Microplate magnetic chemiluminescence immunoassay for detecting urinary survivin in bladder cancer

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Abstract. Survivin is a tumor marker for bladder cancer; however the role of urinary survivin levels has not been fully elucidated due to the limitations of current detection methods. Based on two survivin-specific monoclonal antibodies (McAbs) already confirmed through enzyme linked immunosorbent assays, the present study aimed to establish a microplate magnetic chemiluminescence immunoassay (CLIA) for the detection of urinary survivin levels and evaluate its application for the diagnosis of patients with bladder cancer. Horseradish peroxidase and biotin conjugates were used to label two different anti-survivin McAbs, respectively. The labeled antibodies combined with survivin to form a sandwich immune complex. The streptavidin magnetic particles (MPS) served as the solid phase and the separator. The relevant parameters involved in the immunoassay, including the immunosorbent reagents used and the physicochemical parameters were optimized. Then, urine samples from 130 patients with bladder cancer and 113 healthy controls were detected, and analyzed using the established method. The method was linear to 1,000 ng/ml survivin with a detection limit of 0.83 ng/ml. The intra- and inter-assay coefficients of variation were <8, and <11%, respectively. The concentration of diluted survivin and the dilution ratios gave a linear correlation of 0.9989. The results demonstrated that the urinary survivin levels in patients with bladder cancer were significantly higher (P<0.001) compared with that in healthy controls. At a survivin concentration of 2.0884 ng/ml, the sensitivity and specificity were 86.9 and 61.9%, respectively. Furthermore, the urinary survivin levels were positively correlated with metastatic stage, histological stage and recurrence (P<0.01). In conclusion, the present study preliminarily proposed a microplate magnetic CLIA for survivin detection and further evaluated the value of urinary survivin as a diagnostic marker for bladder cancer.

Introduction

Bladder cancer, the most common malignancy of the urinary tract and the seventh most prevalent cancer worldwide (1), is diagnosed based on the combination of urethrocystoscopy and voided urine cytology (2). While voided urine cytology has an extremely low sensitivity (3) considering its dependence on tumor grades, what makes urethrocystoscopy the ‘golden standard’ for the screening and follow-up of bladder cancer patients (2). However, the procedure of urethrocystoscopy is invasive and discomfort for patients. Therefore, it is important to develop better non-invasive detection methods for the early diagnosis of bladder cancer (4). Several urinary tests including Bladder tumor antigen (BTA) and Nuclear matrix protein-22 (NMP-22) have been approved by FDA (5). Recently additional possible urinary biomarkers have been investigated (6) including Cyfra 21-1 (7), Matrix metalloproteinase-7 (MMP-7) (8) as well as survivin.

Survivin, a 16.5 kDa protein, is one member of the inhibitor of apoptosis protein (IAP) family and has a unique role in apoptosis (9-11). Survivin is expressed in various tumors while in normal adults it is only expressed in tissues as thymus, endothelium and placenta (12-14). Given its differential expression between normal and tumor tissues, survivin is considered to be a unique marker for the diagnosis of cancer. Studies have shown that survivin is associated with poor prognosis in certain kinds of tumors including bladder cancer (15,16). In bladder cancer, survivin is expressed in epithelium and its expression could be detected by immunohistochemistry (IHC). What's more, as a malignancy of the urinary tract, survivin is expressed in urine that could be detected at the protein and mRNA levels (17). Certain studies found that the measurement of survivin mRNA in urine seemed to perform better than voided urine cytology (15). Recent increasing attention has been focused on the value of urinary survivin detection in bladder cancer, while its clinical utility still remains to be elucidated partly due to the limitation of detection methods.

ELISA method is considered accurate and reliable, which has also been used for the detection of urinary survivin (18,19). But the assay is time consuming with poor reproducibility.
Thus alternative methods with better performance are needed. Chemiluminescence immunoassay (CLIA) is a much more effective measurement with the advantages of high sensitivity, uniformity and broad dynamic range (20,21). In this study, we tried to develop a microplate magnetic CLIA with two anti-survivin monoclonal antibodies (McAbs) prepared by our laboratory. Besides, in order to evaluate the potential application of the established method, urinary survivin levels of bladder cancer patients and healthy controls were examined.

Materials and methods

Animals. Female Balb/c mice weighing 18-22 g were purchased from the Laboratory Animal Centre of China Academy of Medical Sciences. The Animal Care Committee of Peking University approved the animal experiments. We performed animal studies in accordance with the Experimental Animal Management Ordinance approved by the Scientific and technological committee of China.

Apparatus. Protein-A/G sepharose (HiTrap Protein G HP, 1 ml) was purchased from GE Healthcare Life Sciences (Buckinghamshire, UK). ALC-B6 peristaltic pump was purchased from Alcott Biotech Co., Ltd. (Shanghai, China). The chemiluminescence detection was carried out using SpectraMax L microplate reader from Molecular Devices, LLC (Sunnyvale, CA, USA). The IKA® MS3 Digital (Staufen, Germany) was employed to blend the solutions in microplates. The incubation procedure at 37°C was carried out at an electric heat constant temperature incubator. The white opaque 96-well flat-bottomed microplates were from Thermo Fisher Scientific (Waltham, MA, USA). A magnetic separation device for 96-well microplate purchased from Beaver Beads Co., Ltd. (Suzhou, China) was used for the separation procedure.

Chemicals and solutions. Incomplete Freund’s adjuvant (IFA), PEG, horseradish peroxidase (HRP; H1759), sodium borohydride (NaBH4; 10H3440), sodium m-periodate (NaIO4; 38F-0860) and (+)-biotin-N-hydroxysuccinimide (NHSB; HMBD0595 V) were from Sigma-Aldrich Co. (St. Louis, MO, USA). Hypoxanthine-aminopterin-thymidine (HAT) and hypoxanthine-thymidine (HT) were from Corning Inc. (Corning, NY, USA). BCA protein assay kit was obtained from Thermo Fisher Scientific.

Generation of hybridoma cell lines. The mouse was immunized with 150 μg purified recombinant human sequence survivin protein MS2-survivin and the same volume of IFA every two weeks until the serum antibody titer was up to 1:10,000. Then 150 μg MS2-survivin was administered into the intra-peritoneal cavity without IFA as the final immunogen boost, 3 days before mice’s spleen was harvested for cell fusion. Isolated splenocytes were mixed with myeloma cells at the ratio of 10:1 with the presence of polyethylene glycol (PEG) and HAT medium. Six to eight days passed and hybrid colonies survived. The colonies were screened by ELISA with MS2-survivin and MS2-PAI for three times. Positive clonal cells were transferred and expanded. Then the hybridoma cell line was established. Cells were passaged in HT medium after HAT selection is completed (2-3 weeks).

Preparation and purification of McAbs to survivin and standard protein. Hybridoma cell lines (C6 and E6) secreting anti-survivin McAbs with high signals on ELISA were expanded as ascetic fluids in BALB/c mice. The McAbs were purified by protein-G affinity chromatography from the ascetic fluids. Antibody concentrations were determined by using the BCA protein assay kit. Our laboratory has already done experiments before comparing the newly generated antibodies with commercial antibodies. We have detected survivin expression in cancer by several methods such as IHC and western blot analysis, using the newly generated antibodies and the commercial antibody (an anti-survivin monoclonal antibody from Santa Cruz Biotechnology, Inc., Dallas, TX, USA) simultaneously. Results showed that the newly generated antibodies were comparable with the commercial antibody (22). Hence, our newly generated antibodies have been identified to be reliable. Recombinant human sequence survivin protein MS2-survivin produced by our laboratory was used as the standard protein (23). The standard series were prepared by diluting MS2-Survivin stock with dilution buffer to target values of 0, 10, 50, 100, 500, 1,000 ng/ml, assigning to S0, S1, S2, S3, S4 and S5, respectively.

Preparation of HRP-labeled antibody. Anti-survivin McAbs (C6, E6) were labeled with HRP, respectively. Briefly, purified McAbs were dialyzed against several changes of carbonate buffer [0.1 M sodium carbonate buffer (NaHCO3/Na2CO3) pH 9.5 at 4°C overnight. The HRP protein dissolved in deionized water at a concentration of 5 mg/ml was pretreated with NaIO4 stirring for 20 min at room temperature in dark, and then was dialyzed against CH3COONa (1 mmol/l sodium acetate buffer) pH 4.4 at 4°C overnight. Equivalent pretreated McAbs and HRP solution were blended and incubated at room temperature for 2 h with gentle stirring in dark. Then NaH2B was added, stirring at 4°C for 2 h. The reaction solution was dialyzed against several changes of PBS buffer (0.01 M sodium phosphate, 0.15 M sodium chloride, pH 7.4) at 4°C overnight. After dialyzing, the reaction mixture was applied to a Sephacryl S-200 column to remove unlabeled HRP. The HRP-conjugated McAbs were stored at -20°C until use.

Preparation of biotinylated antibody. Anti-survivin McAbs (C6, E6) were coupled with biotin, respectively using a standard protocol. Briefly, 1 mg of anti-survivin monoclonal antibody dissolved in 1 ml of 0.1 mol/l carbonate buffer (pH 8.0) was dialyzed against carbonate buffer (pH 8.0) at 4°C for 2 h. 1 mg of NHSB was dissolved in 1 ml of DMSO. 1 ml McAbs solution was added into 120 μl of NHS-D-biotin solution with gentle
stirring and incubated at room temperature for 4 h. Then 9.6 µl of 1 mol/l NH₄Cl was added, stirring at room temperature for 10 min. The reaction solution was dialyzed against several changes of PBS buffer (0.01 M sodium phosphate, 0.15 M sodium chloride, pH 7.4) at 4°C to remove unlabeled biotin. The dialyzed biotinylated McAbs were stored at -20°C until use.

**ELISA procedure.** E6 (100 µl, 2.5 µg/µl) was coated onto a 96-well microplate at 4°C overnight. After blocking with 200 µl 5% skimmed milk in PBS for 1 h at 37°C and washing 3 times with PBST, 100 µl survivin standards and urine samples were added into the wells. After incubating at 37°C for 1 h and washing 3 times, 100 µl HRP-labeled C6 was added to each well. The microplate was incubated at 37°C for 1 h and washed 3 times. Then substrate solution was added to the wells and every pore's absorption was determined at 450 nm. McAbs (C6 and E6) used in this ELISA procedure have been identified to be reliable as mentioned above.

**Microplate magnetic CLIA procedure.** The immunoassay procedure of microplate magnetic CLIA in this study is displayed in Fig. 1 and the detailed steps were as follows: First, 50 µl HRP-labeled anti-survivin McAb, 50 µl biotinylated anti-survivin McAb and 20 µl survivin standard solution were added into the micro-well and incubated for 20 min at 37°C. After the sandwich reaction, 1 µl streptavidin MPs were added to react with immunoassay reagents for another 10 min (capture time) at 37°C. Subsequently, the separation procedure was carried out, during which the magnets attracted streptavidin MPs and any specific captured materials to the bottom of micro-wells. The micro-wells were washed with 200 µl of washing solution three times after removing unwanted materials. Finally, 150 µl of CL substrate was added and the relative light unit (RLU) was measured in the dark.

**Human specimens collection and detection.** All human specimens were obtained from Peking University Cancer Hospital. All of the cancer patients were diagnosed histopathologically and staged according to the tumor node metastasis (TNM) classification released by the American Joint Committee on Cancer (AJCC 7th edition, 2010). Healthy controls with a negative cystoscopy were chosen at the medical examination center. A total of 130 urine samples of bladder cancer patients and 113 urine samples of healthy controls were collected from January to July 2016. All urine samples were collected at the day and centrifuged immediately at 3,000 r/min for 5 min, and the supernatant was aliquoted, and stored at -40°C until detection. The samples were tested for survivin levels using the established method directly without any pretreatment. All patients and healthy controls were informed consent for participation in this study. The study was approved by the Ethics Committee of Peking University Cancer Hospital and Institute. All study procedures were in accordance with the Helsinki Declaration.

**Data analysis.** Standards and samples were measured, and CL intensity values were integrated. Standard curves were obtained by plotting the logarithm of CL intensity (in RLU) against the logarithm of standard concentration and fitting to a linear equation. Student’s t-test was used for the comparison of the variables between groups. Data was expressed as mean values ± standard deviation. Enumeration data was expressed as percentages analyzing by χ² test. A p-value of <0.05 was considered to be significant. Cutoff value was determined by the optimal Youden's index (sensitivity + specificity-1). All the statistical analysis was performed using the SPSS statistical software (SPSS for Mac, version 20).

**Results**

**Determination of the proper antibody combination.** The antibody pair used in this assay has been tested in a sandwiched ELISA assay. One of the two antibodies was labeled with HRP and the other was to be biotinylated. Since there were two possible combinations, the first development step was the selection of the optimal antibody combination. In order to test whether biotinylated E6 and HRP-labeled C6 or biotinylated C6 and HRP-labeled E6 combined better; three different concentrations (10, 100, 1,000 ng/ml) of the survivin standard and a negative control (dilution buffer) were analyzed under both combinations. Then the signal-to-background (S/B) ratio was calculated and compared for the determination of the better antibody combination. Student’s t-test was used for data comparison. According to Fig. 2, HRP-labeled C6 and biotinylated E6 were determined as the proper antibody pair which providing a higher S/B ratio (P<0.05).

**Influence and optimization of immunoassay reagents.** The immunoreaction reagent is an important parameter affecting the sensitivity and accuracy of immunnoassay, especially in a sandwich immunoassay. In this experiment, the dilution ratios of HRP-labeled antibody and biotinylated antibody were studied and optimized. The HRP-labeled antibody and biotinylated antibody were diluted with dilution buffers to a series of dilution ratios with the standard survivin concentration of 100 ng/ml. Data was analyzed using Student's t-test. As shown in Fig. 3, the RLU increased when the dilution ratios of HRP-labeled antibody increased from 1:90,000 to 1:15,000 at all dilution ratios of the examined biotinylated antibody (P<0.05). As for the biotinylated antibody, the RLU was approximately a peak with the dilution ratio of 1:18,000 (P<0.05). Considering both the sensitivity and the assay cost, we selected the dilution ratios of 1:15,000 and 1:18,000 for HRP-labeled antibody and biotinylated antibody respectively.

**Influence and optimization of physicochemical parameters**

**Influence of immunoassay incubation time.** Incubation time of the immunoreagents may have a direct effect on the sensitivity of the immunoassay. Incubation time from 10 to 60 min (10 min as the interval) was studied. As shown in Table I, RLU increased with increasing reaction time, while RLUS/RLU₀ (reflecting sensitivity) and RLUS₀/RLUS (reflecting linear range) increased with time up to 20 min and after 20 min the values tended to decrease. Based on all this and considering nonspecific absorption would improve with a longer incubation time, incubation time of 20 min was selected.

**Influence of the volume of MPs.** The quantity of streptavidin MPs was critical for the immunoassay system. The volumes of the MPs (10 mg/ml) were optimized from 0.2 to 2.0 µl.
Student’s t-test was used for data comparison. As shown in Fig. 4, the RLUs increased with the volume of MPs from 0.2 to 1.0 µl and then decreased gradually as the volume increased from 1.0 to 2.0 µl. An excess of MPs might absorb the emitted light (24). Therefore the optimal volume of MPs was set to 1.0 µl (P<0.05).

**Influence of capture time.** In this experiment, MPs were used as the separation agent. After adding MPs, immunoassay reagents were captured. Capture time from 10 to 60 min (10 min as the interval) was also explored to make clear its effect on the sensitivity of this immunoassay. As shown in Table II, RLUs increased weakly with increasing capture time, meanwhile RLUS/RLUS₀ (reflecting sensitivity) and RLUS₂/RLUS₀ (reflecting linear range) tended to decrease as time gone by. Thus, capture time of 10 min was selected.

**Influence of the volume of the CL substrate and chemiluminescence reaction time.** The CL substrate was a pivotal factor related to the CL intensity and the sensitivity of the assay. In this experiment, the CL substrate volume from 0 to 200 µl (25 µl as the interval) and the chemiluminescence reaction time from 10 to 60 min (10 min as the interval) were examined. Data was compared by Student’s t-test. As shown in Fig. 5, the
RLUs increased with increasing volume of CL substrate up to a maximum at 150 µl and then gradually decreased (P<0.05). As to the chemiluminescence reaction time, RLUs decreased over time (P<0.05). Thus, 150 µl of CL substrate and 10 min of chemiluminescence reaction time were chosen.

**Method evaluation**

**Calibration curve and sensitivity.** Under the optimal conditions, dose-response curve (Fig. 6) was obtained. By first obtaining the average RLU signals for 10 replicates of S₀ and then adding 2 standard deviations (SDs) into the dose-response curve the detection limit was calculated. The obtained detection limit for survivin was 0.83 ng/ml.

**Precision.** In order to obtain the intra-assay precision, three different concentrations of standards were measured 10 times within one assay. Similarly, these standards were analyzed on 5 different days using the same protocol (2 replicates per run) to obtain the inter-assay variation. As shown in Table III, intra-assay and inter-assay CVs were <8 and <11%, respectively.

**Linearity-dilution effect.** The linearity-dilution effect was studied by selecting a certain human sample with relatively high concentration. This sample was then diluted to a series of concentrations with dilution buffer. The results were shown in Fig. 7.

**Comparison with ELISA.** Survivin levels of 60 urine samples of bladder cancer patients were determined simultaneously using ELISA and the newly established method. As can be seen in Fig. 8, two methods were compared and there was a good agreement with the correlation coefficient of 0.9860 (P<0.01).

**Sample analysis**

**General data of groups.** A total of 130 bladder cancer patients and 113 healthy controls were enrolled in the present study. Table IV summarized the general characterization of two groups (χ² test).

**Analysis of urine survivin levels between groups.** Receiver operating characteristic (ROC) curve based on the detection of 130 bladder cancer patients and 113 healthy people was shown in Fig. 9. The area under the curve was 0.799. When survivin concentration was 2.0884 ng/ml, sensitivity and specificity were 86.9 and 61.9%, respectively. We compared the urinary survivin levels between bladder cancer patients and healthy controls (Student’s t-test). The urinary survivin levels were significantly higher in bladder cancer patients than in healthy controls (P<0.001). Table V and Fig. 10 gave the results of comparison and the scatter plot of survivin levels between groups.

**Comparison of urine survivin levels in different clinicopathological categories of bladder cancer patients.** Clinicalpathological features including age, gender, smoke, hypertension, metastasis stage, lymph node status, TNM stage, histological stage, tumor size, tumor thrombus and primary or not were obtained. We compared the urinary survivin levels between different clinicopathological categories of bladder cancer patients using Student’s t-test. Results showed that among all the factors, urinary survivin levels associated with metastatic stage, histological stage and recurrence (P<0.01). The comparison results were summarized in Table VI.
Discussion

Survivin is expressed during fatal development and involved in blocking caspases (25–27) as well as favoring aberrant progression through mitosis (28). Studies have found that overexpression of survivin in human malignancies could be associated with carcinoma metastasis and progression (29). High expression of survivin in bladder cancer is associated with several unfavorable prognostic factors such as increasing recurrence rates, progression and resistance to therapy (30). Though a series of studies have evaluated urinary survivin as a biomarker for bladder cancer, the exact role of urinary survivin is still unclear partly due to the suboptimal measurements (31,32).

Several methods have been used to detect survivin expression including IHC, real-time-PCR, Reverse transcription-polymerase chain reaction (RT-PCR) and ELISA (32–34). However, RT-PCR and Real-time PCR are costly which requires professional facilities. IHC is accurate while tissue specimens are difficult to get. Though ELISA is considered accurate and sensitive, it is to some content limited as time consuming with a bad uniformity. Here in this research, a new measurement was established for the detection of urinary survivin.

In the present study, a microplate magnetic CLIA was established for the analysis of urinary survivin levels. By combining magnetic separation with chemiluminescence detection system, this research first applied a novel sandwich immunoassay to the determination of urinary survivin. The dilution ratios of immunoreagents as well as physicochemical parameters were optimized during the method development.

With the established and refined measurement, 130 samples of bladder cancer patients and 113 samples of healthy controls were detected for urinary survivin levels. The results showed that urinary survivin levels of bladder cancer patients were significantly higher than that of healthy controls, which is consistent with some other studies (13,19,35) while disaccord with another study (18). Compared with ELISA, a main advantage of the novel measurement in this experiment is its rapidity. Besides, by
utilizing microplates, a high flux of analysis and a good uniformity are realized (20,36). At the same time, the microplates were pre-coated with 1% (w/v) BSA in PBS buffer to avoid the high non-specific absorption of the plates. The microplate magnetic technologies were rarely reported (21,37,38) and even few applications were found for the detection of survivin. Here in this study, streptavidin MPs were used as separation reagents, combined with the magnetic separation device, greatly simplifying the separation procedure. Furthermore, the new measurement reduces the amount of immunoassay reagents and the volume of samples, showing great potential in the future.

Meanwhile, it should be noted that this research has some limitations. On one hand, the use of MPs in this research was expected to provide many more active binding sites and increase the sensitivity as well as facilitate larger linear range in the detection (39), but results were not positive. When the cutoff survivin level was 2.0884 ng/ml, sensitivity and specificity were 86.9 and 61.9% respectively, not superior to some other reports of different methods (18,32,40). This may be partly due to the heterogeneity between different studies as well as different cut-off values. While the complexity of urine components and variability of urine pH may be the main factor hinders the development of more sensitive methods (41,42).

On the other hand, the linearity-dilution effect was unstable and varied a lot between different samples using this method. This may be attributed to the imperfect diluent used in this research, which is unable to deal with the complexity of urine components.

The main objective of our study was the establishment of a novel immunoassay for urinary survivin detection; for this purpose urine sample of patients treated recently (from January 2016 to July 2016) in our hospital were collected. It is too close to track the long-term prognosis information of these patients. Since long-term prognosis information of patients is important for clinical course evaluation, our study is limited to some extent considering the deficiency of the information. This study showed that urinary survivin level was correlated with metastatic stage, histological stage and recurrence, which implied survivin as a potential marker of disease progression. It could be better if we could obtain the long-term prognosis information to further evaluate the value of survivin. In the next years, we will follow-up the bladder cancer patients included in this study to do further investigation according to your suggestion. Moreover, we established a new method

Table III. Intra- and inter-assay variability for survivin.

| Sample no. | Times of replication | Concentration (ng/ml) | CV (%) | Days of replication | Concentration (ng/ml) | CV (%) |
|------------|----------------------|-----------------------|--------|---------------------|-----------------------|--------|
| 1          | 10                   | 331.5                 | 0.7    | 5                   | 330.7                 | 1.62   |
| 2          | 10                   | 128.76                | 3.7    | 5                   | 127.83                | 7.87   |
| 3          | 10                   | 22.91                 | 7.54   | 5                   | 23.14                 | 10.46  |

Table IV. General data of bladder cancer patients and healthy controls.

| Variable | Patients (n=130) | Controls (n=113) | P-value |
|----------|-----------------|-----------------|---------|
| Gender, n (%) | 0.331          |                 |         |
| Female    | 36 (27.7)       | 38 (33.6)       |         |
| Male      | 94 (72.3)       | 75 (66.4)       |         |
| Age, n (%) | 0.239          |                 |         |
| >62       | 74 (56.9)       | 73 (64.6)       |         |
| ≤62       | 56 (43.1)       | 40 (35.4)       |         |

Figure 7. The linearity-dilution effect of the high concentration sample.

Figure 8. Correlation between urinary survivin levels measured by the newly established method and ELISA.
and compared it with ELISA, while it is not the end. There are other methods for the evaluation of survivin levels such as methods for mRNA detection and IHC (31,43). Though we have obtained data of these methods from other papers, it is not so accurate considering sample differences (31,43-45).

Therefore it is significant for us to collect more specimens and perform different methods to make better comparisons in the future.

This research implies urinary survivin as a potential tumor marker of metastasis, progression and recurrence in addition to diagnosis for bladder cancer with the novel established method. While considering the small number of patients in this study, a larger series of samples and further studies are needed in order to understand the role of survivin.
in bladder cancer better. Besides, since the molecular mechanism of bladder cancer is complicated, it may be much more predictive to combine other urinary markers with survivin (46,47). Last but not least, the method itself is not perfect given the unsatisfactory sensitivity and linear range. Considering the complexity of urine matrix (48), it is difficult to develop immunoassays with good performance for urinary biomarkers. Therefore much more work should be done to explore a better measurement for urinary survivin detection based on this research.

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