THE CLINICAL EFFICACY AND SAFETY OF SOMATOSTATIN AND THROMBIN COMBINED WITH OMEPRAZOLE FOR THE TREATMENT OF ACUTE UPPER GASTROINTESTINAL BLEEDING

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Abstract

The present study aimed to investigate the clinical efficacy and safety of somatostatin and thrombin combined with omeprazole for the treatment of 64 patients with acute upper gastrointestinal bleeding. 128 patients with acute upper gastrointestinal bleeding in our hospital between January 2015 and May 2017 were enrolled. Patients were randomly divided into two groups according to a random number table: observation group and control group (n = 64, each). The control group received omeprazole; the observation group received somatostatin and thrombin. The total effective rate of the observation group was 93.75%, significantly higher than that of the control group (78.12%, p < 0.05); the time for haemostasis and hospitalisation in the observation group were significantly shorter than in the control group (p < 0.05); the haemoglobin (HGB) after treatment in the observation group significantly increased compared with the control group (p < 0.05); the adverse reaction rates in the observation group was 10.94%; the difference was not statistically significant when compared with the control group (7.81%, p > 0.05). Somatostatin and thrombin combined with omeprazole for treatment of acute upper gastrointestinal bleeding has a significant therapeutic effect and less acute adverse reactions. It is a safe and effective treatment option, and it may be used in clinical practice.

Keywords: acute upper gastrointestinal bleeding, omeprazole, somatostatin, thrombin, clinical efficacy

Introduction

Acute upper gastrointestinal bleeding (UGIB) refers to the bleeding caused by gastrointestinal lesions above the Treitz ligament. The common diseases causing it are: bleeding caused by the rupture and varices of oesophageal veins, emergency ulcers, gastric and duodenal ulcers, gastric mucosal lesions and biliary bleeding [1]. Bleeding caused by jejunal lesions after gastrojejunoostomy also belongs to this scope [2]. The onset of UGTB shows an aging trend [3]. The risk of death of UGB increases six times in patients over 75 years of age [4, 5]. UGB is characterized by sudden onset, rapid change of conditions and high fatality rate [6]. Patients with unstable haemodynamic have higher mortality rates [7, 8]. A study reported that if acute massive haemorrhage occurred, the mortality rate was approximately 10% [9]. Therefore, rapid and effective control of bleeding is the key measure to save UGB patients’ lives. Large sample analysis of UGB clinical data is of great value for improving national health [10]. Omeprazole is a commonly used drug in the treatment of UGB in clinical applications. Its efficacy has been confirmed by many clinical trials, but there are still some patients with unsatisfactory treatment results. Our study aimed to assess the therapeutic potential of omeprazole combined with somatostatin and thrombin in acute UGB.

Materials and Methods

Study design

A total of 128 patients with acute UGB admitted to the Department of Emergency Internal Medicine, Luhe...
Patients were male and 23% sodium-

Electrocardiogram within 24-

Methods

Inclusion and exclusion criteria

Inclusion criteria: (1) Patients with symptoms of UGIB such as melena, hematemesis and abdominal pain within 12 hours and stable vital signs; (2) no haemostatic drugs were used within 24 hours before admission, no drug to inhibit gastric acid secretion was used in the preceding week; (3) the age ranged between 18-80 years old; (4) patients or their family members were voluntary enrolled; all patients or their family members have provided an informed consent. This study was approved by the Ethics Committee of our hospital.

Exclusion criteria: (1) Patients with a history of subtotal gastrectomy and patients with digestive tract tumours; (2) patients with severe complications such as pyloric obstruction or massive haemorrhage; (3) patients with severe heart, liver, lung, kidney and other organ dysfunctions; (4) patients with cerebrovascular diseases, severe trauma, respiratory bleeding and haematological diseases; (5) patients with melena caused by drugs or food and bleeding from the mouth, nose, throat and respiratory tract; (6) patients with incomplete data so that curative effect statistics could not be carried out.

Methods

After admission, patients in both groups received basic treatment to maintain the stability of vital signs, including symptomatic treatment such as vital signs monitoring, fasting and fluid deprivation, maintenance of airway patency and internal environment stability, active blood volume supplementation, adjustment of water and electrolyte disorders. A low-temperature liquid or semi-liquid diet was given to patients 24 h after hematemesis stopped. Patients in the control group were intravenously dripped with 40 mg of omeprazole + 100 mL of saline twice a day for five days. On the basis of the treatment in the control group, patients in the observation group continued to be intravenously pumped with 3.0 mg of somatostatin, which was diluted to 500 mL with 0.9% sodium chloride solution at a dosage rate of 40 mL/hour for five days and were intra-gastrically injected with 500 IU of thrombin, which was diluted with 20 mL of 0.9% sodium chloride solution three times a day, until hematemesis was stopped. During the treatment, no other drugs were used in both groups. No strong tea, coffee, spicy food, smoking and alcohol were allowed.

Before and after treatment, the changes in clinical symptoms and vital signs, respiration, heart rate, blood pressure and continuous electrocardiogram monitoring indexes were observed. The changes in urine volume, haematochezia, hematemesis and haemoglobin (HGB) levels were observed. Improvement of gastrointestinal lesions was observed by gastroscopy. Liver and kidney function examination, routine blood and urine tests and electrocardiogram were performed before and after treatment, and the determination of whether or not adverse reactions occurred in patients was observed and asked.

Curative effect standard

Criteria for bleeding stopping: (1) heart rate ranged within 60 - 100 beats/min; (2) blood pressure was > 90/60 mmHg (1 mmHg = 0.133 kPa); (3) there were no symptoms of melena or hematemesis; and (4) there was no decrease in haematocrit and HGB level.

Clinical curative effect. Excellent: within 24 - 48 hours after treatment, no active bleeding occurred, haematochezia or hematemesis stopped, gastric tube drainage fluid was clear and bloodless, the pulse and blood pressure were stable, and HGB was stable or increased. Effective: within 48 - 72 hours after treatment, bleeding was stopped, gastric tube drainage fluid was bloodless, blood pressure and heart rate were stable, HGB did not decrease. Ineffective: after 72 h of treatment, there were still melena and hematemesis, blood pressure and heart rate were unstable, the gastric tube drainage fluid was still bright red or dark red, the blood volume was continuously decreasing. The total effective rate was expressed as (excellent cases + effective cases)/total number of cases × 100%.

Statistical analysis

Data were statistically analysed using statistical software SPSS18.0. Measurement data were expressed as mean ± standard deviation (x ± SD) and compared using t-test. Count data were expressed as percentage (%) and compared using χ² test. p < 0.05 was considered statistically significant.
Results and Discussion

Comparison of clinical curative effect between two groups of patients

The excellent rate and total effective rate were significantly higher in the observation group than in the control group; the differences were statistically significant (p < 0.05, Table I).

Comparison of haemostatic time and hospitalization time between the two groups

The mean haemostatic time and hospitalization time were significantly shorter in the observation group than in the control group; the differences were statistically significant (p < 0.05, Table II).

Table I

Comparison of clinical curative effect between the two groups of patients

| Index                        | Observation group | Control group | p value |
|------------------------------|-------------------|---------------|---------|
| N                            | 64                | 64            | > 0.05  |
| Male/Female                  | 41/23             | 38/26         | > 0.05  |
| Age (Years)                  | 43.55 ± 8.12      | 44.86 ± 8.75  | > 0.05  |
| Cause of bleeding            |                   |               | > 0.05  |
| Gastric ulcer, duodenal ulcer| 31 (48.44%)       | 32 (50.00%)   |         |
| Duodenal ulcer               | 17 (26.56%)       | 16 (25.00%)   |         |
| Hemorrhagic gastritis        | 8 (12.50%)        | 9 (14.06%)    |         |
| Esophageal and gastric varices| 8 (12.50%)        | 7 (10.94%)    |         |
| Clinical conditions          |                   |               | > 0.05  |
| Black stool only             | 31 (59.38%)       | 35 (54.69%)   |         |
| Black stool with hematemesis | 27 (42.19%)       | 29 (45.31%)   |         |
| Excellent rate (n(%))        | 45 (70.31%)       | 21 (32.81%)   | < 0.05  |
| Effective rate (n(%))        | 15 (23.44%)       | 29 (45.31%)   | < 0.05  |
| Invalid rate (n(%))          | 4 (6.25%)         | 14 (21.88%)   | < 0.05  |
| Total effective rate (n(%))  | 60 (93.75%)       | 50 (78.12%)   | < 0.05  |

Table II

Comparison of haemostatic time and hospitalization time between the two groups

| Groups            | Cases (n) | Haemostatic time (h) | Hospitalization time (d) |
|-------------------|-----------|----------------------|--------------------------|
| Observation group | 64        | 18.11 ± 4.48         | 3.41 ± 2.12              |
| Control group     | 64        | 32.65 ± 7.24         | 5.68 ± 2.39              |
| \( \chi^2 \)      | 13.662    | 5.684                |                          |
| p value           | < 0.05    | < 0.05               |                          |

Comparison of HGB between two groups before and after treatment

The difference in the level of HGB between the two groups before treatment was not statistically significant (p > 0.05). After five days of treatment, HGB levels in both groups were significantly higher than those before treatment, which was significantly higher in the observation group than in the control group; the difference between the two groups was statistically significant (p < 0.05, Table III).

Table III

Comparison of HGB levels between the two groups before and after treatment

| Groups            | Cases (n) | Before treatment | After treatment |
|-------------------|-----------|------------------|-----------------|
| Observation group | 64        | 92.35 ± 15.44    | 115.23 ± 11.34  |
| Control group     | 64        | 93.45 ± 11.89    | 103.72 ± 13.56  |
| \( \chi^2 \)      | 0.452     | 5.209            |                 |
| p Value           | > 0.05    | < 0.05           |                 |

Comparison of adverse reactions between two groups

During treatment, in the observation group, palpitation occurred in two patients, short-term vertigo occurred in two patients and nausea occurred in three patients; the adverse reaction rate was 10.94% (7/64). During treatment, in the control group, palpitation occurred in one patient, short-term vertigo occurred in two patients and nausea occurred in two patients; the adverse reaction rate was 7.81% (5/64). No abnormalities were found in liver and kidney function tests and routine blood and urine tests in both groups. The above-mentioned adverse reactions were all alleviated after symptomatic treatments. The difference in the adverse reaction rate between groups was not statistically significant (\( \chi^2 = 0.267, \ p > 0.05 \)) (Table IV).
Acute UGIB is a common clinical critical illness. The main clinical symptoms are sudden massive hematemesis and melena, accompanied by decreased erythrocyte haematocrit, haemoglobin and blood pressure in varying degrees. Acute UGIB is a critical condition and entails more complications. It extremely easily causes liver and kidney failure and even haemorrhagic shock; if haemostasis is not performed timely and effectively, it will threaten patient's life and safety [2]. Therefore, once the symptoms of UGIB are found, quick and effective haemostasis and rescue measures should be given, and bleeding should be controlled with drugs as soon as possible after this starts, in order to prevent haemorrhagic shock, to find out the causes of bleeding in time and directly create favourable conditions for further treatment. At present, there are many drugs in the treatment of UGIB in clinic, but the curative effects are diverse. It is necessary to find a quick and effective treatment option or drug for haemostasis.

The causes of UGIB are more complicated; digestive system ulcer, acute gastric mucosal injury, oesophageal rupture and gastric varices rupture caused by liver cirrhosis are the most common in clinical practice. Gastric venous rupture usually leads to more severe bleeding and a higher mortality rate than oesophageal varices [11]. The mortality rate of first bleeding of oesophageal and gastric varices bleeding (EGVB) is up to 25 - 50% [12, 13]. Studies revealed that peptic ulcer was the most common cause of UGIB, followed by oesophageal varices and acute gastric mucosal lesions [14]. A large number of previous studies revealed that duodenal ulcer was the most common aetiology of peptic ulcer [15], especially in the case of excessive gastric acid secretion, where it was more likely to occur. Pepsinogen can be activated by gastric acid to produce peptisinase in acidic environment, to rapidly digest blood clots. Additionally, platelet aggregation is also affected by acidic conditions; thereby, the haemostasis function of digestive system is affected [4]. Therefore, in the clinical therapeutic process, acid-making agent is often used to increase the pH value in stomach and in combination with haemostatic drugs to achieve the purpose of haemostasis. Increasing the pH value of gastric secretion is conducive to reduce coagulation and lysis under the influence of pepsin, shorten coagulation time, promote platelet aggregation, delay the dissolution of clots and promote haemostasis [5]. A recent study revealed that intravenous drip of omeprazole could significantly reduce the 24-hour gastric acidity, enhance the haemostatic effect and reduce the mortality rate [16]. Omeprazole is a commonly used proton pump inhibitor in clinical practice. It can selectively act on the cell membrane and inhibit the activity of H⁺-K⁺-ATPase in order to produce strong acid-inhibiting effect, thus increasing the pH value in the stomach. In an alkaline environment, it promotes blood coagulation, thus, fast and effective haemostasis [6]. A clinical trial revealed that omeprazole could rapidly and effectively reduce the nocturnal and 24-hour gastric acidity of patients, maintain the pH value around 3 - 4, thereby improving the symptoms of gastric acid reflux of the oesophagus [17]. Omeprazole can strongly inhibit gastric acid secretion, so it is widely used in the treatment of gastroesophageal reflux and peptic ulcer. Omeprazole can inhibit gastric acid approximately 20 times stronger than an H2 receptor antagonist and has no obvious side effects and is safe and reliable [8]. In addition, omeprazole can produce a negative feedback effect and induce hypergastrinaemia, in turn increasing the blood flow of gastric mucosa, then promoting the repair of ulcer and erosion wounds and promoting the formation of blood scab, to exert a haemostatic effect. Therefore, using omeprazole in the treatment of acute UGIB can effectively inhibit gastric acid secretion, protect the bleeding surface from irritation of gastric acid, and can also trigger the automatic haemostasis mechanism of the body, effectively promoting platelet aggregation and inhibiting the digestion after its aggregation and promoting blood agglutination, thereby exerting a haemostatic effect [9]. Somatostatin is a synthetic cyclic 14-amino acid peptide, having the same structure and physiological effects as natural somatostatin, which can effectively inhibit the production and release of a series of vasodilator peptides such as vasoactive peptide [10]. In addition, it can selectively act on visceral vessels to promote vasoconstriction, reduce visceral blood flow, in turn reducing the symptoms of portal hypertension. However, it has no significant effect on systemic haemodynamics. Somatostatin can inhibit the secretion and release of pepsin and peptic acid,
effectively protecting gastric mucosa and epithelial cells, thus reducing the haemolysis effect of pepsin and peptic acid on blood clots, exerting a synergistic haemostasis effect [11]. Hutchinson et al. revealed that [18] the use of somatostatin can increase the pressure of lower oesophageal sphincter, promotes the effective contraction of the lower oesophageal venous plexus, in turn reducing the blood flow in the varices vein of the oesophagus, and promoting the contraction of blood clots and the agglutination of platelets, which has significant effects on UGIB caused by many factors. Thrombin is an enzymatic haemostatic, is mainly extracted from the venom of the Brazilian spearhead snake, separated and purified. It does not contain toxic components such as neurotoxins. In normal blood vessels, thrombin has no thrombokinase-like effect and plays only its thrombin-like role. When vascular skin lesions occur, both thrombokinase-like and thrombin-like effects can be exerted [13]. Thrombin can stimulate endogenous and exogenous coagulation function after application. Thrombin can directly act on fibrinogen in the blood and promote the conversion of fibrinogen into fibrin, thereby accelerating the coagulation of blood, thus having the effect of haemostasis and coagulation [19, 21]. Different from other haemostatics, thrombin is highly selective and targeted; it can stop bleeding at the site of bleeding, cannot activate the stabilizing factor of fibrin, and cannot affect the platelet aggregation and digestion in other sites of the blood vessel. Therefore, its haemostatic effect is fast and effective [14, 20]. When it reaches the damaged site of the blood vessel, it can quickly produce small thrombus that fills the damaged site, and block the bleeding process. Van Geffen et al. revealed that [22] thrombin can promote mitosis of epithelial cells in order to facilitate wound healing, thereby exerting a haemostatic effect. The present study revealed that the combination of these three drugs could exert a haemostatic effect through different haemostatic mechanisms. The results of the present study revealed that in patients in the observation group who were treated with somatostatin and thrombin combined with omeprazole, the total effective rate was 93.75%, which increased significantly when compared with the control group (78.12%), with the mean haemostatic time and hospitalization time significantly shorter in the observation group than in the control group. The investigators consider that the combination of three drugs can improve the targeting of haemostasis, thereby stopping bleeding more quickly and improving the haemostatic effect. After treatment, HGB levels in both groups were significantly higher than those before treatment, but the level increased more significantly in the observation group than in the control group. Our results suggest that the combination of drugs is helpful to enhance the haemostatic effect. Adverse reaction monitoring revealed that the adverse reaction rate in the observation group was 10.94%, which was not significantly higher when compared with the control group (7.81%). The adverse reaction rate was slightly higher in the observation group as compared to the control group, because the patients' compliance to the doctor's advice was poor and the body position was often changed in the observation group due to the rapid recovery of the patients. In addition, the stimulation of the gastric mucosa by 20 mL saline was another reason. The adverse reactions of the patients were significantly improved after the administration of metoclopramide. Our results revealed that the combination of three drugs does not increase the incidence of adverse reactions.

Conclusions

Somatostatin and thrombin combined with omeprazole in the treatment of acute UGIB can shorten the haemostatic time and the hospitalization, quickly and effectively stop bleeding, and has few adverse reactions, thereby making it a safe and reliable.

Conflict of interest

The authors declare no conflict of interest.

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