving adenosine triphosphate (ATP) content and energy storage in ischemic cardiomyocytes as well [14]. Ischemia reperfusion injury and ventricular remodeling also have been found attenuated in rats that suffer from acute myocardial infarction [15, 16]. And this experiment showed the further beneficial effects of PQS on endothelial function, thrombosis, and platelet activity in combination with dual-platelets in rats with acute myocardial infarction.

PQS + dual antiplatelets showed a tendency in decreasing the ratio between ventricular cavity area and cardiac transverse area, a further significant decrease in plasma ET-1 concentration and a further significant increase in serum NO concentration compared with dual antiplatelets alone.
Figure 1: Cardiac transverse section in different groups (HE staining, general pathologic). (a) Sham group; (b) model group; (c) dual antiplatelets group; (d) PQS + dual antiplatelets group.

Figure 2: The ratio of ventricular cavity area and cardiac transverse area in different groups. The ratio was significantly lower in the PQS + dual antiplatelets group than in the model group. Data are mean ± SD (n = 12 rats/group). *P < 0.05 and **P < 0.01 versus sham group, ##P < 0.01 versus model group.

PQS and dual antiplatelets showed a decreased tendency in PAgT(%), thrombosis time in vivo, thrombus length, wet and dry weight in vitro, TXB_2 concentration, and an increased tendency in 6-keto-PGF_1alpha concentration compared with dual antiplatelets. In addition, there were no significant differences between PQS and dual antiplatelets and dual antiplatelets in coagulation related markers (APTT, PT, TT, and FIB).

ET had great effects on promoting vascular smooth muscle cell proliferation and vasoconstriction, whereas NO could dilate the vessel and inhibit platelet aggregation. There exists a dynamic balance between ET and NO in physiological condition. A large amount of ET release when vascular endothelium injured, which could trigger strong contraction of local vessels and facilitate platelet aggregation through binding to protein kinase isozymes and inhibiting NO release [17, 18]. In this experiment, there was a significant decrease in plasma ET-1 concentration and an increase in serum NO concentration as compared with dual antiplatelets, indicating that PQS might have beneficial effects on rats with AMI, in which process protection of vascular endothelial function and inhibition of platelet aggregation are involved. These effects result from regulation of dynamic balance between ET and NO by PQS which were similar to our previous studies [19].

Very few focuses were fixed on measuring ventricular cavity area and cardiac transverse area ratio, PAgT(%), thrombosis time in vivo, thrombus length, and wet and dry weight in vitro in previous studies. In this experiment, there was no significant difference between PQS + dual antiplatelets group and dual antiplatelets group alone. However, the...
platelet activity, antithrombotic activity, and plasma clotting time are significantly influenced by PQS when distilling PQS in plasma directly [20]. This might be because the rats were not be given enough PQS, or injection not for oral agent should be used.

TXA$_2$ greatly accelerates platelet aggregation and vasoconstriction, while PGI$_2$ could prolong platelet aggregation and promote vasodilation. Because of the short half-life period and unstable characteristics of them, the concentrations of their metabolic products, TXB$_2$ and 6-keto-PGF$_{1α}$, were measured indirectly. Thromboxane synthetase and prostaglandin synthetase existed in platelet and vascular endothelial cell, respectively. Once vascular endothelial cells injured, platelet aggregation easily occurs due to the local PGI$_2$ decrease. In this experiment, TXB$_2$ concentration showed a significant decrease and 6-keto-PGF$_{1α}$ concentration showed a significant increase in PQS + dual antiplatelets group compared with model group, which indicated that PQS + dual antiplatelets might prolong platelet aggregation through regulating the production or dynamic balance of TXB$_2$ and 6-keto-PGF$_{1α}$, but rats in PQS + dual antiplatelets group showed no significant difference compared with dual antiplatelets group on TXB$_2$ and 6-keto-PGF$_{1α}$. However, previous study [21] had showed PQS 25 mg/kg, 50 mg/kg significantly decrease the 6-keto-PGF$_{1α}$ concentration and significantly increase the TXB$_2$ concentration, and the injection of PQS might make the differences.

In conclusion, combining with dual antiplatelets, PQS might improve vascular endothelial function and protect ventricular remodeling in rats with AMI. Although PQS + dual antiplatelets treatments have shown tendency of better antithrombotic effect compared to dual antiplatelets alone, there is no significant difference. Therefore, the antithrombotic effect of combination of PQS and dual antiplatelets treatment will be further investigated, while other forms of PQS such as injection will be used in our future study.
5. Conclusion

We have provided experimental evidence supporting our conclusion that, combining with dual antiplatelets, PQS might improve vascular endothelial function and protect ventricular remodeling in rats with AMI.

Abbreviations

PQS: *Panax quinquefolius* saponin
CHD: Coronary heart disease
AMI: Acute myocardial infarction
LAD: Left anterior descending coronary artery
NO: Nitric oxide
TXB$_2$: Thromboxane B$_2$
APTT: Activated partial thromboplastin time
PT: Prothrombin time
TT: Thrombin time
FIB: Fibrinogen
PCI: Percutaneous coronary intervention
CHM: Chinese herbal medicine
WM: Western medicine
ET-1: Endothelin-1
ADP: Adenosine diphosphate
PAGT(%): Platelet aggregation rate
LVEF: Left ventricular ejection fraction
SD: Standard deviation.

Conflict of Interests

The authors declare that there is no conflict of interests.

Authors’ Contribution

Baojun Wang and Yue Liu are the co-first authors. Dazhuo Shi and Jiangang Liu conceived and designed the animal experiments and helped to draft the paper. Baojun Wang, Qingxiang Zhang, and Lei Zhang performed the experiments. Baojun Wang and Yue Liu analyzed the data and prepared the paper with Qinghua Shang.

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