Immunochromatography Detection of Ricin in Environmental and Biological Samples

Junhua Wu*, Yuxia Wang*, Peiyuan Jia, Chenyu Wang*, Yu Zhao*, Hui Peng, Wenqing Wei, Hua Li*

*Beijing Institute of Pharmacology and Toxicology, Beijing 100850, China
†Beijing Institute of Basic Medical Sciences, Beijing 100850, China
‡General Hospital of Beijing Military Area Command, Beijing 100700, China

* Corresponding author: wangyuxia1962@hotmail.com (Yuxia Wang); amms_hli@126.com (Hua Li)

Abstract

Ricin has been considered as a potential agent of biological warfare or terrorist attack. This work is aimed to develop and evaluate a rapid test strip for the assay of ricin in environmental and biological samples. Two specific antibodies were used respectively as the capture antibody (Mab 4C13) coated on a nitrocellulose membrane and the tracing antibody (MAb 3D74) labeled with colloidal gold. The immunochromatography strips were tested to detect ricin in porridge, salted vegetable, water and other drinks. The detection could be finished within fifteen minutes and its sensitivity was at 10 ng/ml of ricin in water. When no ricin control samples were tested in parallel, some kind of drinks, such as Sprite and Coca cola showed false positive bands even without toxin pollution, indicating the negative control is essential to avoid possible false positive results. The strips could give positive signals after loading human serum sample mixed with 25 ng ml$^{-1}$ of ricin in vitro. When rats were intramuscularly treated with 100 µg kg$^{-1}$ of ricin, the residual ricin in the serum samples could be successfully detected by the strips even at 24 h after intoxication. The immunochromatography test strip is a useful tool to check the possible pollution of ricin in environmental samples such as drink and food, and even to detect the residual toxin in human serum before victim developing intoxication symptoms.

Keywords: Ricin, Antibody, Immunochromatographic Assay, Colloidal Gold, Strip Assay


1. Introduction

Ricin is a heterodimeric ribosome inactivating protein which inhibits protein synthesis, consequently resulting in cell death [7, 17]. The lethal oral dose of ricin in human has been estimated to be 1-20 mg kg$^{-1}$ body weight [1]. In our experimental studies 5µg kg$^{-1}$ of ricin was found to be lethal in mice administered intravenously. Ricin has been considered as a potential agent of biological warfare or terrorist attack, because it can be easily isolated from castor beans and produced in sufficient quantities for weapons [13]. The well known use of ricin might have been the assassination of Georgi Markov, a Bulgarian dissident in 1978 [12, 14].

For the need of antiterrorism, various technologies have been established for ricin detection. Many of them are highly sensitive, such as biosensors based on magnetoelastic and surface Plasmon resonance [15, 19, 4], and other methods that depend on immuno-polymerase chain reaction assay [10] and microelectrode array [3]. They are useful for ricin detection in various environmental samples. Enzyme-linked immunosorbent assay (ELISA) had also been used in ricin assay with its high sensitivity and specificity [8, 11]. Based on the mechanism of ELISA and the engineering design of colloidal gold-based immunochromatographic assay, the strip has been developed and widely used in detection of many kinds of antigens, including the detection of ricin in water and powder samples. Its sensitivity reached 10-50 ng mL$^{-1}$ of toxin in water [6, 16] and be well used in the environmental example assay because of its high sensitivity and rapid operation.

In this work, a strip based colloidal gold-based immunochromatographic assay was established. It was further evaluated for its use in ricin detection in environmental and biological samples.

2. Materials and methods

2.1. Material

Ricin was obtained from the Laboratory of Toxicant Analysis, Beijing Institute of Pharmacology and
Toxicology. Mouse ascites containing monoclonal antibodies (4C13, 3D74) were produced from Beijing Institute of Basic Medical Sciences. The antibodies were purified using protein G-sepharose 4 fast flow (Amersham). Colloidal gold particles were made in our laboratory. Nitrocellulose membrane Unisart CN 140 (Sartorius) was used as membrane and coated with capture antibody and control antibody. Male Wistar rats were supplied by the Animal Center of the Institute. They were housed in a controlled environment (21±2 °C; 55±5% of humidity; 12 h light/dark cycle with light provided between 6 am and 6 pm). Food and water were given ad libitum. All the animal experiments were carried out in the Beijing Center for Drug Safety Evaluation, in accordance with a protocol approved by the Institutional Animal Care and Use Committee of the Center, which is in compliance with the guidelines of the Association for Assessment and Accreditation of Laboratory Animal Care International (AAALAC). Purified goat against mouse IgG was from KENGEN BIOTECH. CO., LTD.

2.2. Preparation of colloidal gold conjugated with antibody 3D74

Colloidal gold was prepared according to a technique of monodispersion colloidal gold [18, 5]. Chlorauric acid was reduced by sodium citrate to produce colloidal gold particles which have a diameter about 11 nm. Newly prepared colloidal gold (40 ml) was conjugated with 5 mg of monoclonal antibody 3D74 at room temperature for 20 minutes. The conjugating solution was centrifuged at 12000 rpm and the precipitation was dissolved in 2 mM boric buffer pH 7.2, supplemented with 1% BSA.

2.3. Establishment of immunochromatographic test strips

The test strip was established following the application guide of Rapid Lateral Flow Test Strips on the Website of Millipore (http://www.millipore.com/techpublications/tech1/tb5000en00). As the capture antibody, purified monoclonal antibody 4C13 (1 mg mL⁻¹) dissolved in 10 mM phosphate buffer pH 7.2 was bound onto the membrane at the position of test line. The same amount of goat against mouse IgG was loaded at the control line. The glassfiber impregnated with colloidal gold-antibody 3D74 conjugate was dried at 37°C for 2 h and put under the sample pad. The strip was made by consistent laminating the membrane, sample pad, conjugating pad and absorbent pad on a support backing.

2.4 Establishment of intoxicated animal mode

For assessment of ricin in serum by the test strips, the rats were injected intramuscularly with 100 µg kg⁻¹ ricin prepared in saline. Blood samples were collected at 0.5, 1, 2, 4, 6, 8, 12, 16 and 24 h post-dose, respectively, and the sera were harvested.

2.5. Detection of ricin in various samples by test strips

2 mM boric buffer pH 7.4, containing 5% sucrose, 0.2% BSA and 0.04% NaN₃ was used as loading buffer. Ricin was diluted with the loading buffer to prepare the testing samples at the concentration range of 0-100 ng mL⁻¹. The testing samples were assayed with the strips. Comparing with negative control, a dark red test line appeared within 15 min was considered as a positive result. The storage stability of the strips was checked after they were kept in a refrigerator at 4 °C for more than 1 year. Water, Sprite, Coca cola and the supernatant of porridge were respectively used as testing samples for the scenarios of the toxin pollution. Serum samples from rats were diluted with the loading buffer and then an aliquot of 100 µL was dropped onto the samples pads of strips.

3. Results

3.1. The detection limit of test strips

According to the method mentioned above, the colloidal gold particles were prepared and labeled with purified antibody against ricin. The maximum absorption of colloidal gold particles with brownish red color was at 517 nm and 524 nm respectively before and after antibody labeling. The glassfiber impregnated with colloidal gold conjugate was put under the sample pad which was overlapped with the nitrocellulose membrane to establish the test strips. The labeled particles could bind with ricin molecules and then be captured by another antibody 4C13 bound at the test line on membrane to give the positive band when they flew from the sample port to the absorption pad.

Ricin in the loading buffer was assayed by test strip. The strips showed two clear red bands for samples with ricin concentrations of 10-100 ng mL⁻¹, while only one control band was observed for the control samples (Fig. 1).

3.2. The storage stability of test strips

The test strips were sealed in aluminium foil bags and kept at 4 °C in a refrigerator for 1 to 3 years. The stability of these strips was tested using ricin standard solutions in loading buffer. The detection limit of 25, 50 and 100 ng mL⁻¹ were obtained for the strips reserved at 4 °C for 1, 2 and 3 years, respectively (Fig. 2.).

3.3. Detection of ricin in food and drink samples by test strips

Poisoning through digestive tract is the one of major paths of ricin poisoning. In order to detect whether the test strips could be used to assay the ricin pollution in food and drink samples, ricin was spiked into water, rice porridge, Coca cola and Sprite. respectively and then these samples were loaded onto the strips. The ricin in water, supernatant of porridge and stir-frying tomato could be well measured with the detection limit of 25 ng mL⁻¹. However, Coca cola and Sprite gave false positive bands for ricin negative samples. Using loading buffer to dilute these two drink samples could eliminate the false band or decrease its intensity of staining, After the
Fig. 1 Ricin detection in loading buffer by test strips. Different concentrations of ricin solutions (100 μL) were dropped onto the sample pad of test strips directly (A) and onto the sample port of test strips with housing (B).

Fig. 2 Storage stability of test strips. Test strips were kept at 4 °C for 1(A), 2(B), 3(C) years in a refrigerator and were tested with ricin spiked in loading buffer at levels over 0-100 ng mL⁻¹.

Fig. 3 Ricin detection in the food and drink samples. Samples of food or drink were diluted with equal volume of the loading buffer spiked with 25 ng mL⁻¹ of ricin. 100 μL sample was dropped onto the test strip. The diluted samples were made from diluting Coca Cola and Spring with loading buffer at the ratio of 1:1 and then mixed with 25 ng mL⁻¹ of ricin in loading buffer before being determined.

Fig. 4 Detection of ricin in human serum. Ricin dissolved in human serum at 1-50 ng mL⁻¹ was used to detect the limit of detection (A). 50 ng mL⁻¹ of ricin in serum was saved in incubator at 37 °C for 0-8 hours and then diluted with equal volume of loading buffer before detected by test strips.

Fig. 5 Ricin detection in serum samples of rats after poisoned with a lethal dose. Ricin was intramuscularly injected at 100 μg kg⁻¹ to the gluteus maximus of rats. Three serum samples at different time points were collected and then diluted (1:1) with loading buffer before being detected.

human serum samples was spiked with ricin in vitro and kept at 37 °C for different time periods before tested with the strips. The result indicated that ricin in human serum could be detected at the sensitivity of 25 ng mL⁻¹ and the ricin was stable in serum at 37 °C for 8 hours without significantly degradation (Fig. 4).

3.5. Detection of ricin in sera of the ricin exposed rats

The test strip could be well used to detect the toxin in environmental samples. The method is rapid and sensitive. It could also capture the ricin in human serum with the detection limit of 25 ng mL⁻¹. In this study, serum samples from ricin exposed rats were also tested. The serum samples were collected after rats received a lethal ricin dose (100 μg kg⁻¹) by intramuscular injection and then diluted with the loading buffer at the ratio of 1:1. The brownish red test lines on the strips were observed for the serum samples harvested at 1 to 24 h post-exposure, suggesting that the strips might be used as a field detection tool for checking the toxin in biological samples taken from exposed victims (Fig. 5).

4. Discussion

Various technologies have been developed for ricin detection [9]. Among them the colloidal gold-based immunochromatographic assay based strip [6, 16] was widely used for field or daily detection. It is sensitive, rapid and easy to handle, especially for detection of environmental samples. In this study a test strip was established using immunochromatographic method. Two antibodies bond with two different epitopes of...
ricin were used for colloidal gold particle labeling and capturing antibody respectively. Using this strip the ricin in boric buffer (pH 7.4) was successfully assayed within 15 minutes with the detection limit of 10 ng/mL (Fig.1). After stored at 37 °C for 3 years, the strip could still give positive response at the ricin level of 50-100 ng mL−1 (Fig. 2).

As previously reported, there were some contents in samples that could disturb ricin detection [2]. In this study various types of samples, including food and beverages were selected to test the possible interferences on ricin detection by strips. When spiked in water, porridge, the supernatants of bean curd and fried tomato ricin was well detected at the level of 25 ng mL−1. However, when ricin-free samples of Coca Cola and Sprite were loaded, false positive bands appeared, indicating some contents in these samples could affect the immunoassay. Dilution with loading buffer would eliminate or reduce the interference (Fig. 3). The strip is able to detect ricin in these samples when the samples are appropriately diluted with loading buffer and the negative controls is assayed parallelly. Other detection method can also be used to reconfirm the result by the test strip.

So far, the immunochromatographic strips are mainly used for ricin detection in environmental samples, such as water and solid powder. No report for their use in biological samples has been seen, despite of the fact that the early detection of ricin intoxication in human is extremely important for taking necessary medical treatment. Generally, the latency of clinical symptom of ricin poisoning is several hours depending on the route of intoxication and the toxin dosage applied. A rapid identification of poisoning source and a correct diagnosis are very important for successful clinical treatments against ricin poisoning. In this study the strip was tested with serum samples from human and rat to determine whether it could be used to detect ricin in biological samples. The human serum spiked with ricin was maintained in 37 °C incubator for different time periods before loading onto the strips. Results of in vitro test indicated that the strip was able to detect ricin in human serum. This result encouraged us to conduct a further test with the serum samples taken from the rats that were intramuscularly injected with a lethal dose of ricin. Positive test bands were observed for the sera collected at 2 to 24 hour post intoxication. The response time was only 10 min. The result indicates that this method is rapid, sensitive and suitable for the emergency detection of human intoxication with ricin. The results of immunochromatographic strips can be reconfirmed using our previously reported Sandwich ELISA method, which could quantitatively determine ricin in serum samples of poisoned mice [11]. The combination of the strip and the ELISA assay will provide quick qualitative and quantitative detection for ricin poisoning cases. Our further work will focus on developing a ricin enrichment method for serum or plasma samples to increase the assay sensitivity, which is also very important for the ricin assay especially in the cases of sublethal poisoning.

Acknowledgements

We are grateful to Dr. Liming Guo and Ming Yu for the gift of mouse ascites to prepare Mabs against ricin. This work was supported by a Grants from the National Natural Science Foundation of China (NO. 30772593, 30973562), the National Basic Research Program of China (NO. 2010CB933904).

References


Copyright: (c) 2011 J. Wu, et al. This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.