Reevaluation of the post-marketing safety of Shuxuening injection based on real-world and evidence-based evaluations

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Aim: To evaluate the factors influencing suspected hypersensitivity and adverse systemic reactions after Shuxuening injection and to provide innovative ideas and methods for the reevaluation of post-marketing safety of Shuxuening.

Methods: This study used a prospective, nested case–control study design, combined with a prescription sequence analysis design method. It classified patients who exhibited trigger signals after administration of Shuxuening injection as suspected allergic patients and made comparisons with patients who did not report adverse effects to calculate the correlation between relevant risk factors and suspected allergic reactions. Randomized controlled studies and cohort studies of the adverse drug reaction (ADR) of Shuxuening were performed using a computer database. Data retrieval was carried out by the foundation governing the individual database. Meta-analysis was performed by using R3.2.3 software to evaluate the ADRs of Shuxuening.

Results: The results of real-world study showed that administration of Shuxuening in combination with potassium aspartate and magnesium, atorvastatin calcium, Shengmai injection, pantoprazole sodium, or high-dose medication was a risk factor for suspected allergic reactions. Meta-analysis showed that the incidence of adverse events was 5.84% (95% CI 0.0499; 0.0674), and serious adverse reaction rate was 4.36% (95% CI 0.0188; 0.0760) when Shuxuening was used in combination with these drugs. The incidence of allergic reaction was also influenced by the vehicle, duration of treatment, single dose, and indicated vs off-label use.

Conclusion: Risk factors for adverse reaction following the use of Shuxuening injection in patients are associated with a single dose, vehicle, type of disease, and combination with potassium aspartate, atorvastatin calcium, Shengmai injection, injection with pantoprazole sodium, and other drugs. Physicians should be careful to follow guidelines when administering this drug. We further propose that the unique methodology used in this study may be useful for reevaluation of the safety of other traditional Chinese medicines.

Keywords: real-world study, evidence-based evaluation, Shuxuening injection, post-marketing safety reevaluation, allergic reaction

Introduction

Shuxuening is a traditional Chinese medicine delivered by injection. It is an aqueous extract of ginkgo leaf, with sorbitol, 95% ethanol, and methyl sulfide as the excipients. The active ingredients of Shuxuening are total flavonoid glycosides and ginkgolides, whose main effect is to cause vasodilation thereby improving microcirculation. Principal indications include use in the treatment of ischemic cardiovascular and cerebrovascular diseases, coronary heart disease, angina, cerebral embolism, and cerebral
vasospasm. In 2013, the China Medical Insurance Research Association released a report titled “2009 – 2011 National Urban Basic Medical Insurance in patients with traditional Chinese medicine utilization analysis”. They reported that the use of Shuxuening injection ranked first among that of internal medical drugs. Unfortunately, Shuxuening is often misused, and clinical adverse reaction reports are not uncommon. The National Drug Adverse Reaction Monitoring Annual Report mentioned that the incidence of adverse reactions/events associated with Shuxuening injection ranked relatively high. At present, the study of adverse drug reactions (ADRs) after Shuxuening injection is mainly based on case reports, literature reviews, and analyses. No reports exist regarding the incidence of allergic reactions to Shuxuening injection or the incidence of resulting anaphylactic shock. Thus, analysis and investigation of real-world user information from a large sample of patients who received Shuxuening injections is important. A comprehensive analysis of adverse reactions to Shuxuening reported in the literature was carried out to provide a reliable reference for the clinical use of Shuxuening.

This study is based on capturing trigger signals through combined nested case–control studies (NCCSs) with prescription sequence analysis (PSA) to perform real-world evaluation of the incidence of adverse reactions to Shuxuening. PSA, as first introduced by Petri et al, is a research method based on drug prescription records to detect drug reactions. Its use requirements are based on existing, complete prescription record databases and can be applicable when the use of certain medications signal an ADR to another drug. The hospital information system (HIS) records a large amount of real-world clinical data as well as a complete record of all medication information of patients undergoing hospitalization. Although the HIS database does not document specifically whether the patient had an allergic reaction, such reactions can be inferred, if, for example, certain medications, such as dexamethasone, are administered subsequent to Shuxuening injection.

The NCCS was first proposed by the American epidemiologist Mantel in 1973. Also called the synthetic case–control study or case–control study within a cohort, cases of a disease that occur in a defined cohort are identified in the NCCS and, for each, a specified number of matched controls are selected among those in the cohort who have not developed the disease by the time of disease occurrence in the case. Then, the collected information and biological specimens of the 2 groups of patients are compiled for necessary testing and sorted for statistical analysis.

To fully investigate the incidence of adverse reactions following Shuxuening injection, we combined PSA with NCCS to extract and study a large data sample in the real world. Then we combined the information of the large data sample with the evidence-based evaluation, and also combined patient medical records and the literature database to explore factors that affect adverse reactions, incidence of serious adverse reactions, and rare adverse reactions to Shuxuening injection.

**Methods**

**Data sources**

**Real-world data sources**

Data were collected from medical records of patients of the Bengbu Medical College First Affiliated Hospital from July 1, 2015, to June 30, 2017. During that time, a total of 7,208 patients were treated with Shuxuening injections. An allergic group and a matched control group were selected from this pool of available records.

**Evidence-based data sources**

For evidence-based analyses, we searched PubMed, Embase, the Cochrane Library, CNKI, VIP, Wanfang Data, and CBM from the foundation date of each database to June 30, 2017. We combined the keywords and free words (“Shuxuening injection”, “side effect”, “safety”, and “adverse drug reaction”) to retrieve data and then screened the literature according to the inclusion criteria.

**Research design**

**Realistic research design**

We used a forward-looking, single-center design method of NCCS combined with PSA to perform real-world analyses. Based upon the existing literature, the following signals were set as triggers for defining a hypersensitive response in this study. The patient pool was divided into an exposure group and a nonexposed group, depending on whether individual patients were administered anti-allergic drugs, such as dexamethasone, promethazine, or loratadine, while using vitamin C and calcium gluconate or other allergic shock drugs from the onset of adverse reactions to Shuxuening injection to reverse its effects. Adverse reactions to Shuxuening were divided into delayed-type hypersensitivity and rapid-onset allergic reactions. Most of the rapid allergic reactions appeared within 30 min of administration, whereas most of the late-onset ADR occurred within a day of medication administration. Therefore, we focused on suspected allergic reactions.
occurring within 24 h of treatment. Specific groups were as follows.

Exposure group
This group comprised patients who discontinued use of Shuxuening in the first 24 h, those who had a record of only 1 injection of Shuxuening, and those who used anti-allergic drugs within 24 h after stopping Shuxuening due to a suspected allergic reaction. There were no records of using anti-allergic drugs before and during the Shuxuening medication period. The exposure group eventually included 104 cases.

Nonexposed group
This group comprised patients who did not use any anti-allergic drugs after using Shuxuening injection and continued use of Shuxuening injection for >7 d. The average age of patients with allergy was within 5 years in the nonexposed group, and both groups had the same sex proportion. The ratio of exposure group patients to nonexposed patients was 1:4. The nonexposed group eventually included 416 cases.

Evidence-based research design
The studies that met the following inclusion criteria were included: 1) The study types were randomized controlled trial (RCT), case–control study, cohort study, and observational research. Language was limited to Chinese and English. 2) The subjects were patients who used Shuxuening injection and were not limited to age, sex, or disease types. 3) The test drug was Shuxuening injection; the medication method was intravenous infusion, or intravenous or intramuscular injection, as a single medication or in combination with other drugs. 4) Outcome indicators reported the incidence of adverse events/incident rates or provided statistics for adverse events/incident rates.

Exclusion criteria included the following: 1) interventions using other drugs that interfere with determination of the attribution of adverse reactions; 2) adverse reactions were not described; 3) incomplete data that could be analyzed; 4) short reports; 5) reviews; 6) animal models; and 7) basic medical research.

The RCTs, observational studies, and other studies were analyzed using the Cochrane System Evaluator Manual 5.1.0 offset risk assessment tool16 and MOOSE guidelines for quality assessment17 and included 7 criteria. Each criterion was counted as 1 point, and a total score ≥4 indicated high-quality literature. Rating criterion included the following: 1) clearly described hypothesis or purpose of the study; 2) whether the study object is explicitly included in the exclusion criteria; 3) whether the treatment program had a detailed description; 4) when a definitive diagnosis or definition of the outcome of the measurement was made; 5) whether the sample size was estimated in advance, or whether the patients were recruited consecutively; 6) whether the description of the main findings was clear, or whether the end-point evaluation was objective; 7) whether the statistical analysis was appropriate, or whether the outcome required stratified analysis and reporting, such as according to the disease stage, abnormal test results, and the characteristics of patients. Two reviewers independently extracted the data according to the inclusion and exclusion criteria and cross-checked the results for accuracy. When there was an inconsistency, the research team assisted in whether to include the study. The general information extracted included the author’s name, article title, publication date, type of study method, observation object, intervention measure, and quality evaluation score. Care was taken to avoid possible offsets and to perform sensitivity analysis if offsets could not be avoided.

Statistical analysis
Real-world statistical analysis
In this study, the main analytical methods used were the Chi-square test, the Wilcoxon test, and logistic regression analysis. SPSS18.0 software was used for statistical processing. \( P < 0.05 \) was the criterion for statistical significance.

Statistical analysis of evidence-based evaluation
Meta-analysis was performed using R3.2.3 software. In the meta-analyses, the CI for each effect was expressed as 95% CI. The evaluation of data heterogeneity was performed by a homogeneity test (\( Q \) test), and the test level was \( \alpha = 0.1 \). At the same time, the quantitative \( F \) was combined to determine the degree of heterogeneity (the judgment criterion was \( P \geq 0.1 \) or \( F \leq 50\% \)). When the heterogeneity was low, the fixed-effect model was used for analysis; in other cases, the random-effect model was used. Sensitivity analysis was used to investigate the stability of the results of the meta-analysis.

Data extraction analysis flowchart
The extraction and analysis process of the hospital case data are shown in Figure 1, and the extraction and analysis process of the literature data are shown in Figure 2.
A total of 7,208 patients who used Shuxuening injection were identified from HIS

Patients who were administered anti-allergic medication within 24 h of halting Shuxuening injection

Suspected class of allergic reactions
- allergic group n = 104
- control group n = 416

Gender, age
- 1:4 match

The time of using Shuxuening from start to finish > 7 d and did not use anti-allergic drugs during medication

Analysis of influencing factors of suspected allergic reaction

- Admission condition
- Allergic history
- Solvent type
- Single dose
- Combined medication

Figure 1 Flowchart of case assignments.
Abbreviation: HIS, hospital information system.

Retrieving relevant literature in database
(n = 3,819)

Remove the duplicate documents
(n = 1,729)

Reading the titles and abstracts
(n = 2,090)

Eliminate these nonqualifying literature citations
- (n = 963)
  - Studies on pharmacological action (n = 38)
  - Based on literature research (n = 74)
  - Unrelated literature (n = 582)
  - Animal experiment (n = 116)
  - Case report (n = 135)
  - System evaluation (n = 18)

The full text is qualified
(n = 1,127)

Eliminate these nonqualifying literature citations
- (n = 1,063)
  - No description of ADR (n = 759)
  - The research type is unclear (n = 98)
  - ADR analysis of TCM injection (n = 104)
  - Intervention did not match (n = 49)
  - The data reported is not complete (n = 53)

Final included literature citations
(n = 64)

Figure 2 Process and results of literature screening.
Abbreviations: ADR, adverse drug reaction; TCM, traditional Chinese medicine.
Ethical approval
Ethical approval was given by the Clinical Medicine Research Ethics Committee of the First Affiliated Hospital of Bengbu Medical College, and written informed consent was obtained from the patients. The information from the medical records of patients was confidential.

Real-world research results
General patient characteristics
Table 1 shows the sex and age differences of patients in the exposure group and the nonexposed group. The mean age of the exposure group was 68.76 ± 13.97 years after NCCS matching, and the mean age of the nonexposed group was 65.20 ± 14.87 years. Analysis using the Wilcoxon test showed no significant difference in age between the 2 groups (P = 0.229). The sex ratios in the exposure group and the nonexposed group were identical (1:4). There was no significant difference between the 2 groups in terms of sex as assessed by the Chi-square test (P = 0.861). The above results show that the 2 groups were well matched demographically, with comparable distribution of age and sex.

Analysis of the influencing factors in suspected allergic reactions
A previous study has shown that the influencing factors associated with allergic reactions were admission, single dose, and combination therapy. This study explored the impact of the above variables on suspected allergic reactions.

Admission conditions
Hospitalization analysis results are shown in Table 2. According to the patient case information, patient admissions were divided into general admissions and emergency admissions. The results show that there was no significant difference between the 2 groups (P > 0.05), and the admission conditions were not the factors in Shuxuening injection allergic reactions.

Disease type
The results of disease type analysis of the 2 groups are shown in Table 3. According to the patient case information and Shuxuening injection instructions, patient disease types were divided into 2 groups: indicated treatment and off-label treatment. The Chi-square test results showed that there was no significant difference between the 2 groups (P > 0.05). Therefore, allergic reactions were not influenced by whether Shuxuening treatments were administered for indicated diseases or off-label use.

Single dose
The Wilcoxon test and Chi-square test of single dose of the 2 groups are shown in Tables 4 and 5, respectively. The single dose of Shuxuening administered through intravenous infusion was 5–30 mL/d. The maximum dose recommended for the drug specification was 20 mL/d. We compared the single dose between the exposure group and the nonexposed group to determine the effect of the single dose on suspected allergic reactions. The Wilcoxon test results and the Chi-square test results showed that the difference between treatment groups was statistically significant (P < 0.05, the exposure group > the nonexposed group). Therefore, from a statistical point of view, it can be considered that a single-dose volume of >20 mL is an influencing factor in allergic reactions.

Combined medication
In this study, conditional logistic regression was used to model the effect of combined medication. For screened variables, the stepwise selection method was used. According to the analysis of the frequency of combined medications, we initially determined what types of drugs are used in conjunction with Shuxuening and how frequently were they associated with subsequent allergic reactions. The top 50 drugs were selected based on their frequency of use in combination therapy. Drugs with resulting values of P > 0.05 were omitted. The analysis showed that the risk of allergic reactions was small when Shuxuening injection was used in conjunction with aspirin, metoprolol, low-molecular-weight heparin calcium or sodium, or Danshen compound solution (Chinese medicine). However, there was an increased risk of allergic reactions when Shuxuening was used in conjunction with potassium aspartate, atorvastatin calcium tablets, Shengmai injection, or injection of pantoprazole sodium (OR values were >1, and 95% CIs were >1) (Table 6).

Table 1 The distribution of patient age and sex in two groups, n (%)

<table>
<thead>
<tr>
<th>Groups</th>
<th>n</th>
<th>Age (years)</th>
<th>Sex</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>≤40</td>
<td>41–60</td>
</tr>
<tr>
<td>Exposure group</td>
<td>104</td>
<td>4 (3.85)</td>
<td>16 (15.38)</td>
</tr>
<tr>
<td>Nonexposed group</td>
<td>416</td>
<td>20 (4.81)</td>
<td>108 (25.96)</td>
</tr>
</tbody>
</table>
Evidence-based evaluation results

Literature search results
A total of 3,819 related literature citations were retrieved from the database. There were 3,795 articles derived from Chinese databases, including 974 articles from VIP, 550 articles from Wanfang database, and 2,271 articles from CNKI. There were 24 articles cited in English, including 1 in the Cochrane Library and 23 in PubMed. Those articles were screened according to the inclusion and exclusion criteria, resulting in 64 articles being selected for the meta-analysis, including 56 articles of RCT, 6 articles of non-RCT, and 1 article of other observational studies.

Profile of included studies
The 64 articles contained information of 6,171 patients. Out of the 3,222 reported cases of Shuxuening injection use, adverse reactions were reported in 213 patients. Serious adverse reactions were reported in 5 journal articles, including 11 cases of serious adverse reactions and 7 cases of allergic shock. The ADR of Shuxuening injection was mainly associated with skin and associated tissues. The most common adverse reactions were face flushing, rash ecchymosis, and itching, accounting for 42.1% (85/202) of reported adverse reactions, followed by nervous system reactions, accounting for 29.2% (59/202). Of the 11 reported cases of severe adverse reactions, allergic shock accounted for 63.6% (7/11).

Meta-analysis results

Incidence of severe allergic reactions following Shuxuening injection
There were 5 articles that reported serious adverse reactions. The incidence of severe ADR was 0.34%, with an incidence of anaphylactic shock of 0.22%. Meta-analysis included

<table>
<thead>
<tr>
<th>Groups</th>
<th>Number of cases (%)</th>
<th>Number of cases (%)</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Exposure group (104)</td>
<td>28 (26.92)</td>
<td>76 (73.08)</td>
<td>1.598</td>
</tr>
<tr>
<td>Nonexposed group (416)</td>
<td>88 (21.15)</td>
<td>328 (78.85)</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Groups</th>
<th>Indicated (%)</th>
<th>Off-label (%)</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Exposure group (n=104)</td>
<td>101 (97.12)</td>
<td>3 (2.88)</td>
<td>0.325</td>
</tr>
<tr>
<td>Nonexposed group (n=416)</td>
<td>399 (95.91)</td>
<td>17 (4.09)</td>
<td></td>
</tr>
</tbody>
</table>

Table 2 Analysis of hospitalization

<table>
<thead>
<tr>
<th>Groups</th>
<th>Emergency (%)</th>
<th>General (%)</th>
<th>χ²</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Exposure group (104)</td>
<td>17 (4.09)</td>
<td>77 (95.91)</td>
<td>0.056</td>
<td>0.813</td>
</tr>
<tr>
<td>Nonexposed group (416)</td>
<td>399 (95.91)</td>
<td>1 (0.22)</td>
<td>0.005</td>
<td>0.944</td>
</tr>
</tbody>
</table>

Discussion

Methodology
In recent years, there are increasing numbers of real-world studies (RWSs) to evaluate post-market drug efficacy and safety. These RWSs were based on a large sampling of recorded clinical treatment data from the HIS and a complete record of the drug information from patients during hospitalization. In the application of traditional Chinese medicine, safety reevaluation has broad prospects. Combined PSA with NCCS was used to analyze the screened suspected allergic reaction cases caused by Shuxuening injection. The PSA method was based upon the existing, complete prescription record database to achieve, in line with clinical practice, good external authenticity. When using certain drugs that are indicated for treatment of ADR of other drugs, the requirements of the PSA method are met. Although there is no record of allergic reactions in the HIS data, the use of anti-allergic indications is consistent with PSA conditions, indicating the patient has experienced an allergic reaction. The advantage of using NCCS method is that we can collect data first, save manpower and material resources, and decrease survey selection bias. There is a good balance and comparability between the allergic group and control group.
resulting from matching of interference from confounding factors. This research method is economical and allows accurate extrapolation.²⁵

In this study, HIS data were used to explore the influencing factors of adverse reactions, and by using the collected data, we were able to avoid the common recall bias of the classic case–control study.²⁶ The selected controls were randomly selected from the same cohort obtained from patients who used Shuxuening; the data were highly comparable; and to a certain extent, selection bias can be reduced when the effect is estimated.

Most of the studies on evidence-based medicine for traditional Chinese medicine injections are based on clinical RCTs. It is well known that RCT studies have many advantages: they can effectively prevent selection bias, they are highly comparable, and the results exhibit high authenticity and an increased level of confidence.²⁷,²⁸ Meta-analysis of widely searched RCT research data can result in a more objective evaluation of the evidence, and a more accurate and objective evaluation of the effect indicators, which improves study credibility.

Therefore, the innovation of this study was the combined analysis of database literature for the incidence of ADR, with the analysis based on RWSs to understand the factors influencing suspected adverse reactions of Shuxuening. Such an approach can reveal much regarding the factors that influence the adverse reactions of Shuxuening, the incidence of serious adverse reactions, and the occurrence of rare adverse reactions, and may provide a way for further research and ideas. Based on the literature database analyses, this study should be helpful to understand the adverse events related to Shuxuening injection, and when combined with the HIS data analysis, we can grasp the occurrence of clinical adverse reactions. These 2 methods complement each other and allow for a comprehensive comparison. Furthermore, this approach allows for a more in-depth and innovative investigation of the safety of clinical applications of Shuxuening. Lastly, our approach resulted in a more reliable evidence chain to provide references for clinical use of Shuxuening. A search of literature databases found no similar literature published.

### Risk factors

There have been several reports related to the influencing factors of adverse reactions following Shuxuening injection,²⁶,²⁹–³¹ and the influencing factors were reported in combination, solvent type, high-dose medication, and type of disease treatment. The results of this study show that there was a statistically significant correlation between single dose and adverse events. There is also a greater risk of allergic reactions when Shuxuening was combined with potassium magnesium aspartate, atorvastatin calcium, Shengmai injection, and pantoprazole sodium. It has been reported that there is a contraindication between Shuxuening injection and pantoprazole sodium, anti-infective drugs, traditional Chinese medicine injections, and other drugs.³² The interaction of drugs in the body may lead to adverse reactions and even occurrence of allergic reactions. Thus, care should be taken when combining Shuxuening with drugs, and an appropriate time interval should be maintained between administration of drug combinations.

Due to the inclusion of a large single-center study, there were only a few instances where 0.9% sodium chloride solution was used as a vehicle. Thus, we were unable to perform real-world research analysis on the vehicle. Meta-analysis of evidence-based medicine showed that the vehicle used was a possible factor for adverse events along with high dose and type of treatment (indicated vs off-label): The incidence of adverse reactions with 0.9% sodium chloride solution (5.60%) was higher than that with 5% glucose solution (2.62%), and the difference was significant (P < 0.05). Therefore, the type of vehicle is a suspected factor in Shuxuening adverse reactions. Thus, these results suggest that 5% glucose is best used as a vehicle for Shuxuening. In addition, the incidence of ADR increased significantly (P < 0.05) when the dose was >20 mL/d (10.15%) when compared to doses <20 mL/d (5.57%). Therefore, doses >20 mL/d may be a key factor

### Table 4 The Wilcoxon test results for single dose

<table>
<thead>
<tr>
<th>Groups</th>
<th>Average (mL)</th>
<th>SD</th>
<th>Median (mL)</th>
<th>Minimum (mL)</th>
<th>Maximum (mL)</th>
<th>Wilcoxon test (P-value)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Exposure group (n=104)</td>
<td>20.38</td>
<td>1.93</td>
<td>20</td>
<td>20</td>
<td>30</td>
<td>0.015</td>
</tr>
<tr>
<td>Nonexposed group (n=416)</td>
<td>19.51</td>
<td>2.83</td>
<td>20</td>
<td>5</td>
<td>30</td>
<td></td>
</tr>
</tbody>
</table>

### Table 5 The Chi-square test results for single dose

<table>
<thead>
<tr>
<th>Groups</th>
<th>≤20 mL/d</th>
<th>&gt;20 mL/d</th>
<th>χ²</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of cases (%)</td>
<td>Number of cases (%)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Exposure group (n=104)</td>
<td>96 (92.31)</td>
<td>8 (7.69)</td>
<td>22.941</td>
<td>0.000</td>
</tr>
<tr>
<td>Nonexposed group (n=416)</td>
<td>412 (99.04)</td>
<td>2 (0.48)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
leading to the occurrence of Shuxuening ADR, which is consistent with a previous study.33

Through the analysis of HIS data, the Chi-square test showed that off-label use of Shuxuening is not an influencing factor in hypersensitivity induced by Shuxuening injection. However, the results of the meta-analysis showed that the incidence of ADR with the off-label use of Shuxuening was greater than the incidence of adverse reactions when treatment with Shuxuening was indicated. This may be due to the data sources and necessitates further study.

Incidence of allergic reactions and severe anaphylactic shock

Following review of the existing literature, we found cases of severe adverse reactions following Shuxuening injection.34–36 Symptoms of severe adverse events following Shuxuening injection were usually rapid and severe, including anaphylactic shock, which is life threatening. Patient medical records in the HIS showed that the incidence of suspected allergy-related adverse events was 1.44%. Meta-analysis of evidence-based medicine studies showed that the incidence of severe ADR was 0.34%, and the incidence of anaphylactic shock was 0.22%. The incidence of severe adverse events and the incidence of anaphylactic shock are rarely reported in other studies regarding the safety of Shuxuening injection. Some research reports stated that as a traditional Chinese medicine injection, Shuxuening has a high incidence of serious adverse reactions.26,37,38 If serious adverse reactions were detected early, however, the patient’s symptoms usually disappear following timely administration of a series of anti-allergic emergency treatments.39 To reduce the incidence of serious adverse reactions, a thorough patient history regarding allergies should be obtained by the clinician. In addition, because the drug contains a small amount of ethanol, it

Table 6 Screening table of suspected risk factors

<table>
<thead>
<tr>
<th>Medicine</th>
<th>Regression coefficient</th>
<th>SE</th>
<th>P-value</th>
<th>OR</th>
<th>95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Potassium aspartate and magnesium</td>
<td>0.922</td>
<td>0.608</td>
<td>0.002</td>
<td>2.513</td>
<td>1.386; 4.559</td>
</tr>
<tr>
<td>Atorvastatin calcium tablets</td>
<td>1.361</td>
<td>0.354</td>
<td>0.033</td>
<td>1.042</td>
<td>0.669; 1.621</td>
</tr>
<tr>
<td>Shengmai injection</td>
<td>1.088</td>
<td>0.687</td>
<td>0.001</td>
<td>2.970</td>
<td>1.514; 5.824</td>
</tr>
<tr>
<td>Pantoprazole sodium</td>
<td>0.456</td>
<td>0.449</td>
<td>0.041</td>
<td>1.578</td>
<td>1.016; 2.451</td>
</tr>
<tr>
<td>Low-molecular-weight heparin</td>
<td>0.092</td>
<td>0.477</td>
<td>0.700</td>
<td>1.096</td>
<td>0.687; 1.749</td>
</tr>
<tr>
<td>Salvia miltiorrhiza</td>
<td>0.426</td>
<td>0.575</td>
<td>0.136</td>
<td>1.531</td>
<td>0.872; 2.688</td>
</tr>
<tr>
<td>Aspirin</td>
<td>–0.610</td>
<td>0.509</td>
<td>0.011</td>
<td>0.530</td>
<td>0.322; 0.871</td>
</tr>
<tr>
<td>Metoprolol tartrate tablets</td>
<td>–0.778</td>
<td>0.658</td>
<td>0.016</td>
<td>0.459</td>
<td>0.241; 0.875</td>
</tr>
</tbody>
</table>

Note: OR is used to measure the size of the risk factor; the smaller the OR value, the smaller the risk.

Table 7 Meta-analysis results of total incidence rate and different influencing factors of ADRs

<table>
<thead>
<tr>
<th>Analysis type</th>
<th>Studies</th>
<th>Total sample size</th>
<th>Number of ADR cases</th>
<th>Heterogeneity test</th>
<th>Effect models</th>
<th>Meta-analysis results</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>Total incidence</td>
<td>64</td>
<td>3,222</td>
<td>213</td>
<td>P² = 36.8%, P = 0.0022</td>
<td>Fixed</td>
<td>0.0584</td>
</tr>
<tr>
<td>Incidence of general ADR</td>
<td>61</td>
<td>3,088</td>
<td>202</td>
<td>P² = 36.5%, P = 0.003</td>
<td>Fixed</td>
<td>0.0577</td>
</tr>
<tr>
<td>Incidence of severe ADR</td>
<td>5</td>
<td>236</td>
<td>11</td>
<td>P² = 0%, P = 0.5952</td>
<td>Fixed</td>
<td>0.0436</td>
</tr>
<tr>
<td>5%–10% GS</td>
<td>22</td>
<td>1,198</td>
<td>71</td>
<td>P² = 10.6%, P = 0.3211</td>
<td>Fixed</td>
<td>0.0517</td>
</tr>
<tr>
<td>0.9% NS</td>
<td>16</td>
<td>725</td>
<td>45</td>
<td>P² = 8.7%, P = 0.3545</td>
<td>Fixed</td>
<td>0.0560</td>
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<tr>
<td>Treatment time ≤7 d</td>
<td>6</td>
<td>398</td>
<td>23</td>
<td>P² = 42.7%, P = 0.1203</td>
<td>Fixed</td>
<td>0.0514</td>
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<tr>
<td>7 d &lt; treatment time ≤14 d</td>
<td>40</td>
<td>1,913</td>
<td>117</td>
<td>P² = 18.3%, P = 0.1588</td>
<td>Fixed</td>
<td>0.0549</td>
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<tr>
<td>Treatment time &gt;14 d</td>
<td>17</td>
<td>811</td>
<td>71</td>
<td>P² = 52.2%, P = 0.0064</td>
<td>Fixed</td>
<td>0.0700</td>
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<tr>
<td>Single medication</td>
<td>9</td>
<td>448</td>
<td>28</td>
<td>P² = 0%, P = 0.7465</td>
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<td>0.0569</td>
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<tr>
<td>Combined medication</td>
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<td>2,481</td>
<td>166</td>
<td>P² = 44.3%, P = 0.0005</td>
<td>Fixed</td>
<td>0.0616</td>
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<tr>
<td>Dose ≥20 mL/d</td>
<td>56</td>
<td>2,806</td>
<td>176</td>
<td>P² = 28.7%, P = 0.0261</td>
<td>Fixed</td>
<td>0.0557</td>
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<tr>
<td>Dose &gt;20 mL/d</td>
<td>5</td>
<td>202</td>
<td>22</td>
<td>P² = 48.0%, P = 0.1037</td>
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<td>0.1015</td>
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<tr>
<td>Indications</td>
<td>57</td>
<td>2,930</td>
<td>189</td>
<td>P² = 39.3%, P = 0.0016</td>
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<td>0.0566</td>
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<tr>
<td>Super indications</td>
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<td>292</td>
<td>24</td>
<td>P² = 0%, P = 0.4729</td>
<td>Fixed</td>
<td>0.0775</td>
</tr>
</tbody>
</table>

Abbreviations: ADR, adverse drug reaction; GS, glucose solution; NS, sodium chloride solution.
should be used with caution in patients intolerant to alcohol. During treatment, vigilance for adverse reactions must be maintained, and once these occur, treatment should be halted and appropriate interventions should be applied.

**Conclusion**

RWSs combined with meta-analysis showed that the suspected influencing factors of adverse reactions have a relationship with the type of solvent, the dosage, off-label use, applications, and combination therapy. There was no clear relationship between the condition of the patient upon admission and suspected hypersensitivity reactions to Shuxuening. Other risk factors for suspected hypersensitivity reactions include combination with potassium aspartate, atorvastatin calcium tablets, Shengmai injection, pantoprazole sodium, and other drugs. Thus, clinicians should follow the instructions regarding dosage, be careful when combining Shuxuening with other medication, and resist the use of Shuxuening injection for nonindicated purposes. In this study, RWSs extracted single-center sample data that are not representative of the overall clinical use of Shuxuening injection, and therefore, a larger sample of data is required to verify the results of this study.

In conclusion, our study combined real-world data with analysis of evidence-based literature. We have shown here that this approach provides for a more scientific and intuitive reference for evaluating the clinical application of Shuxuening injection and the control of adverse reactions. We propose that this novel approach for reevaluation of the post-marketing safety of Shuxuening may be extended to the reevaluation of the safety of other traditional Chinese medicines.

**References**


