**Huangqi Guizhi Wuwu Decoction for treating diabetic peripheral neuropathy: a meta-analysis of 16 randomized controlled trials**

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**Funding:** This work was supported by a grant from the National Basic Research Program of China (973 Program), No. 2010CB530600; and Institutes Project from Guang’anmen Hospital of China Academy of Chinese Medical Sciences, No. 2011261.

**Graphical Abstract**

Clinical effects of *Huangqi Guizhi Wuwu* Decoction for treating diabetic peripheral neuropathy

**Abstract**

**OBJECTIVE:** This meta-analysis was performed to systematically assess the efficacy and safety of the Chinese herbal medicine *Huangqi Guizhi Wuwu* Decoction (HGWWD) for treating diabetic peripheral neuropathy.

**DATA SOURCES:** Six electronic databases, including the Cochrane Library, MEDLINE database, Chinese Biomedical Database, Chinese National Knowledge Infrastructure Database, Chinese Science and Technique Journals Database, and the Wanfang Database, were searched on the internet for randomized controlled trials published up until 1 December 2015. The search terms included “Chinese herbal medicine”, “diabetic peripheral neuropathy” and “randomized controlled trials” in Chinese and in English.

**DATA SELECTION:** We included randomized controlled trials using HGWWD/modified HGWWD for the treatment group, without restriction for the control group. We assessed literature quality in accordance with the Cochrane Review Handbook. A random or a fixed effects model was used to analyze outcomes using RevMan 5.2 software.

**OUTCOME MEASURES:** The primary outcomes were changes in symptoms and nerve conduction velocities. The secondary outcomes...
were fasting blood glucose and hemorheological indexes.

RESULTS: Sixteen randomized controlled trials, with a total of 1,173 patients, were included. Meta-analysis revealed that the efficacy of HGWWD for diabetic peripheral neuropathy was significantly superior compared with the control treatment (i.e., control group) (risk ratio = 0.36, 95% confidence interval (CI): 0.29–0.46, Z = 8.33, P < 0.00001) Compared with the control group, there was an increase in median motor nerve conduction velocity (mean difference (MD) = 3.46, 95% CI: 1.88–5.04, Z = 4.30, P < 0.01) and median sensory nerve conduction velocity (MD = 3.30, 95% CI: 2.04–4.56, Z = 5.14, P < 0.01). There was also an increase in peroneal motor nerve conduction velocity (MD = 3.22, 95% CI: 2.45–3.98, Z = 8.21, P < 0.01) and peroneal sensory nerve conduction velocity (MD = 3.05, 95% CI: 2.01–4.09, Z = 5.75, P < 0.01) in the treatment groups. No significant difference in fasting blood glucose was found between the treatment groups and the control groups (MD = −0.12, 95% CI: −0.42–0.19, Z = 0.76, P = 0.45). Plasma viscosity was significantly decreased after treatment (MD = −0.42, 95% CI: −0.21 to −0.02, Z = 2.30, P = 0.02). No significant difference in fibrinogen was detectable (MD = −0.53, 95% CI: −1.28–0.22, Z = 1.38, P = 0.17). Four trials reported that treatment groups experienced no adverse reactions. Adverse events were not mentioned in the other 12 trials. No trial reported the incidence of complications, quality of life outcomes, or health economics.

CONCLUSION: HGWWD treatment improves diabetic neurologic symptoms and ameliorates nerve conduction velocities. Our study suggests that HGWWD may have significant therapeutic efficacy for the treatment of diabetic peripheral neuropathy. However, the methodological quality of the randomized controlled trials was generally low. Larger and better-designed randomized controlled trials are required to more reliably assess the clinical effectiveness of HGWWD.

Key Words: nerve regeneration; meta-analysis; diabetic peripheral neuropathy; randomized controlled trials; Huangqi Guizhi Wuwu Decoction; traditional Chinese medicine; mecobalamin; efficacy; nerve conduction velocities; fasting blood glucose; hemorheology; neural regeneration

Background

Diabetic peripheral neuropathy (DPN) is a common and severe complication of diabetes mellitus (Dong et al., 2016). DPN can cause progressive nerve fiber degeneration as well as sensory and motor nerve dysfunction, which is characterized by unrelenting pain (hyperalgesia), sensory loss and limb impairment that can progress to muscular dystrophy in late DPN stages (Boulton et al., 2005). Recent reports suggest that ~50% of those with diabetes mellitus develop nerve damage within 10–20 years after diagnosis (Spruce et al., 2003; Tesfaye et al., 2012). Tesfaye’s group estimated that DPN may affect as many as 236 million people worldwide (Tesfaye et al., 2012). DPN symptoms worsen at night, and result in sleep and mood disturbance, and poor treatment causes further serious complications (diabetic foot ulcers, infections, and gangrene) (American Diabetes Association, 2013). Thus, DPN represents a major problem to the health care system (Davies et al., 2015). At present, treatment for DPN chiefly includes diabetes management via tight glucose control and pain relief for neuropathy (Hemmingson et al., 2013; Davies et al., 2015). Therapeutic strategies include aldose reductase inhibitors, antioxidant therapy, neurotrophic drugs and analgesics, which all have potential toxicity and side-effects, as well as poor tolerability. Moreover, many of these therapies may be ineffective for many diabetic patients (Gao et al., 2012). Thus, traditional Chinese medicine may hold some promise for specific DPN patients.

DPN may fall under the traditional Chinese medicine categories of “blood impeding” (China Association of Traditional Chinese Medicine, 2007). Clinical studies indicate that qi asthenia-associated blood stasis, and vessel and collateral obstruction are common in DPN (Wang et al., 2011; Zhou et al., 2013). Huangqi Guizhi Wuwu Decoction (HGWWD) is a classical prescription described in Jingui Yaoled, written by Zhang Zhong-jing during the Han dynasty. HGWWD is composed of Radix Astragali seu Hedysari, Ramulus Cinnamomi, Radix Paeoniae Alba, Rhizoma Zingiberis Recens and Fructus Jujubae, all of which are reported to improve qi and promote the flow of yang, invigorating the blood to promote coronary circulation (Bai et al., 2015). HGWWD is used to treat “blood impeding” characterized by a feeling of numbness and pain, and rough pulse, consistent with the features of DPN. Recent studies (Hua et al., 2008; Bian, 2010; Wang, et al., 2012) suggest that the mechanisms of action of HGWWD in the treatment of DPN in rats include an amelioration of morphological changes and an improvement of nerve function, probably through improving expression of multiple neurotrophic factors, enhancing blood flow and reducing free radical production. HGWWD may improve subjective symptoms associated with this progressive disabling disorder as well (Tong et al., 2006; Zhang et al., 2014). HGWWD exerts multiple therapeutic effects by acting on multiple targets. Although previous studies (Li, 2002; Liu, 2005; Chen et al., 2006; Ji et al., 2007; Bian et al., 2010; Wang and Luan, 2011; Hu et al., 2013) suggest that HGWWD is therapeutically effective for DPN, a systematic evaluation of these studies is required to more accurately assess the efficacy and safety of the medicine.

Systematic evaluation of randomized controlled trials (RCTs) is a reliable method for verifying the validity and safety of certain therapies. A few systematic reviews and meta-analyses (Chen et al., 2013) have examined the efficacy of Chinese herbal medicines for DPN, but these studies did not focus on a specific formulation. In the present systematic study, we evaluate the efficacy of a classical prescription—HGWWD—by evaluating clinical trials. Our findings should serve as a reference for clinicians seeking effective treatments for DPN.

Data and Methods

Database and search strategy

Trials were identified from the following six electronic databases: Cochrane Library, MEDLINE database, Chinese Biomedical Database, Chinese National Knowledge Infrastructure Database, Chinese Science and Technique Journals Database, and the Wanfang Database. We searched all trials published before 1 December 2015. Search terms (free words search) were as follows: “diabetes” OR “diabetic” OR “diabetes mellitus”; (“diabetic peripheral neuropathy” OR “peripheral nervous system disease” OR “diabetic neuropathies”) AND...
The following six items: random sequence generation (selection bias), allocation concealment (selection bias), blinding of participants and personnel (performance bias), blinding of outcome assessment (detection bias), incomplete outcome data (attrition bias), selective reporting (reporting bias), and other sources of bias. We categorized each item into one of three levels—“high risk”, “low risk” or “unclear—as follows: low risk of bias (all the items had a low risk of bias), high risk of bias (at least one item had a high risk of bias), and unclear risk of bias (at least one item had an unclear risk of bias). Discrepancies in interpretation were resolved by consensus or by a third party (Xiao-lin Tong).

Outcome measures
Clinical efficacy was the main outcome, and was based on changes in symptoms and nerve conduction velocities (NCVs) (Ministry of Health, China, 2002). The secondary outcomes were fasting blood glucose and hemorheological indexes at the end of treatment. Improvements were categorized as significantly effective (disappearance of subjective symptoms, recovered tendon reflex, and NCV increased significantly), effective (alleviated subjective symptoms, improved tendon reflex, and NCV increased) or ineffective (no improvement in symptoms, tendon reflex or NCV). NCV was assessed by median MNCV, median SNCV, peroneal MNCV and peroneal SNCV.

Statistical analysis
RevMan 5.2 software (Cochrane Collaboration, Oxford, UK) was used for data analysis. Publication bias was examined by funnel plots. In the outcomes, the data for efficacy were dichotomous, and others were continuous. Dichotomous data were expressed as the risk ratio (RR), and continuous outcomes between groups as mean difference (MD), both with a 95% confidence interval (95%CI), using a fixed-effects or random-effects model. A significant difference for the heterogeneity test was considered when $P < 0.05$, and substantial heterogeneity was indicated by an $I^2$ of 50% or more. In such cases, a random effects model was used; otherwise a fixed effects model was used (i.e., when $P > 0.05$ or $I^2 < 50%$).

Results
Description of trials
Our primary searches identified 372 references from the databases. Of these, 209 references were repeated literature and were excluded. After reading the titles and summaries, another 117 references were excluded because they were repeated literature, case reports, experimental studies, retrospective studies or literature reviews. A total of 46 references were retrieved for further assessment. After a detailed evaluation of the full text, an additional 30 references were excluded. Among these, seven trials were excluded because they used another traditional Chinese medicine therapy in the treatment group. Four trials were excluded because of incomplete or erroneous baseline or treatment data. Seven studies claimed that the trials were RCTs, but they lacked a control
group. The therapies in six trials were not in accordance with the inclusion criteria. The remaining six excluded trials were theoretical or non-clinical experiments. The search results are displayed in Figure 1.

Included trials
We included 16 trials (Li, 2002; Liu, 2005; Chen and Liu, 2006; Ji, 2007; Lian et al., 2007; Shu, 2007; Bian, 2010; Lu and You, 2010; Ye, 2010; Zhu, 2010; Wang and Luan, 2011; Hu and Wang, 2012; Jiang, 2012; Han, 2013; Yang, 2013; Zhang and Fu, 2013). All were RCTs with three parallel arms. All trials were conducted and published in China before 1 December 2015. The characteristics of the included trials are summarized in Table 1.

HGWWD decoction for treating DPN
Participants
In this meta-analysis, a total of 1,173 participants with DPN were included (619 patients and 554 controls). All trials were hospital-based and included inpatients and/or outpatients. Trial samples ranged from 36 to 225 participants, and a total of 622 males and 551 females were included. The age of participants ranged from 25 to 78 years. In six trials (Li, 2002; Liu, 2005; Lian et al., 2007; Lu and You, 2010; Zhu, 2010; Yang, 2013), enrolled patients suffered from type 2 diabetes, and in one trial (Shu, 2007), enrolled patients with either type 1 or type 2 diabetes were included. In nine trials, the type of diabetes was not reported. Diagnostic criteria for diabetes were necessary. Thirteen trials (Li, 2002; Liu, 2005; Chen and Liu, 2006; Lian et al., 2007; Shu, 2007; Bian, 2010; Ye, 2010; Zhu, 2010; Wang and Luan, 2011; Hu and Wang, 2012; Han, 2013; Yang, 2013; Zhang and Fu, 2013) mentioned World Health Organization diabetes mellitus diagnostic criteria (1999) or the American Diabetes Association criteria. In these 13 trials, acquired nerve symptoms, including pain, tingling, numbness, weakness, paralysis, abnormal tendon reflex and delayed nerve conduction were reported, and other neuropathic diseases were excluded. In three trials (Ji, 2007; Lu and You, 2010; Jiang, 2012), internal diabetes mellitus diagnostic criteria were used, and acquired nerve symptoms, including pain, tingling, numbness, weakness, paralysis, abnormal tendon reflex and delayed nerve conduction were reported, and other neuropathic diseases were excluded. Six trials (Li, 2005; Bian, 2010; Lu and You, 2010; Zhu, 2010; Hu et al., 2012; Han, 2013) mentioned that the pattern of the syndrome was consistent with qi asthenia caused by blood stasis, while the remaining 10 trials did not classify the syndrome according to traditional Chinese medical theory.

Treatments
Seven trials (Li, 2002; Liu, 2005; Shu, 2007; Bian, 2010; Lu and You, 2010; Zhu, 2010; Hu and Wang, 2012) used modified HGWWD, and compared this treatment group with a group given mecobalamin. In two trials (Wang, 2011; Jiang, 2012), modified HGWWD was compared with patients given no treatment. In seven trials (Chen and Liu, 2006; Ji, 2007; Lian et al., 2007; Ye, 2010; Han, 2013; Yang, 2013; Zhang and Fu, 2013), modified HGWWD plus mecobalamin was compared with mecobalamin alone. Hypoglycemic therapy was concomitantly given in both groups to control glycemia. Trial durations varied from 2 weeks to 12 weeks.

Outcomes
All trials used clinical efficacy based on changes in symptoms and NCV reported at the end of treatment, which was used as the main outcome index. We used FBG and hemorheological indexes as secondary outcome measures in the present meta-analysis. Adverse events were also recorded. None of the trials reported complications, reduced quality of life, or economic effects.

Risk of bias in the included trials
The quality assessment of the included trials is shown in Table 2. No trial reported details of sample size calculations, and none of the 16 trials were double-blind, placebo-controlled trials. Eight trials (Li, 2005; Chen, et al., 2006; Shu, 2007; Ji, 2007; Hu and Wang, 2012; Jiang, 2012; Han, 2013; Zhang and Fu, 2013) described methods of randomization using a random number table. The remaining eight trials indicated "randomly allocating", but the method of randomization was not provided. No trial stated how allocation concealment or blinding was performed. All the 16 trials provided complete baseline information, and described similarities between comparison groups. No trial reported participant losses, so attrition bias was uncertain. Selective reporting was difficult to assess, and trial protocols were unavailable. All trials were generally assessed low in quality and contained risk of bias, suggesting that future studies might influence the confidence intervals in this meta-analysis and require revising the conclusion.

Meta-analysis results
Efficacy
The results of the 16 trials were included in our meta-analysis and demonstrated a significant difference in clinical efficacy between the treatment groups and the control groups. These trials showed insignificant heterogeneity of the trial results ($\chi^2 = 8.49$, $P = 0.83$, $I^2 = 0\%$). Thus, a fixed effects model was used for statistical analysis. Treatment groups were superior to control groups in terms of efficacy ($n = 1,173$, $RR = 0.36$, 95%CI: 0.29–0.46, $Z = 8.33$, $P < 0.00001$) ($n = 1,173$, $RR = 1.33$, 95%CI: 1.24–1.42, $Z = 8.39$, $P < 0.01$). To compare the clinical efficacy of modified HGWWD with the control group, subgroup analysis was performed. There was a significant difference between HGWWD alone and the control groups ($n = 566$, $RR = 0.31$, 95%CI: 0.22–0.45, $Z = 6.42$, $P < 0.00001$), without heterogeneity (chi-square=1.70, $P = 0.99$, $I^2 = 0\%$). A significant difference in clinical efficacy was also found between HGWWD plus conventional Western drug and control groups ($n = 607$, $RR = 0.41$, 95%CI: 0.30–0.57, $Z = 5.34$, $P < 0.00001$; $n = 607$, $RR = 1.28$, 95%CI: 1.17–1.40, $Z = 5.39$, $P < 0.01$), without heterogeneity ($n = 607$, $RR = 0.41$, 95%CI: 0.30–0.57, $Z = 5.34$, $P < 0.00001$) ($\chi^2$...
The pooled analysis of NCVs taken as a continuous measurement was not different between treatment groups in any of the trials. After 2 to 12 weeks, NCVs differed significantly between treatment groups and control groups. To compare NCVs in the modified HGWWD groups with those in the control groups, subgroup analysis was performed.

Eight trials, involving a total of 711 patients, reported median MNCV as an outcome ($MD = 3.46, 95\% CI: 1.88–5.04, Z = 4.30, P < 0.01$) (Figure 3). Significant heterogeneity between trials was observed ($\chi^2 = 50.75, P < 0.01, I^2 = 86\%$), and accordingly, a random effects model was used for statistical analysis. Four trials compared the median MNCV between HGWWD alone and a control group ($n = 286, MD = 3.56, 95\% CI: 0.76–6.36, Z = 2.49, P = 0.01$), with heterogeneity ($\chi^2 = 32.71, P < 0.01, I^2 = 91\%$), and four trials compared the median MNCV between HGWWD plus conventional Western drug and a control group ($n = 425, MD = 3.31, 95\% CI: 1.46–5.15, Z = 3.51, P < 0.01$), with heterogeneity ($\chi^2 = 14.87, I^2 = 80\%, P < 0.01$).

Nine trials, involving a total of 765 patients, reported mean SNCV as an outcome ($MD = 3.30, 95\% CI: 2.04–4.56, Z = 5.14, P < 0.01$) (Figure 4). Significant heterogeneity between trials was observed ($\chi^2 = 38.39, P < 0.01, I^2 = 79\%$), and therefore, a random effects model was used for statistical analysis. Five trials compared the median SNCV between HGWWD alone and a control group ($n = 340, MD = 3.47, 95\% CI: 1.61–5.32, Z = 3.66, P < 0.01$), with heterogeneity ($\chi^2 = 16.18, I^2 = 75\%, P < 0.01$), and four trials compared the median SNCV between HGWWD plus conventional Western drug and a control group ($n = 425, MD = 3.14, 95\% CI: 1.17–5.12, Z = 3.11, P < 0.01$), with heterogeneity ($\chi^2 = 22.21, I^2 = 86\%, P < 0.01$).

Nine trials, involving a total of 765 patients, reported peroneal MNCV as an outcome ($MD = 3.22, 95\% CI: 2.45–3.98, Z = 8.21, P < 0.01$) (Figure 5). Heterogeneity between trials was significant ($\chi^2 = 16.02, P = 0.04, I^2 = 50\%$), and accordingly, a random effects model was used for statistical analysis. Five trials compared the peroneal MNCV between HGWWD alone and a control group ($n = 340, MD = 3.87, 95\% CI: 3.07–4.66, Z = 9.54, P < 0.01$), without heterogeneity ($\chi^2 = 2.77, P = 0.60, I^2 = 0\%$), and four trials compared the peroneal MNCV between HGWWD plus conventional Western drug and a control group ($n = 425, MD = 2.52, 95\% CI: 1.56–3.48, Z = 5.15, P < 0.01$), with heterogeneity ($\chi^2 = 5.82, P = 0.12, I^2 = 48\%$).

Eight trials, involving a total of 711 patients, reported peroneal SNCV as an outcome ($MD = 3.05, 95\% CI: 2.01–4.09, Z = 5.75, P < 0.01$) (Figure 6). Significant heterogeneity between trials was observed ($\chi^2 = 26.65, P = 0.04, I^2 = 74\%$), and therefore, a random effects model was used for statistical analysis. Four trials compared peroneal SNCV between HGWWD alone and a control group ($n = 286, MD = 4.27, 95\% CI: 2.73–5.82, Z = 5.43, P < 0.01$), without heterogeneity ($\chi^2 = 9.83, P = 0.02, I^2 = 69\%$), and four trials compared the peroneal SNCV between HGWWD plus conventional Western drug and a control group ($n = 425, MD = 1.94, 95\% CI: 1.27–2.62, Z = 5.68, P < 0.01$), without heterogeneity ($\chi^2 = 0.48, P = 0.92, I^2 = 0\%$).

### Glycemic control

In four trials (Li, 2002; Liu, 2005; Bian, 2010; Lu and You, 2010), changes in fasting blood glucose levels were assessed, and they did not show heterogeneity ($\chi^2 = 1.78, I^2 = 0\%, P = 0.62$). Thus, a fixed effects model was used for statistical analysis. A meta-analysis of four trials revealed no significant difference in changes in fasting blood glucose levels between the treatment groups and the control groups ($n = 204, MD = –0.12, 95\% CI: –0.42–0.19, Z = 0.76, P = 0.45$) (Figure 7).
Figure 2 Clinical efficacy of HGWWD in the treatment of DPN.
Treatment groups (HGWWD/HGWWD + mecobalamin) were superior to control groups (mecobalamin/no treatment) for efficacy ($RR = 0.36$, 95% CI: 0.29–0.46, $Z = 8.33$, $P < 0.00001$). HGWWD: Huangqi Guizhi Wuwu Decoction; DPN: diabetic peripheral neuropathy; $RR$: risk ratio; CI: confidence interval.

Figure 3 Effects of HGWWD on median MNCV in patients with DPN.
Treatment groups (HGWWD/HGWWD + mecobalamin) were superior to control groups (mecobalamin/no treatment) for improvement of median MNCV ($MD = 3.46$, 95% CI: 1.88–5.04, $P < 0.01$). HGWWD: Huangqi Guizhi Wuwu Decoction; MNCV: motor nerve conduction velocity; DPN: diabetic peripheral neuropathy; $MD$: mean difference; CI: confidence interval.
Figure 4 Effects of HGWWD on median SNCV in patients with DPN. Treatment groups (HGWWD/HGWWD + mecobalamin) were superior to control groups (mecobalamin/no treatment) for improvement of median SNCV ($MD = 3.30, 95\% CI: 2.04–4.56, P < 0.01$).

HGWWD: Huangqi Guizhi Wuwu Decoction; SNCV: sensory nerve conduction velocity; DPN: diabetic peripheral neuropathy; $MD$: mean difference; CI: confidence interval.

Figure 5 Effects of HGWWD on peroneal SNCV in patients with DPN. Treatment groups (HGWWD/HGWWD + mecobalamin) were superior to control groups (mecobalamin/no treatment) for improvement of peroneal SNCV ($MD = 3.05, 95\% CI: 2.04–4.09, P < 0.01$).

HGWWD: Huangqi Guizhi Wuwu Decoction; SNCV: sensory nerve conduction velocity; DPN: diabetic peripheral neuropathy; $MD$: mean difference; CI: confidence interval.

Figure 6 Effects of HGWWD on peroneal MNCV in patients with DPN. Treatment groups (HGWWD/HGWWD + mecobalamin) were superior to control groups (mecobalamin/no treatment) for improvement of peroneal MNCV ($MD = 3.22, 95\% CI: 2.45–3.98, P < 0.01$).

HGWWD: Huangqi Guizhi Wuwu Decoction; SNCV: sensory nerve conduction velocity; DPN: diabetic peripheral neuropathy; $MD$: mean difference; CI: confidence interval.
Table 1 Characteristics of the trials included in the meta-analysis

<table>
<thead>
<tr>
<th>Source</th>
<th>Sample (T/C)</th>
<th>Age (year)</th>
<th>Sex (male/female)</th>
<th>Intervention</th>
<th>Treatment group</th>
<th>Control group</th>
<th>Treatment period (week)</th>
<th>DPN duration (year)</th>
<th>Outcome measure</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bian, 2010</td>
<td>20/20</td>
<td>T: 62.6±9.9</td>
<td>C: 60.4±9.1</td>
<td>Modified HGWWD (200 mL/d, bid)</td>
<td>Mecobalamin (500 μg, tid)</td>
<td>8</td>
<td>T: 5.6±3.8</td>
<td>C: 5.2±3.3</td>
<td>CER+NCV+Glu</td>
</tr>
<tr>
<td>Hu and Wang, 2012</td>
<td>30/30</td>
<td>T: 65.3±7.0</td>
<td>C: 64.3±6.2</td>
<td>Modified HGWWD</td>
<td>Mecobalamin (0.5 mg, tid)</td>
<td>8</td>
<td>T: 5.4±2.8</td>
<td>C: 5.5±3.7</td>
<td>CER+symptom scores</td>
</tr>
<tr>
<td>Li, 2002</td>
<td>34/20</td>
<td>T: 58.4±8.4</td>
<td>C: 56.7±8.6</td>
<td>Modified HGWWD (bid)</td>
<td>Mecobalamin (500 μg, tid)</td>
<td>8</td>
<td>T: 3.4±0.9</td>
<td>C: 3.6±0.8</td>
<td>CER+NCV+</td>
</tr>
<tr>
<td>Liu, 2005</td>
<td>24/24</td>
<td>T: 51.3±6.5</td>
<td>C: 51.8±7.0</td>
<td>Modified HGWWD (300 mL/d, bid)</td>
<td>Mecobalamin (500 μg, tid)</td>
<td>8</td>
<td>T: 4.5±1.2</td>
<td>C: 4.5±1.1</td>
<td>CER+Glu+ HR</td>
</tr>
<tr>
<td>Lu and You, 2010</td>
<td>32/30</td>
<td>T: 57.2±5.2</td>
<td>C: 61.7±11.1</td>
<td>Modified HGWWD (bid)</td>
<td>Mecobalamin (0.5 mg, tid)</td>
<td>8</td>
<td>T: 6.5±4.8</td>
<td>C: 5.6±4.4</td>
<td>CER+NCV+</td>
</tr>
<tr>
<td>Shu, 2007</td>
<td>34/34</td>
<td>T: 49.6±7.2</td>
<td>C: 51.7±9.3</td>
<td>Modified HGWWD (400 mL/d, bid)</td>
<td>Mecobalamin (500 μg, tid)</td>
<td>8</td>
<td>T: 1.8±1.49</td>
<td>C: 1.9±1.33</td>
<td>CER+NCV</td>
</tr>
<tr>
<td>Zhu, 2010</td>
<td>38/32</td>
<td>T: 49.5±14.5</td>
<td>C: 47.5±14.5</td>
<td>Modified HGWWD (bid)</td>
<td>Mecobalamin (500 μg, tid)</td>
<td>2</td>
<td>Not reported</td>
<td>CER</td>
<td></td>
</tr>
<tr>
<td>Wang and Luan, 2011</td>
<td>158/50</td>
<td>T: 46.3±8.7</td>
<td>C: 45.8±5.4</td>
<td>Modified HGWWD (bid)</td>
<td>Blank</td>
<td>10</td>
<td>T: 2.5±0.6</td>
<td>C: 2.6±0.7</td>
<td>CER+NCV</td>
</tr>
<tr>
<td>Jiang, 2012</td>
<td>33/33</td>
<td>T: 53.4±6.3</td>
<td>C: 54.2±5.0</td>
<td>Modified HGWWD (300 mL/d, bid)</td>
<td>Blank</td>
<td>12</td>
<td>1 month — 4 years</td>
<td>CER</td>
<td></td>
</tr>
<tr>
<td>Chen and Liu, 2006</td>
<td>34/28</td>
<td>T: 58.6±5.2</td>
<td>C: 54.6±6.5</td>
<td>Modified HGWWD (200 mL/d, bid)+ mecobalamin (500 μg, tid)</td>
<td>Mecobalamin (500 μg, tid)</td>
<td>6</td>
<td>T: 3.6±3.2</td>
<td>C: 7.8±3.6</td>
<td>CER+NCV</td>
</tr>
<tr>
<td>Han, 2013</td>
<td>31/31</td>
<td>T: 55.3±10.1</td>
<td>C: 54.2±9.6</td>
<td>Modified HGWWD+ mecobalamin (0.5 mg, tid)</td>
<td>Mecobalamin (500 μg, tid)</td>
<td>8</td>
<td>Not reported</td>
<td>CER+NCV+ symptom scores</td>
<td></td>
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<tr>
<td>Ji, 2007</td>
<td>33/32</td>
<td>T: 56.0±7.9</td>
<td>C: 55.0±8.0</td>
<td>Modified HGWWD (400 mL/d, bid)+ mecobalamin (500 μg, tid)</td>
<td>Mecobalamin (500 μg, tid)</td>
<td>12</td>
<td>T: 5.1±1.2</td>
<td>C: 4.9±1.1</td>
<td>CER</td>
</tr>
<tr>
<td>Lian et al., 2007</td>
<td>122/103</td>
<td>T: 53.9±6.9</td>
<td>C: 53.1±7.3</td>
<td>Modified HGWWD (300 mL/d, bid)+ mecobalamin (0.5 mg, tid)</td>
<td>Mecobalamin (500 μg, tid)</td>
<td>8</td>
<td>T: 5.1±2.9</td>
<td>C: 5.2±3.1</td>
<td>CER+NCV</td>
</tr>
<tr>
<td>Zhang and Fu, 2013</td>
<td>38/38</td>
<td>T: 49.2±6.5</td>
<td>C: 48.6±6.3</td>
<td>Modified HGWWD (200 mL/d, bid)+ mecobalamin (500 μg/d)</td>
<td>Mecobalamin (500 μg/d)</td>
<td>4</td>
<td>T: 4.2±1.4</td>
<td>C: 4.0±1.7</td>
<td>CER+NCV</td>
</tr>
<tr>
<td>Ye, 2010</td>
<td>18/18</td>
<td>T: 62.3</td>
<td>C: 62.23</td>
<td>Modified HGWWD+ mecobalamin (500 μg/d)</td>
<td>Mecobalamin (500 μg/d)</td>
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<td>T: 3.65</td>
<td>C: 3.34</td>
<td>CER</td>
</tr>
<tr>
<td>Yang, 2013</td>
<td>40/41</td>
<td>T: 52.6±14.7</td>
<td>C: 53.2±15.1</td>
<td>Modified HGWWD (400 mL/d, bid)+ mecobalamin (0.5 mg, tid)</td>
<td>Mecobalamin (500 μg, tid)</td>
<td>8</td>
<td>Not reported</td>
<td>CER</td>
<td></td>
</tr>
</tbody>
</table>

T: Treatment group; C: control group; CER: clinical effectiveness; NCV: nerve conduction velocity; Glu: blood glucose; HR: hemorheological indexes; HGWWD: Huangqi Guizhi Wuwu Decoction; DPN: diabetic peripheral neuropathy; bid: bis in die; tid: ter in die.

Hemorheology

Two trials (Li, 2002; Liu, 2005) evaluated changes in hemorheological indexes. In this review, we mainly assessed plasma viscosity and fibrinogen variations. Two trials did not show evidence of heterogeneity in plasma viscosity (χ² = 0.48, P = 0.49, I² = 0%). Thus, a fixed effects model was used for statistical analysis. There was a statistically significant difference between the two groups in plasma viscosity (n = 102, MD = -0.11, 95% CI: -0.21 to -0.02, Z = 2.50, P = 0.012) (Figure 8). Heterogeneity in fibrinogen between trials was significant (χ² = 3.93, P = 0.05, I² = 75%), and therefore, a random effects model was used for statistical analysis. The meta-analysis of the two trials showed that the treatment groups were not statistically different from the control groups in the reduction in fibrinogen (n = 102, MD = -0.53, 95% CI: -1.28 to -0.22, Z = 1.38, P = 0.17) (Figure 9).

Adverse and other effects

In four trials (Liu, 2005; Bian, 2010; Wang and Luan, 2011; Jiang, 2012), the treatment groups were reported to have experienced no adverse reactions. Adverse events were not mentioned in the other 12 trials, and therefore, the safety of the Chinese herbal medicine therapy could not be assessed. Only one trial (Liu, 2005) reported follow-up.
The funnel shape of the plot was not completely symmetrical, indicating a potential publication bias. Although we conducted comprehensive searches and attempted to avoid bias, all trials were published in China, and accordingly, we could not exclude potential publication bias (Figure 10).

**Discussion**

DPN is a chronic, progressive disease. Treatment for DPN is aimed at improving the quality of life and preventing further nerve damage and complications (diabetic foot ulcers, infection, and gangrene) (American Diabetes Association, 2013). Because treatment by conventional Western medicine is of limited clinical efficacy, safer and more effective therapies are urgently needed. Clinical practice has shown that the combination of traditional Chinese medicine and Western medicine has substantial therapeutic potential for treating diabetes and its complications (Tong, 2012). Clinical trials and research on traditional Chinese medicine for treating DPN, including herbs and acupuncture, have been reported (Perkins et al., 2001; Boulton et al., 2005). However, no trial evaluated DPN, and the scores correlate with NCV examinations. The Toronto Clinical Scoring System is widely used to evaluate DPN, and the scores correlate with NCV examinations (Perkins et al., 2001; Boulton et al., 2005). However, no trial used this scoring system, and three trials (Lu et al., 2010; Hu et al., 2012; Han, 2013) adopted a questionnaire based on the HGWWD for DPN was significantly superior to that of the treatments for the control group. This finding suggests that HGWWD has therapeutic efficacy for DPN. However, the methodological quality of the trials was generally low, and there is the possibility of bias. Therefore, a more definitive statement of the clinical efficacy of HGWWD requires larger and better-designed RCTs.

There are a number of limitations of this meta-analysis. Although the decoction of HGWWD in the included trials was generally similar, and administration was twice per day, the herb ingredients, dosage and treatment periods varied among the trials. The period of treatment ranged from 2 weeks to 12 weeks. No trial reported sample size calculations, so assessment of clinical efficacy was made difficult, which might have negatively impacted the reliability of the outcomes.

We assessed clinical efficacy based on changes in symptoms and NCV as the main outcome indexes. Only half of the clinical trials provided NCV examination results, and those that did, examined different nerves. Consequently, clinical trials should standardize DPN NCV examinations to enhance the reliability of analysis. Limb discomfort is an important clinical characteristic, and clinical efficacy assessment should include evaluation of symptomatic variation. The Toronto Clinical Scoring System is widely used to evaluate DPN, and the scores correlate with NCV examinations (Perkins et al., 2001; Boulton et al., 2005). However, no trial used this scoring system, and three trials (Lu et al., 2010; Hu et al., 2012; Han, 2013) adopted a questionnaire based on the

### Table 2 Quality assessment of the trials included in the meta-analysis

<table>
<thead>
<tr>
<th>Study</th>
<th>Random sequence generation</th>
<th>Allocation concealment</th>
<th>Blinding of participants, personnel and outcome assessment</th>
<th>Incomplete outcome data</th>
<th>Selective reporting</th>
<th>Other bias</th>
<th>Risk of bias</th>
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<td>Bian, 2010</td>
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<td>High</td>
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<tr>
<td>Liu, 2005</td>
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<tr>
<td>Lu and You, 2010</td>
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<tr>
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<tr>
<td>Chen and Liu, 2006</td>
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<tr>
<td>Han, 2013</td>
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<tr>
<td>Ji, 2007</td>
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<tr>
<td>Zhang and Fu, 2013</td>
<td>Random number table</td>
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<tr>
<td>Ye, 2010</td>
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<tr>
<td>Yang, 2013</td>
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</table>
Figure 7 Effects of HGWWD on fasting blood glucose in patients with DPN.

Treatment groups (HGWWD) were not statistically different from control groups in decreasing fasting blood glucose ($MD = -0.12, 95\% CI = -0.42–0.19, Z = 0.76, P = 0.45$). HGWWD: Huangqi Guizhi Wuwu Decoction; DPN: diabetic peripheral neuropathy; $MD$: mean difference; $CI$: confidence interval.

Figure 8 Effects of HGWWD on plasma viscosity in patients in DPN.

Treatment groups (HGWWD) were not statistically different from control groups in decreasing plasma viscosity ($MD = -0.11, 95\% CI = -0.21–0.02, P = 0.02$). HGWWD: Huangqi Guizhi Wuwu Decoction; DPN: diabetic peripheral neuropathy; $MD$: mean difference; $CI$: confidence interval.

Figure 9 Effects of HGWWD on fibrinogen in patients with DPN.

Treatment groups (HGWWD) were not statistically different from the control groups in decreasing fibrinogen ($MD = -0.33, 95\% CI = -1.28–0.22, P = 0.17$). HGWWD: Huangqi Guizhi Wuwu Decoction; DPN: diabetic peripheral neuropathy; $MD$: mean difference; $CI$: confidence interval.

Guiding principles for clinical research of new Chinese patient medicine (Ministry of Health, China, 2002). Thus, it is urgent to standardize the assessment of symptoms. The clinical trials mainly assessed outcomes using substitutive indexes (NCV, and response rate of clinical symptoms); however, no trials assessed quality of life in patients with endpoint criteria (incidence of diabetic foot ulcers, amputation rates, and disability rates).

In the included trials, only four trials (Li, 2002; Liu, 2005; Bian, 2010; Lu and You, 2010) examined fasting blood glucose changes. Although our meta-analysis revealed no significant difference in this variable between the treatment and control groups, we could not fully evaluate the influence of blood glucose. Only a few trials reported hypoglycemic effects, but some reports suggest that Huangqi (Radix Astragali seu Hedysari) and Baishao...
(Radix Paeoniae Alba) have hypoglycemic effects, and these herbs are components of HGWWD. Therefore, it is unclear whether the HGWWD formulation affects blood glucose and whether the amelioration of NCV is related to improvement of blood glucose. Although one trial (Zhou et al., 2013) proposed that the HGWWD formulation may improve blood glucose, they did not provide support for this statement.

In the present meta-analysis, we found a statistically significant difference in the reduction in plasma viscosity, although there was no statistically significant difference in the reduction in fibrinogen. Previous studies (Hua et al., 2008; Wang et al., 2009, 2010, 2012) suggest that HGWWD inhibits coagulation and platelet aggregation. One trial (Wang, et al., 2009) showed that Huangqí (Radix Astragali seu Hedysari) decreases blood viscosity and inhibits coagulation and platelet aggregation. Guízhi (Ramulus Cinnamomum) dilates blood vessels and regulates the circulation. Nevertheless, clinical data are presently lacking, and additional clinical trials and experiments are needed to clarify the effect of HGWWD on the circulatory system.

Quality of the evidence
All of the RCTs were of very low quality in terms of design, reporting and methodology. Eight trials (Liu, 2005; Chen and Liu, 2006; Shu, 2007; Ji, 2007; Jiang, 2012, Hu and Wang, 2012; Han, 2013; Zhang and Fu, 2013) mentioned the use of a random number table, while another eight trials only indicated “randomly allocating”; with no detailed information. No trial mentioned allocation concealment, suggesting selection bias. No trials mentioned blinding methods that might rule out performance and detection biases.

In this meta-analysis, NCV was an objective index that should not be affected by blinding, but patients and researchers were aware of the treatments, which might affect subjective symptoms. Herbs were not randomized, and the traditional Chinese medicine treatment was not compared to an approved medication for DPN. Instead, it was compared to an alternative or complementary method for treating DPN. No trials reported participant losses, so attrition bias was unclear. As such, the improvements in qualitative outcomes should be regarded with caution. All trials were published in China.

Potential biases in the review process
The asymmetrical funnel plot demonstrates the potential publication bias. Funnel plots are a visual aid to identify publication bias or systematic heterogeneity. All of the 16 trials were included in these funnel plots, recognizing the substantial heterogeneity of the treatment, trial size and design. However, none of the trials found a negative effect, indicating publication bias. Although we undertook extensive searches for unpublished literature, we found no negative trials. However, trials with large positive results are often much easier to publish than trials with negative results. Therefore, it is likely that publication bias is present, affecting the reliability of the meta-analysis.

Safety and follow-up
Inadequate reporting on adverse events in the included trials was a problem. Four trials reported that the treatment groups experienced no adverse reactions. Adverse events were not mentioned in other 12 trials. Only one trial (Liu, 2005) reported follow-up. Therefore, long-term effectiveness, side effects, and health outcomes could not be evaluated.

Future clinical trials should put emphasis on the following aspects: (1) Enhancing the quality of the methodology, including assessing sample size, increasing the power of the test, and improving randomization, allocation concealment, the blinding method, as well as safety reporting and detailed follow-up. (2) Side effects and long-term effects of HGWWD should be studied along with an objective endpoint (diabetic foot ulcers, amputation and disability rates) to evaluate the quality of life. (3) Future trials should standardize the composition of the herbal mixture, and examine the dose-effect relationships. This should help optimize the efficacy of the Chinese herbal medicine, either when used alone or in combination with a Western medical treatment.

Conclusion
Our meta-analysis suggests that the Chinese herbal medicine HGWWD is better for improving NCVs and alleviating subjective DPN symptoms compared with control treatment. However, the long-term effectiveness and safety of HGWWD for DPN are uncertain, and most trials included in this review were of poor quality. Thus, the limitations of this meta-analysis should be taken into consideration. Well-designed, large-scale, high-quality multicenter RCTs are required to provide better outcome assessments. Furthermore, long-term outcome observations, including mortality, quality of life and adverse events, are also necessary to more reliably assess the clinical efficacy of HGWWD for DPN.

Author contributions: BP and TYZ conceived the study and wrote the paper. LHZ and FW did data collection. RY and QZ were responsible for the statistical analysis. XLT participated in the design of the study and coordination. All authors approved the final version of the paper.

Conflicts of interest: None declared.

Plagiarism check: This paper was screened twice using Cross-Check to verify originality before publication.

Peer review: This paper was double-blinded and stringently reviewed by international expert reviewers.

References


