Abstract. The effects of different doses of folic acid and vitamin B12 on rabbits with deep vein thrombosis (DVT) and hyperhomocysteinemia were investigated. In total, 60 New Zealand rabbits were divided into untreated control, low-dose and high-dose groups. After inducing DVT, hemorheology and coagulation indexes were measured 3 and 10 days later. We found that both treatment groups performed better than the control group, and the high-dose performed better than the low-dose. Ten days after thrombosis, the levels of Hcy and D-dimer were lower in the high-dose group. Moreover, the changes of lower extremity deep venous thrombosis were significantly reduced in both high- and low-dose groups, but the high-dose group showed the most improvement. The effective rate of the high-dose group was 100%, higher than the rate in the low-dose and control groups. Overall, high-dose of folic acid and vitamin B12 can significantly improve plasma Hcy, coagulation indexes, and pathological changes in the venous thrombosis of the lower extremity.

Introduction

Deep venous thrombosis (DVT) of the lower extremity is a haemal disease involving abnormal circumfluence (1). Once the thrombus occurs, a small fraction of them ablate on their own; however, most thrombus spread gradually to the venous main body of the limbs (2). If timely clinical diagnosis and treatment are not provided, the situation will aggravate with thrombus sequelae, with serious health consequences. Research suggests the increase of homocysteine (Hcy) levels in plasma is closely related to DVT (3,4). Some studies propose that the metabolic process of hyperplasma Hcy is associated with folic acid and vitamin B12, and that folic acid and vitamin B12 can lower the plasma levels of Hcy (5-7). Thus, can folic acid and vitamin B12 be used to treat hyperhomocysteinemia with DVT? If so, what are the effects of different doses of folic acid and vitamin B12 on DVT with hyperhomocysteinemia?

In this study, we created a rabbit model with DVT and hyperhomocysteinemia and used different doses of folic acid and vitamin B12. We then examined the effect of each treatment to provide experimental support for the treatment of DVT with hyperhomocysteinemia with folic acid and vitamin B12.

Materials and methods

Experiment animals and treatments. Sixty male New Zealand rabbits of grade SPF were provided by the experimental animal center of Lanzhou University, with body mass of 2.1-3.1 kg. We created the rabbit model of DVT with hyperhomocysteinemia and randomly divided them into three groups: control (treated with normal saline), low-dose (treated with 5 mg/day folic acid + 0.25 mg/day vitamin B12) and high-dose (treated with 15 mg/day folic acid + 0.5 mg/day vitamin B12). Normal saline was obtained from Shanghai Di Ran Dang Cheng Pharmaceutical, folic acid tablets from Fujian Minghua Pharmaceutical, Fujian, China (National Medicine permission number: H19993229), and vitamin B12 from Datong Changxing Pharmaceutical, Shanxi, China (National Medicine permission number: H14022782). The rabbits were kept in a cage with controlled temperature (24°C) and free access to water. The study was approved by the Ethics Committee of the First Hospital of Lanzhou University.

Hemorheology detection. Three days and 10 days after modeling, 5 ml of whole blood were obtained from the ear artery from each animal and hemorheology indexes such as plasma viscosity, whole blood reduced viscosity at low shear, whole blood reduced viscosity at high shear, and red cell assembling index were detected using a hemorheology detector (Beijing Succeeder Science and Technology Development).

Coagulation function detection. Three days and 10 days after modeling, 3 ml whole blood were obtained from the ear artery from each animal, and coagulation function indexes such as fibrinogen, activated partial thromboplastin time, thrombin...
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Biochemical tests. The fully automated biochemical analyzer (Shanghai Hengsheng Medical Apparatus and Instruments Limited Company, Shanghai, China) was used to measure Hcy levels before and 10 days after DVT modeling by enzymatic cycling assay. Latex agglutination test was used to measure plasma D-dimer in days 1 and 10 days after DVT modeling. 10 days after DVT modeling, 5% thipentone of 0.1 ml/100 g body mass was injected to enterocoeleia for anesthesia. Then, rabbits were dissected to expose the saphenous artery, vein, saphenous vein, artery part of left hind limbs, to check thrombus and record if there is bump in the lower limbs. After blood vessel of left hind limb was separated, formalin was applied to fix and conventional pathological section was conducted, with hematoxylin and eosin (H&E) stain and pathologic histology change of deep venous thrombosis was inspected of rabbit lower extremity under microscopic examination. Color Doppler ultrasound (Jiangsu Jiahua Electronic Equipment Limited Company, Jiangsu, China) was applied to observe prognosis of thrombus.

Statistical analysis. Software SPSS 20.0 (IBM, New York, NY, USA) was used to process data. Enumeration data are shown by rate, \( \chi^2 \) was used to detect and measurement data are presented by mean ± standard deviation. Comparison between two groups was done by t-test and comparison among several groups applied F in detection. P<0.05 was considered to indicate a statistically significant difference.

Results

Hemorheology indexes. We measured hemorheology indexes at days 3 and 10 days after DVT modeling. At day 3, plasma viscosity, whole blood reduced viscosity at low shear, whole blood reduced viscosity at high shear, and red cell assembling
index were significantly lower in the low-dose and high-dose groups compared to controls (Table I). At day 10, all the values improved in the three groups, but the low- and high-dose groups still showed lower values than the control group (Table II). Additionally, the hemorheology indexes for the high-dose group were superior to those of the low-dose group (Tables I and II).

Coagulation function. We next measured coagulation indexes in the rabbits 3 and 10 days after DVT modeling. The levels of fibrinogen at days 3 and 10 were significantly lower in the two treatment groups compared with controls (Tables III and IV). The coagulation indexes APTT, PT, and TT were significantly higher in both treatment groups compared with controls (Tables III and IV). The coagulation function indexes of the high-dose group were superior to those of the low-dose group (Tables III and IV).

Hcy level. Before DVT modeling, the Hcy levels were comparable in the control, low-dose, and high-dose groups (Fig. 1). 10 days after DVT modeling, Hcy levels in the low-dose group were lower than in the control group (Fig. 2). The Hcy levels in the high-dose group were even lower than those in the low-dose group (Fig. 2).

D-dimer. Before DVT modeling, the level of D-dimer level was comparable in the three groups (Fig. 3). Ten days after modeling, the level of D-dimer in the low-dose group was lower than in the control group (Fig. 4). The Hcy levels in the high-dose group were even lower than those in the low-dose group (Fig. 4).

Table IV. Coagulation function indexes 10 days after modeling.

| Groups      | No. | Fibrinogen (g/l) | Activated partial thromboplastin time (s) | Thrombin time (s) | Prothrombin time (s) |
|-------------|-----|------------------|------------------------------------------|-------------------|----------------------|
| Control     | 20  | 4.16±0.41        | 15.05±0.62                               | 28.11±0.54        | 9.01±0.47            |
| Low-dose    | 20  | 3.64±0.75        | 17.83±1.12                               | 35.45±0.43        | 12.16±0.22           |
| High-dose   | 20  | 3.32±0.54        | 19.05±0.76                               | 38.33±0.46        | 14.43±0.55           |

\^p<0.05 compared with model control group; \^p<0.05 compared with low-dose group.

Figure 1. Hcy level before DVT modeling.

Figure 3. D-dimer level before DVT modeling.

Figure 2. Hcy level 10 days after DVT modeling.

Figure 4. D-dimer level 10 days after DVT modeling.
Vein pathology. Ten days after DVT modeling, we compared the change of lower extremity deep venous thrombosis of high-dose group and low-dose group were significantly reduced, while the improvement level of high-dose group was the best. Each layer of the normal femoral vein under light microscope recovered well in the high-dose group, without edema in loose connective tissue of adventitia (Fig. 5A). Partial serious intima damage was visible in the low-dose group, with endothelial falling partially (Fig. 5B). In the control group, endothelial cells of vein blood vessel fell off, with obvious adventitia edema, a great amount of fibrous protein aggregation in 'latticed' fibrin, full of red cells and a little deciduous endothelial cells, forming thrombus filling the whole lumen (Fig. 5C).

Thrombus recovery. Ten days after DVT modeling, the effective rate of treating lower extremity deep venous thrombosis with the high doses of folic acid and vitamin B12 was 100%, whereas the effective rate of the low-dose treatment was 75% (Table V). The control group showed no benefits and had an effective rate of 0% (Table V). Ten days after DVT modeling, the femoral venous wall echo was enhanced in the control group, with uneven intima echo incrassation and a little blood flow signal in the femoral vein (Fig. 6A). The femoral venous wall of the low-dose group thickened, with full solid and medium echo, blood flow signal in lumen around, dark color suggesting that flow rate was low (Fig. 6B). The femoral venous wall of the high-dose group thickened with strong echo in lumen, colorful blood flow presenting irregular fullness (Fig. 6C).

**Discussion**

Thrombus is caused by blood solidification and abnormal viscosity due to obstacles affecting its fluidity; this results in the formation of thrombocytes and activation of blood coagulation factor (8-10). DVT is one of the results where abnormal hemorheology and viscosity induce ischemia and anoxia of organs (11-13). Therefore, correcting blood viscosity and improving blood coagulation function indexes have an extremely important role in the prevention and treatment of DVT.

Our study showed that for 3 days and 10 days after DVT modeling, hemorheology and coagulation indexes were the best.
in the high-dose folic acid and vitamin B12 group. Some reports have proposed that Hcy is a thrombus-forming agent, and high Hcy could induce deep venous thrombosis (14-17). In addition, plasma D-dimer has been proposed to have a forewarning function for thrombus and hypercoagulable state (18,19). Ten days after DVT modeling, we found that the levels of Hcy and D-dimer improved the most in the high-dose group.

Through histopathologic examination, we found that 10 days after DVT modeling the high-dose and low-dose groups showed significant improvement in the change of lower extremity deep venous thrombosis. Deep venous thrombosis of lower extremity with color Doppler ultrasound diagnosis has some advantages, such as simple operation, safety and high efficiency, low cost, repeatability and direct observation, and the detection of related lesions and bleeding features around blood vessels (20). Our results also showed that after DVT modeling, the effective rate of lower extremity deep venous thrombosis treatment of high-dose group was 100%. The results of color Doppler ultrasound diagnosis of DVT corresponded to our findings in histopathologic examination. This validated the feasibility and high efficiency of color Doppler ultrasound diagnosis for deep venous thrombosis of the lower extremity.

In summary, folic acid and vitamin B12 have clear therapeutic effects in a rabbit model of DVT with hyperhomocysteinemia. The high-doses we tried improved various relevant symptoms, including the levels of Hcy and D-dimer, homocysteinemia. The high-doses we tried improved various relevant symptoms, including the levels of Hcy and D-dimer, homocysteinemia. The high-doses we tried improved various relevant symptoms, including the levels of Hcy and D-dimer, homocysteinemia. The high-doses we tried improved various relevant symptoms, including the levels of Hcy and D-dimer, homocysteinemia.

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