Plasma D-dimer Can Effectively Predict the Prospective Occurrence of Ascites in Advanced Schistosomiasis Japonica Patients

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Abstract: China still has more than 30,000 patients of advanced schistosomiasis while new cases being reported consistently. D-dimer is a fibrin degradation product. As ascites being the dominating symptom in advanced schistosomiasis, the present study aimed to explore a prediction model of ascites with D-dimer and other clinical easy-achievable indicators. A case-control study nested in a prospective cohort was conducted in schistosomiasis-endemic area of southern China. A total of 291 patients of advanced schistosomiasis were first investigated in 2013 and further followed in 2014. Information on clinical history, physical examination, and abdominal ultrasonography, including the symptom of ascites was repeatedly collected. Result showed 44 patients having ascites. Most of the patients’ ascites were confined in the kidney area with median area of 20 mm². The level of plasma D-dimer and pertinent liver function indicators were measured at the initial investigation in 2013. Compared with those without ascites, cases with ascites had significantly higher levels of D-dimer (0.71±0.24 μg/L vs 0.48±0.12 μg /L, P = 0.006), as well ALB (44.5 vs 46.2, g/L) and Type IV collagen (50.04 vs 44.50 μg/L). Receiver operating characteristic curve analyses indicated a moderate predictive value of D-dimer by its own area under curve (AUC) of 0.64 (95% CI: 0.54-0.73) and the cutoff value as 0.81 μg/L. Dichotomized by the cutoff level, D-dimer along with other categorical variables generated a prediction model with AUC of 0.76 (95% CI: 0.68-0.89). Risks of patients with specific characteristics in the prediction model were summarized. Our study suggests that the plasma D-dimer level is a reliable predictor for incident ascites in advanced schistosomiasis japonica patients.

Key words: Schistosoma japonicum, D-dimer, ascites, prediction, schistosomiasis japonica, China

INTRODUCTION

Schistosomiasis japonica, caused by the infection of Schistosoma japonicum, is a widely distributed parasitic zoonoses that threatens human’s health and social economic development [1,2]. The epidemic situation of the disease used to be serious in China. Comprehensive schistosomiasis control strategies in the past decades decreased a concrete number of infected cases from 1.52 million in 1989 to 0.12 million in 2013, and most of the endemic areas are controlled within the status of transmission interruption [2]. However, new challenges emerged that the remaining disease transmission in the geographically complex areas turns to be more challenging and that more infected patients who were not treated thoroughly are now developing into the late stage, i.e., advanced schistosomiasis [3-5]. Meanwhile, a few patients who received the standard clinical treatment timely still keep on the development to advanced schistosomiasis [3,5-8]. According to a 2014 national wide report, 115,614 cases of S. japonicum infection were estimated, and 30,880 patients are suffering from advanced schistosomiasis. Advanced schistosomiasis is characterized by hepatosplenic conditions, such as periportal liver fibrosis, spleen enlargement and congestion, portal hypertension, and other serious sequelae. The late stage schistosomiasis is hitherto difficult, if not impossible, to be fully recovered. Thus, with an increasing proportion of advanced cases in the overall disease spectrum, researchers should provide effective prevention strategies with early detection, early diagnosis, and early treatment.
Ascites is a common symptom in the late stage of schistosomiasis, which is caused by venous obstruction and portal hypertension with accumulation of granulomatous inflammation and fibrosis [3,9]. More frequency of ascites in the advanced schistosomiasis patients were reported from different endemic areas with prevalence of ascites as 60-90% in advanced cases [10,11]. In advanced schistosomiasis cases, the interaction of reinfection and failure in fibrosis treatment destroys the patient’s liver and portal vein system, causing the symptoms of ascites, and eventually depletes patient’s health and quality of life [12]. Therefore, it is important to pre-diagnose and provide prevention and treatment to those advanced patients who have great risk of ascites. However, few studies have done this research.

The pathophysiological facts of vascular injury and fibrosis underlying the development of advanced schistosomiasis may be capable to predict the risk of ascites occurrence. D-dimer, one of the degradation products of cross-link fibrin, has been validated as an accurate marker of vascular injury and fibrinolytic activity [13-15]. Studies have confirmed the potential sensitivity and specificity of D-dimer in early diagnosis of pulmonary embolism, lung cancer, and esophageal varix bleeding [16-18]. It was found regarding the mechanisms of advanced schistosomiasis that peripheral portal vasculature destructions in liver fibrosis increases the intrahepatic portal stress and causes ascites [19]. Thus, it is logical and plausible to employ D-dimer as a sensitive marker in the prediction of ascites. In the present study, we aimed to explore the predictive value of D-dimer in the incident ascites in patients of advanced schistosomiasis during 1-year follow up.

MATERIALS AND METHODS

Participants

A case-control study nested in a prospective cohort was conducted within 4 fields of Xinjian, Nanchang, Duchang, and Yugan Counties in Jiangxi Province, P. R. China, all of which are serious endemic areas of schistosomiasis japonica. Using stratified multistage sampling method, a total of 291 advanced schistosomiasis patients, diagnosed by the criteria made by Ministry of Health of the People’s Republic of China [20], were first investigated in March through May 2013. To minimize the confounding effects of other liver diseases, patients with hepatic cirrhosis or steatosis, or active consumption of alcohol were excluded. Advanced patients were treated with standard clinic practices which included basic treatment, symptomatic treatment, and etiology treatment in the following year. All patients were further followed up in June through September 2014 for clinical symptoms and abdominal ultrasound test. In the subsequent design of case-control study, 44 patients with ascites in year 2014, diagnosed by abdominal ultrasonography [21], were compared as cases with the other 247 patients without ascites as controls. This study was approved by the Human Research Ethics Committee of the Sun Yat-Sen University (no. 2014032) in China. All patients received an explanation about the scope of the study, such as objectives, procedures, and potential risks, and signed an informed consent statement before inclusion in the study.

Data collection

Clinical history query, physical examination, and abdominal ultrasonography were conducted in the interviews of 2013 and 2014. Information on demographics, history of schistosomiasis, physical features, hepatic fibrosis, portal internal diameter, and ascites was assessed. Additionally, in the initial investigation in 2013, venous blood samples were drawn from all of the investigated individuals under aseptic conditions without stasis using vacuum tubes (Lucky Nation Medical, China) containing 0.562 M EDTA-K3. Sera were obtained after rapid centrifugation for 10 min at 2,000 g, stored in 0.5 ml aliquots at −80°C. Further, a set of biomedical tests measured the serum levels of SEA, value of D-dimer, albumin (ALB), globulin (GLB), thrombin time (TT), and coagulation factor VIII (CF-VIII), hyaluronic acid (HA), procollagen type III (PC-III), type IV collagen (C-IV), laminin (LN), etc. following the standard procedures.

Biomedical measurement

The test sera firstly were defrosted rapidly to room temperature before use. Then, the samples were tested by ELISA using secretory-excretory antigen (SEA) derived from eggs of S. japonicum. Meanwhile, the following control sera were used on each plate: a reference positive sera (serum from a patient with microscopically confirmed S. japonicum infection), and a tropical negative serum (serum from an individual with no evidence of Schistosoma infection). D-dimer was assayed by immunoturbidimetric method (Siemens, no. 412056). The liver function tests, including ALB, GLB, TT, and CF-VIII, were measured by automated biochemistry analyzer (Roche, Basel, Swiss). Additional hepatic fibrosis indices, including HA, PC-
III, C-IV, and LN, were measured by commercially available radioimmunoassay (RIA kits; Shanghai Navy Medical Institution, Shanghai, China). To control the confounding effects of hepatitis B virus (HBV) infection, determinations of HBsAg and anti-HBc were made by automatic light detector and test reagents (Abbott, North Chicago, Illinois, USA). Abdominal ultrasonography was employed to identify the activity of ascites and to observe the morphology of the liver and portal system. Accordingly, hepatomegaly was defined when the thickness of the left lobe was more than 70 mm or the major diameter of the right lobe was over 140 mm; portal hypertension was defined when the width of the inner portal vein was no less than 12 mm.

Diagnostic criteria of schistosomiasis
According to the criteria made by the Ministry of Health of P. R. China, schistosomiasis is defined as (1) long-term or repeating water contact, or (2) finding eggs or miracidia by stool examination, or (3) finding eggs by transrectal biopsy, or (4) serum immunological tests positive. Advanced schistosomiasis is defined as (1) schistosomiasis patients suffering from portal hypertension of schistosomal liver fibrosis or colon granuloma, or (2) schistosomiasis patients with hepatomegaly (mainly in the left lobe).

Data analysis
Continuous variables were described by means and standard deviation or median with quartiles and were further compared between groups with appropriate statistical methods based on their distributions. Categorical variables were described with absolute values and percentages and further tested by a series of χ² tests to examine the associations with ascites in the year 2014.

Defining occurrence of ascites as the state variable, we employed Receiver Operating Curve (ROC) to analyze the predictive value of each significantly different factor between groups. The greatest sum of specificity and sensitivity in the coordinator of the curve then determined the D-dimer’s cutoff value for ascites, which further divided D-dimer levels into 2 groups. Then, we combined all of those categorical variables in a predictive logistic regression model whose dependent variable was the occurrence of ascites in 2014. Thus, predicted probability values were used as the prediction variable in a final ROC analysis of ascites. The Area under the Curve (AUC) was used to judge the performance of this predictive model. The probability of ascites occurrence in 2014 in different combination of risk factors was finally estimated using the final prediction model.

All P-values were 2-sided, and type I error was set as 0.05. All statistical analyses were conducted with SPSS version 18.0 (SPSS Inc. Chicago, Illinois, USA).

RESULTS
Characteristics of advanced patients and liver functions
In a total of 291 patients we recruited in 2013, ascites occurred in 44 of them in 2014. Except for 3 cases (6.8%) with ascites in the liver area, all other patients’ ascites were confined in the kidney area. The median area of the liquid grey area detected by ultrasound was 20 mm² with a range of 8 to 61 mm². As shown in Table 1, patients’ D-dimer levels in 2013 were significantly higher in patients who developed to ascites in 2014 (0.71 ± 2.44 μg/L), compared to the control group (0.48 ± 2.12 μg/L). Additionally, the case group had more proportions of males and level 2 hepatic fibrosis in 2013 than in the control group. We also found significantly lower ALB while higher type IV collagen in the case group. Patients with obstinate ascites are also more likely developing to collateral circulation with splenomegaly and subsequent splenectomy (see an advanced case of schistosomiasis with ascites in the endemic area in 2014 in Fig. 1A and excised spleen in Fig. 1B). No significant differences in other liver function indicators, as well as age, occupation, HBV infection status, treatment during 2014, and portal hypertension were found between the case and control groups (Table 1).

Predicted ascites occurrence with D-dimer and other factors as potential prognostic indicators
Gender, hepatic fibrosis level, ALB, type IV collagen, and D-dimer, which were significantly different between cases and control groups, were used as independent variables to predict the occurrence of ascites in 2014. As shown in Table 2, the AUC of D-dimer was 0.64 with a 95% confidence interval of 0.54 to 0.73. The AUC of other indicators were also significantly greater than 0.5.

Combined prediction model for ascites in 2014
The logistic model which combined all the 5 significantly differentiated indicators of gender, hepatic fibrosis level, ALB, type IV collagen, and D-dimer was significant with R square as
Table 1. Comparison of the general, infective, and hepatic characteristics in baseline (2013) between the cases of advanced schistosomiasis with/without ascites in 2014

| Baseline measurement                        | Ascites occurrence in 2014 | P-value |   |
|---------------------------------------------|-----------------------------|---------|---|
|                                             | Yes (n= 44)                 | No (n= 247) |   |
| Age (mean± SD, year)                        | 62.83± 9.68                 | 61.80± 9.71 | 0.443 |
| Gender                                      |                             |         |   |
| Male                                        | 33 (75.0)                   | 141 (57.1) | 0.026 |
| Female                                      | 11 (25.0)                   | 106 (42.9) |       |
| Occupation                                  |                             |         |   |
| Fisher                                      | 27 (61.4)                   | 163 (66.0) | 0.483 |
| Farmer                                      | 15 (34.1)                   | 65 (26.3)  |       |
| Others                                      | 2 (4.5)                     | 19 (7.7)   |       |
| HBV infection in 2013                       |                             |         |   |
| Negative                                    | 11 (25.0)                   | 61 (25.4)  | 0.996 |
| Antibody                                    | 26 (59.1)                   | 142 (59.2) |       |
| Positive                                    | 7 (15.9)                    | 37 (15.4)  |       |
| Hepatic fibrosis level in 2013              |                             |         |   |
| Level 1                                     | 4 (9)                       | 36 (15)    | 0.075 |
| Level 2                                     | 13 (30)                     | 104 (42)   |       |
| Level 3                                     | 27 (61)                     | 105 (43)   |       |
| Portal hypertension in 2013                 |                             |         |   |
| Yes                                         | 10 (22.7)                   | 41 (16.7)  | 0.337 |
| No                                          | 34 (77.3)                   | 204 (83.3) |       |
| Treatment in 2014                           |                             |         |   |
| Yes                                         | 33 (75.0)                   | 154 (62.3) | 0.107 |
| No                                          | 11 (25.0)                   | 93 (37.7)  |       |
| D-dimer (mg/L)                              | 0.71± 2.44                  | 0.48± 2.12 | 0.005 |
| ELISA (SEA IgG, OD value)                   | 0.34 (0.15-0.58)            | 0.27 (0.17-0.47) | 0.248 |
| ALB (g/L)                                   | 44.5 (40.8-45.8)            | 46.2 (44.0-48.6) | <0.001 |
| GLB (g/L)                                   | 29.3 (24.8-31.8)            | 27.6 (24.7-30.3) | 0.101 |
| A/G                                         | 1.5 (1.4-1.8)               | 1.7 (1.5-1.9) | 0.006 |
| HA (μg /L)                                  | 186.5 (122.8-260.8)         | 151.8 (105.4-234.6) | 0.109 |
| PC-III (μg/L)                               | 83.5 (71.3-113.5)           | 77.6 (63.8-93.4) | 0.053 |
| Type IV collagen (μg/L)                     | 50.0 (43.2-73.3)            | 44.5 (27.1-57.6) | 0.006 |
| LN (μg/L)                                   | 124.7 (108.7-152.3)         | 125.9 (102.6-147.2) | 0.453 |

Fig. 1. (A) A patient of advanced schistosomiasis with ascites in the endemic area (2014, photo by one of the authors). (B) Excised spleen of a patient which developed to collateral circulation with splenomegaly (2009, photo by one of the authors).
As shown in Table 3, hepatic fibrosis level fell out of a significant predictor (OR=1.34, 95% CI: 0.62-2.90) when included with other factors into the final inclusive model. Nevertheless, the performance of this model was better than any individual predictors with the AUC as 0.76 with a 95% confidence interval of 0.68-0.89. In addition, the prediction model can be translated into the following equation for the probability of the occurrence of ascites:

\[ P = \frac{1}{1 + e^{-2.516 + 0.94 \times \text{Gender} + 0.82 \times \text{D-dimer} + 0.29 \times \text{Hepatic Fibrosis level} + \text{ALB} \times (0.609 \times \text{Low} \text{er OR} - 1.551 \times \text{Higher}) + 0.879 \times \text{Collagen}}} \]

This equation would be easily used for clinics since all of the predictors were binary. Practitioners can input the value as 0 or 1 into the equation then get the probability of the occurrence of ascites.

Finally, the probability of incident ascites was found to be as largest as 73.5% when an advanced case is a male, with higher level D-dimer, ALB, C-III, and level 3 hepatic fibrosis. Followed by is a probability of 67.5% when a patient has all these positive risk factors except for level 3 hepatic fibrosis. The probability will decrease to 60.2%, 55.0%, and 53.6 when patients have all risk factors except for 1 symptom of higher ALB, higher D-dimer, or higher TIC, respectively. We listed 7 combinations of the risk factors with the probability in Table 4 for clinical practitioners to refer when these easy-achievable test results are available.

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**Table 2.** The area under curve of each individual predictor for the occurrence of ascites in 2014

| Indicators       | Area  | Std. error | P-value | Asymptotic 95% confidence interval          |
|------------------|-------|------------|---------|---------------------------------------------|
|                  |       |            |         | Lower bound                                 |
|                  |       |            |         | Upper bound                                 |
| D-dimer          | 0.635 | 0.050      | 0.004   | 0.537                                       |
| Gender           | 0.591 | 0.045      | 0.058   | 0.537                                       |
| Hepatic Fibrosis level | 0.595 | 0.046      | 0.045   | 0.505                                       |
| ALB              | 0.697 | 0.042      | <0.001  | 0.615                                       |
| Type IV collagen | 0.641 | 0.046      | 0.006   | 0.550                                       |

**Table 3.** The final inclusive prediction model of ascites occurrence in 2014

| Beta        | Standardized error | Wald     | P-value | OR (95% CI)       |
|-------------|--------------------|----------|---------|-------------------|
| Constant    | -2.516             | 0.444    | 32.060  | 0.000             |
| Gender      | Female Reference   | -        | -       | -                 |
|             | Male               | 0.942    | 0.450   | 4.376             |
| D-dimer     | ≤0.81 ng/mL Reference | -   | -       | -                 |
|             | >0.81 ng/mL        | 0.82     | 0.399   | 4.219             |
| Hepatic Fibrosis level | Class 2 Reference | -   | -       | -                 |
| ALB         | <34 g/L (normal)   | 0.609    | 1.461   | 0.174             |
|             | ≥34 g/L (higher)   | -1.551   | 0.638   | 5.906             |
| Type IV Collagen | <75 µg/L Reference | -   | -       | -                 |
|             | ≥75 µg/L           | 0.879    | 0.512   | 2.950             |

**Table 4.** Probability of ascites occurrence in combinations of risk factors

| Combinations of risk factors | Gender | Higher level of D-dimer | Level 3 hepatic fibrosis | Higher ALB | Higher TIC | Probability (%) |
|------------------------------|--------|-------------------------|--------------------------|------------|-----------|-----------------|
| 1                            | Male   | Yes                     | Yes                      | Yes        | Yes       | 73.5            |
| 2                            | Male   | Yes                     | No                       | Yes        | Yes       | 67.5            |
| 3                            | Male   | Yes                     | Yes                      | Yes        | No        | 60.2            |
| 4                            | Male   | No                      | Yes                      | Yes        | Yes       | 55.0            |
| 5                            | Male   | Yes                     | Yes                      | Yes        | No        | 53.6            |
| 6                            | Male   | Yes                     | No                       | No         | Yes       | 53.1            |
| 7                            | Female | Yes                     | Yes                      | Yes        | Yes       | 52.0            |
DISCUSSION

In this nested case-control study, we observed 44 cases of incident ascites in 291 patients with advanced schistosomiasis japonica and found higher levels of plasma D-dimer as an independent risk factor. D-dimer may predict the prospective occurrence of ascites in combination with other traditional clinical risk factors of gender as male, grade III hepatic fibrosis, high level of ALB, and high level of type IV Collagen.

Ascites is a dominant symptom in advanced schistosomiasis at the current endemic control stage in China. In our 1-year prospective study, 44 out of 291 patients revealed ascites at the end of the follow-up period with an incidence rate of 15.1%. Given the high proportion around 65% of ascites subtype in advanced cases across the whole endemics [22,23], the incidence rate in the present study was even lower than estimated in the transmission-active areas. Ascites was regarded as the trend symptom in advanced schistosomiasis based on an analyses of historical medical records of all patients of advanced schistosomiasis from 2004 to 2013 in Jinmen county, Hubei province [24]. We found males as a risk factor of ascites, which is consistent with previous studies conducted in the same regions but inconsistent with studies conducted in Yunnan where women are the major labor [25]. However, we did not find significant differences in the distributions of occupations across cases and control groups. Occupation was previously regarded as an indicator of exposure to Schistosoma-infected water and occupations with more contacts, such that fishers and farmers of paddy fields are in higher risk [26]. However, with the popularity of machine-farming practice and inhibition of disease transmission, people in those traditional high-risk jobs may be well protected now. Hepatitis B virus infection was previously found to be interacting with schistosomiasis in the liver damaging but we did not find a significance association of HBV infection status with incident ascites in the present study [27]. However, we did find there still remaining 15% patients with active HBV infection while another half (59%) were immunized. This finding indicates a remaining need of HBV treatment in patients with co-infection of both S. japonicum and HBV.

With abdomen ultrasound, we found patients having grade III hepatic fibrosis were more tend to have ascites than those without. Even though the association was marginally significant \(P = 0.075\), the trend was still meaningful and provided evidence for the need of treatment to fibrosis in advanced S. japonicum. Traditional treatment to S. japonicum was mostly targeting at antigen, especially with an extensive usage of praziquantel when apparent symptoms are absent. However, more and more pieces of evidence are indicating that liver damage is constantly advancing along with fibrogenesis caused by Schistosoma eggs. Yin et al. [28] investigated 75 newly diagnosed advanced S. japonicum in transmission-blocked areas and found that 82.7% patients had grade III hepatic fibrosis. They suggested the liver of patients with infection history kept being insulted by the eggs’ residuals, such as proteins and lipopolysaccharide, albeit the larva in the eggs were killed by antigen treatment and patients were free from reinfection. Similarly, we also found worse liver function in patients with incident of ascites that their ALB was significantly lower than those without, but lack significant differences in SEA IgG levels. We also found an indicator of hepatic fibrosis, type IV collagen level, was significantly higher in the case group. All these findings indicated the association of ascites with enduring liver damage and fibrogenesis.

The mechanisms of ascites in advanced schistosomiasis have been widely studied, including physical obstruction caused by eggs and periovular granulomas, liver fibrosis, and portal vein hypertension, along with decreased plasma colloid osmotic pressure [1]. However, the mechanism of liver fibrosis is relatively less studied. In the current study, we found increased levels of D-dimers in patients with incident ascites compared with those without. D-dimer is the primary product when cross-lined fibrin is degraded by plasmin-induced fibrinolytic activity, which is correlated with intravascular levels of fibrin turnover [17]. Thus, D-dimer was regarded as a marker of fibrin turnover, which reflects impairment in coagulation and fibrinolysis pathways. This abnormal function may lead to hepatic fibrosis, as we observed in those patients with incident ascites. Infection of Schistosoma may lead to blood vessel injury and subsequent activate systems involving inflammation, coagulation, and fibrinolysis [29,30]. Additionally, the liver damage will also confine the synthesis of anti-coagulation proteins and thus further aggravate the imbalance to the tendency of thrombosis [29] and increase the risk of liver fibrosis. All of these malfunctions eventually lead to the occurrence of ascites.

Given the critical role of D-dimer in the aforementioned mechanism of ascites and hepatic fibrosis, we combined D-dimer with traditional clinical easy-achievable indicators to predict the incidence of ascites in patients with advanced schistosomiasis japonica. The combination is more capable in pre-
dicting future incidence of ascites than any other indicators on their own. Conventionally a prediction with an area under the ROC curve (AUROC) greater than 0.7 would be regarded as usable in clinics [31]. Thus, the AUROC of the combination of D-dimer with other indicators as 0.76 with a 95% interval of 0.68-0.89 demonstrated its usefulness for a clinical value. As such, we calculated the 7 highest possibilities of future ascites for clinical practitioners to make better decisions when providing treatments to patients with advanced schistosomiasis japonica. Clinical practitioners could use Table 4 as a reference to check the possibility of future ascites incidence risk in patients who may not have any symptoms presently. However, our results are still limited with a small number of samples and relatively short observation time. So, further studies with more representative samples and longer study duration would guarantee to verify the prediction model.

Our study has its own limitations. First, we obtained patients’ sera only once in the year 2013. Although no patients had the symptom of ascites in 2013 when sera were collected, more measurements of D-dimer during the follow-up may provide more accuracy in the prediction model of ascites. Second, our study sample was small in number and follow-up duration was short, which may limit the power to detect more meaningful indicators of incident ascites. Third, we included a number of patients (15%) co-infected with HBV. Patients of advanced schistosomiasis associated with HBV infection usually show persistent viremia and exhibit chronic active hepatitis, which can progress to cirrhosis [13,14]. Even though we set HBV infection status as a confounding factor throughout the analyses process, the proportion of ascites contributed by HBV cannot be thoroughly eliminated.

In conclusion, we suggest that plasma D-dimer is a reliable predictor for incident ascites in advanced schistosomiasis japonica in combination with other clinical easy-achievable indicators. However, further studies with more representative samples and longer study duration would guarantee to verify the prediction model.

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CONFLICT OF INTEREST

We have no conflict of interest related to this work.

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