A Renewable and Ultrasensitive Electrochemiluminescence Immunosenor Based on Magnetic RuL@SiO$_2$-Au~RuL-Ab2 Sandwich-Type Nano-Immunocomplexes

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Abstract: An ultrasensitive and renewable electrochemiluminescence (ECL) immunosensor was developed for the detection of tumor markers by combining a newly designed trace tag and streptavidin-coated magnetic particles (SCMPs). The trace tag (RuL@SiO$_2$-Au~RuL-Ab2) was prepared by loading Ru(bpy)$_3^{2+}$ (RuL)-conjugated secondary antibodies (RuL-Ab2) on RuL@SiO$_2$ (RuL-doped SiO$_2$) doped Au (RuL@SiO$_2$-Au). To fabricate the immunosensor, SCMPs were mixed with biotinylated AFP primary antibody (Biotin-Ab1), AFP, and RuL@SiO$_2$-Au~RuL-Ab2 complexes, then the resulting SCMP/Biotin-Ab1/AFP/RuL@SiO$_2$-Au~RuL-Ab2 (SABAR) sandwich-type immunocomplexes were absorbed on screen printed carbon electrode (SPCE) for detection. The immunocomplexes can be easily washed away from the surface of the SPCE when the magnetic field was removed, which made the immunosensor reusable. The present immunosensor showed a wide linear range of 0.05–100 ng mL$^{-1}$ for detecting AFP, with a low detection limit of 0.02 ng mL$^{-1}$ (defined as S/N = 3). The method takes advantage of three properties of the immunosensor: firstly, the RuL@SiO$_2$-Au~RuL-Ab2 composite exhibited dual amplification since SiO$_2$ could load large amount of reporter molecules (RuL) for signal amplification. Gold particles could provide a large active surface to load more reporter molecules (RuL-Ab2). Accordingly, through the ECL response of RuL and tripropylamine (TPA), a strong ECL signal was obtained and an amplification analysis of
protein interaction was achieved. Secondly, the sensor is renewable because the sandwich-
type immunocomplexes can be readily absorbed or removed on the SPCE’s surface in a
magnetic field. Thirdly, the SCMP modified probes can perform the rapid separation and
purification of signal antibodies in a magnetic field. Thus, the present immunosensor can
simultaneously realize separation, enrichment and determination. It showed potential
application for the detection of AFP in human sera.

Keywords: [Ru(bpy)$_3$]$^{2+}$@SiO$_2$-Au; alpha-fetoprotein; sandwich-type immunoreaction;
screen printed carbon electrode; electrochemiluminescence

1. Introduction

The clinical analysis of cancer biomarkers is critical to the early diagnosis of cancer and proteomics
research, which will also promote the understanding of cancer diseases’ related biological
processes [1-3]. Alpha-fetoprotein (AFP) is a well-known tumor marker related to hepatocellular
carcinoma (HCC) [4]. The average concentration of AFP in serum often increases under cancer
conditions from 5–20 ng mL$^{-1}$ [5]. Thus the development of ultrasensitive detecting methods for trace
levels of AFP in human serum is important for the early diagnosis of cancers. Compared with
conventional immunoassays such as enzyme-linked immunosorbent assay (ELISA) and
chemiluminescence immunoassay, the ECL assay not only shows high sensitivity and wide dynamic
concentration response range, but also is potential and spatially controlled [6-10]. Among various ECL
systems, Ru(bpy)$_3^{2+}$-based ECL has gained importance owing to its superior properties of high
sensitivity and good stability in aqueous solution under moderate conditions [11,12]. Recently, some
stable and sensitive ECL biosensors based on Ru(bpy)$_3^{2+}$ doped SiO$_2$ nanoparticles have been
intensively investigated [13-15]. On one hand, SiO$_2$ nanoparticles are considered a good matrix
because their surfaces are easily functionalized and conjugated to bioactive molecules, which shows
great potential in bioanalysis. On the other hand, the SiO$_2$ matrices can resist dye molecules coming off
and this increases their photostability [13,16]. In addition, gold nanoparticles (Au NPs) have gained
attention in the last years due to the unique structural, electronic, optical, and catalytic properties which
have made them become very attractive materials for biosensor systems and bioassays [17]. Yuan and
co-workers have proposed a novel sensitive ECL immunoassay with AFP antibody labeled
RuL@SiO$_2$-Au as probes [18]. However it required complex modification procedures on the electrode.
Furthermore the sensor cannot be renewed after the immune products are formed on electrode.

Conventional electrodes, such as glassy carbon electrode (GCE), Au electrode or magnetic carbon
paste electrode (MCPE), are relatively expensive and bulky. Moreover, they are not convenient to add
or remove the magnetic iron on the long cylindrical electrode to attract the magnetic nanoprobe.
However, amperometric biosensors fabricated with screen printed carbon electrodes (SPCEs) have the
advantages of integration of electrodes, simple manipulations, low cost and low consumption of sample
which can be used for one step determination, and then discarded [19,20]. Recently, magnetic
nanoparticles have also gained increasing interest and have been widely applied in immunoassays [21,22]
due to their biocompatibility, superparamagnetism and good electron conductivity [23], which can
simplify the process of protein immobilization and separation [24]. The magnetic nanoprobes strategy
developed recently has proven to be a highly sensitive technique for detecting human tumor cells, and is
especially well suited to separate and at the same time detect low-concentrations of proteins [25,26]. More importantly, the magnetic probes can be modified and
removed from the surface by a magnetic field added on the flat bottom of SPCEs. All these steps can
make the electrode’s surface renewable and simplify the electrode modification steps. Furthermore, if a
RuL labeled antibody is employed instead of pure antibody to be labeled on RuL@SiO$_2$-Au nanoparticles
as probes (RuL@SiO$_2$-Au–RuL–Ab2), the relevant ECL signal could greatly improve due to the
increased number of reporter molecules (RuL) that are modified on the electrode surface.

Our present work is motivated by the promising applications of the SBAR sandwich-type
immunocomplexes for ultrasensitive detection of AFP and renewable surface of SPCEs. In this study,
the streptavidin-coated magnetic particles (SCMP), as supporting material, not only perform the rapid
separation and purification of signal antibody on magnetic field, but they also enhance the fixed
capacity of Biotin-Ab1 to improve the detection range. Furthermore, the SBAR sandwich-type
immunocomplexes were readily immobilized on the working electrode of SPCEs by magnets,
therefore, unlike traditional electrochemical immune sensors, the electrodes do not require complex
modification and cleaning. In addition, the RuL@SiO$_2$-Au–RuL–Ab2 composite exhibited dual
amplification since SiO$_2$ could absorb large amount of reporter molecules (RuL) and gold particles
could provide large active surface to load more reporter molecules (RuL–Ab2). Thus, the SCMP/Biotin-
Ab1/AFP/RuL@SiO$_2$-Au–RuL–Ab2 (SBAR) sandwich-type immunocomplexes were easy to control
and have strong ECL signals.

2. Experimental Section

2.1. Reagents and Materials

Alpha-fetoprotein antibody (Anti-AFP, 12 mg L$^{-1}$) was from Biocell Company (Zhengzhou, China). Elecsys AFP Kits were from Roche Diagnostics GmbH. Tris(2,2’-bipyridyl)ruthenium(II) chloride Hex hydrate-(Ru(bpy)$_3$Cl$_2$6H$_2$O) was from Strem Chemicals. Tripropylamine (TPA) was from Tokyo Kasei Kogyo Co, Ltd. Gold chloride (HAuCl$_4$), BSA (96–99%) and Triton X-100 (TX-100) were bought from Sinopharm Chemical Reagent Co.Ltd (Shanghai, China). Tetraethylorthosilicate (TEOS) and 3-aminopropyltriethoxysilane (APTES) were from Aladdin Chemistry Co. Ltd. Gold nanoparticles (Au-NPs, ~16 nm in diameter) were prepared according to the procedures reported by Eunüstan et al. [27]. Phosphate buffered solution (PBS, pH 7.4) was prepared using 0.1 M Na$_2$HPO$_4$, 0.1 M KH$_2$PO$_4$ and 0.1 M KCl. Blocking buffer solution consisted of a PBS with 3% (w/v) BSA and 0.05% (v/v) Tween 20. Washing buffer solution consisted of a PBS with 0.1 M NaCl and 0.05% (v/v) Tween 20 (PBST). All other chemicals were of analytical grade and all solutions were prepared with
doubly distilled water.

2.2. Apparatus

ECL experiments were carried out using a MPI-B model electrochemiluminescence analyzer (Xi’an Remax Electronic Science &Technology Co. Ltd., Xi’an, China) with the voltage of the photomultiplier
A three-electrode system was used, which consists of a screen printed carbon working electrode (SPCE), a carbon auxiliary electrode and an Ag/AgCl reference electrode (DropSens Corporation, Spain). A H600 transmission electron microscope (Hitachi, Japan) was employed to characterize the nanoparticles.

2.3. Preparation of RuL@SiO$_2$-Au Nanoparticles

RuL@SiO$_2$ nanoparticles were prepared according to the literature [16,28]. In brief, to a mixture of TX-100 (1.77 mL), cyclohexane (7.5 mL), n-hexanol (1.8 mL), 0.1 M Ru(bpy)$_3^{2+}$ (80 µL), tetraethylorthosilicate (TEOS, 100 µL) and water (340 µL) was added NH$_4$OH (60 µL, 25%). The mixture was stirred for 24 h, and the generated solid was isolated with acetone, sonicated for 10 min, followed by centrifuging and washing with ethanol and water. Then, the orange RuL@SiO$_2$ nanoparticles were collected and dried in vacuum oven. To prepare the RuL@SiO$_2$-Au nanoparticles, RuL@SiO$_2$ nanoparticles suspension in ethanol (6 mL, 2 mg mL$^{-1}$) was mixed with APTES (200 µL) and vigorously stirred for 30 min. The mixture was centrifuged and washed with water and ethanol to produce aminoterminated RuL@SiO$_2$ nanoparticles. A mixture of Au-colloid (2 mL) and APTES-modified RuL@SiO$_2$ suspension (1 mL) was then prepared and shaken for 1 h to let Au NPs be absorbed by the surface of the amino-terminated RuL@SiO$_2$ nanoparticles. The thus generated RuL@SiO$_2$-Au nanoparticles were collected by centrifugation, washed with water and suspended in water. The schematic graph of the fabrication process of RuL@SiO$_2$ and RuL@SiO$_2$-Au nanoparticles was shown in Scheme 1.

Scheme 1. Schematic illustration of the preparing procedures of RuL@SiO$_2$ nanoparticles and RuL@SiO$_2$-Au composite.

2.4. Preparation of RuL@SiO$_2$-Au~RuL-Ab$2$ and RuL@SiO$_2$-Au~Ab$2$ (RuL@SiO$_2$-Au Composite Nanoparticles Labeled AFP Secondary Antibody)

The RuL@SiO$_2$-Au~RuL-Ab$2$ bionanocomposite were prepared according to the literature [17]. In brief, a mixture of RuL@SiO$_2$-Au nanoparticles suspension and excess RuL-Ab$2$ was stirred for 24 h at 4 °C. The mixture was centrifuged and washed with water to obtain the RuL@SiO$_2$-Au~RuL-Ab$2$ bionanocomposite. The as-prepared bionanocomposite was incubated with BSA to block the unreacted and nonspecific sites, and then centrifuged and washed with PBS, dispersed in 0.1 M PBS (2 mL, pH 7.4) and stored at 4 °C for further use. The RuL@SiO$_2$-Au~Ab$2$ bionanocomposite was fabricated by using the same procedure, as shown in Scheme 2.
2.5. *Preparation of the SBAR Sandwich-Type Immunocomplexes*

The schematic of the fabrication process is shown in Scheme 3. The immunocomplexes were prepared as follows: a mixture of Biotin-Ab1 (50 µL), different concentrations of AFP (50 µL) and RuL@SiO₂-Au–RuL-Ab2 (50 µL) was prepared and held for 10 min at room temperature. After that, streptavidin-coated magnetic particles (SMP, 50 µL) were added into the mixture, allowing the RuL@SiO₂-Au–RuL-Ab2-AFP-Ab1-Biotin sandwich type immune-complexes to be captured on the surface of magnetic particles via biotin-streptavidin interaction. Then the sandwich-type immunocomplexes were isolated by a magnet, washed with PBST solution three times, dispersed in PBS (150 µL, pH 7.4) and stored at 4 °C for ECL tests.
2.6. ECL Measurements

The three-electrode system and ECL measurement process are shown in Scheme 4. For each test, sandwich-type immunocomplex solution (5 µL) prepared with different concentrations of target AFP was attached on the cleaned SPCE surface with a NdFeB permanent magnet, ECL measurements were then performed in PBS (30 µL, pH 7.4) containing $10^{-5}$ M TPA with a photomultiplier tube voltage of 800 V.

**Scheme 4.** (a) Schematic illustration of three-electrode SPCE system; (b) The process of ECL measurements; (c) the ECL measurement system.

3. Results and Discussion

3.1. Characterization of RuL@SiO$_2$ and RuL@SiO$_2$-Au Nanoparticles

In this work, $[\text{Ru(bpy)}_3]^{2+}$-doped silica matrix loaded with Au-NPs, named RuL@SiO$_2$-Au, was prepared as ECL signal amplification labels and immobilization substrates for AFP secondary antibody (Ab2). RuL@SiO$_2$ nanoparticles were first fabricated by using the well-established water-in-oil (W/O)
microemulsion method. Figure 1(A) shows the TEM image of RuL@SiO$_2$ nanoparticles with a uniform size distribution (~120 nm diameter). Incorporation of RuL molecules inside the silica matrix protects them from the surrounding environment, increases photostability and provides signal enhancement due to an increasing amount of RuL molecules doped per nanoparticle [29]. Furthermore, the ease of assembling functional groups such as amines, thiols and carboxyls on the surface of [Ru(bpy)$_3$]$^{2+}$-doped silica nanoparticles enables their use as ideal amplification labels for bioanalysis applications [30]. To immobilize AFP secondary antibody on the RuL@SiO$_2$ matrix, the surface of RuL@SiO$_2$ nanoparticles was aminoterminated with APTES and further reacted with Au-NPs. Figure 1(B) demonstrates that some individual Au-NPs (~16 nm diameter) and cluster-shape Au-NPs were successfully assembled on the surface of RuL@SiO$_2$ nanoparticles. These attached Au-NPs could provide a biocompatible, accessible matrix for immobilization of AFP secondary antibody.

**Figure 1.** TEM images of (A) RuL@SiO$_2$; (B) RuL@SiO$_2$-Au.

3.2. Optimization of Experimental Conditions

The ECL behavior of the sandwiched immunoassay was caused by the TPA and Ru(bpy)$_3$$^{2+}$. Thus TPA plays an important role in enhancement of the ECL signal. Furthermore, the enhanced ECL signal was related to the concentration of TPA. As can be seen from Figure 2, the ECL signal increased with increasing concentration of TPA. The effective enhancement occurred at the TPA concentration of $10^{-5}$ M. Thus 0.1 M of PBS containing $10^{-5}$ M TPA was selected for our measurements.

**Figure 2.** Effect of the different concentrations of TPA on the ECL intensity. Experimental parameters: initial potential = 0.0 V, high potential = 1.2 V, scan rate = 100 mV/s.
3.3. ECL Characterization of RuL@SiO$_2$-Au~RuL-Ab2

Prior to the use of RuL@SiO$_2$-Au~RuL-Ab2 as labels for the preparation of ECL immunosensors, we investigated the ECL performance of RuL@SiO$_2$-Au~RuL-Ab2 in the presence of TPA. For sandwich-type immunosensors, the sensitivity is mainly determined by the sensitivity of the label. In this work, the signal of the ECL immunosensor was mainly from the encapsulated RuL toward TPA oxidation. Since a large number of RuL molecules were incorporated into SiO$_2$ nanoparticles, and due to the intrinsic good electrocatalytic activity of RuL toward TPA oxidation, we hypothesized the sensitivity of the ECL immunosensor could be greatly enhanced when RuL@SiO$_2$-Au~RuL-Ab2 was used as recognition element of target AFP and signal amplification label. Figure 3 shows the different modified electrodes in PBS in the presence of $10^{-5}$ M TPA. It can be seen that only a small ECL intensity change was observed at the RuL-Ab2 modified electrode (curve c). On the contrary, a remarkable ECL intensity increase was observed for the RuL@SiO$_2$-Au~Ab2 modified electrode (curve b) and the largest ECL intensity response occurred at the RuL@SiO$_2$-Au~RuL-Ab2 modified electrode (curve a). These results indicated that a large amount of RuL molecules were incorporated into the silica matrix and RuL-Ab2 could be efficiently captured on the surface of RuL@SiO$_2$-Au nanoparticles.

![Figure 3. ECL-potential curves of, (a) SCMP/Biotin-Ab1/AFP/RuL@SiO$_2$-Au~RuL-Ab2; (b) SCMP/Biotin-Ab1/AFP/RuL@SiO$_2$-Au~Ab2; and (c) SCMP/Biotin-Ab1/AFP/RuL-Ab2 in pH 7.4 PBS containing $10^{-5}$ M TPA. Experimental parameters: initial potential = 0.0 V, high potential = 1.2 V, scan rate = 100 mV/s.](image)

3.4. Performance of the ECL Immunosensor

The highly sensitive label of the SBAR sandwich-type immunocomplexes was then used to construct ECL immunosensors for AFP detection. The double-conjugated Ru(bpy)$_3^{2+}$ sandwich-type immunocomplexes named Biotin-Ab1/AFP/RuL@SiO$_2$-Au~RuL-Ab2 were formed through antigen-antibody interaction in the presence of different concentrations of AFP. They were further interacted with streptavidin-coated magnetic particles (SCMP) and attached on SCPEs by magnet for ECL measurements. Then the ECL response of RuL and TPA was recorded in 0.1 M PBS (pH 7.4) containing $10^{-5}$ M TPA.
As shown in Figure 4, EI increased with the increasing of AFP concentration ranging from 0.05 to 100 ng mL\(^{-1}\). A linear relation between the logarithm of EI and the logarithm of AFP concentration was obtained [Log (ΔEI) = 3.3706 + 0.4075 Log (c_{AFP}/ng mL^{-1})] with a correlation coefficient R = 0.9962. The detection limit was 0.02 ng mL\(^{-1}\) (3σ). Such a low detection limit is better than those of previously reported AFP immunosensors. The comparison of results is shown in Table 1. The improved detection limit may be attributed to two aspects: (1) the high loading level of RuL molecules into the silica nanoparticles improved the detection limit; (2) the large amount of RuL-Ab2 absorbed onto the Au-NPs surface enhanced the access chance of the antibody-antigen interaction, especially when the AFP concentration is very low.

**Figure 4.** The schematic illustration of the ECL intensity versus the concentration of AFP (0.05 to 100 ng mL\(^{-1}\)) in 0.1 M PBS (pH 7.4) containing 10\(^{-5}\) M TPA. Insert: the relationship between Log of ΔECL signal towards log of different AFP concentrations. Experimental parameters: initial potential = 0.0 V, high potential=1.2 V, scan rate = 100 mV/s.

### Table 1. Comparison of analytical properties of various AFP immunosensors and immunoassays.

| Assay method            | Linear range (ng mL\(^{-1}\)) | LOD (ng mL\(^{-1}\)) | Detection antibody                          | Reference |
|-------------------------|--------------------------------|-----------------------|---------------------------------------------|-----------|
| Electrochemistry assay   | 1.00–500.0                     | 0.8                   | Au-alkaline phosphatase-labeled antibody    | [31]      |
|                         | 0.50–80.0                      | 0.25                  | HRP-antibody                                | [32]      |
| Voltammetric ELISA      | 0.10–200                       | 0.10                  | HRP-antibody                                | [33]      |
| Chemiluminescence       | 1.0–800                        | 0.23                  | HRP-antibody                                | [34]      |
| Photoelectrochemistry   | 0.05–50                        | 0.04                  | Label-free                                  | [35]      |
| Electrochemiluminescence| 0.05–50                        | 0.03                  | Ru-silica@Au-antibody                       | [17]      |
|                         | 0.05–100                       | 0.02                  | RuL@SiO\(_2\)-Au-RuL-antibody              | This work |

3.5. Specificity for the Detection of AFP

The selectivity of the immunosensor was also tested by adding possible interfering substances in the AFP-mediated sandwich-type immunoreaction. Different immunocomplexes were prepared with AFP (10 ng mL\(^{-1}\)) or AFP (10 ng mL\(^{-1}\)) together with the following individual interferent: carcinoembryonic antigen (CEA, 10 ng mL\(^{-1}\)), human IgG (HlgG, 1 μg mL\(^{-1}\)), carbohydrate antigen 19-9 (CA19-9, 10 ng
mL\(^{-1}\)), human chorionic gonadotropin antigen (HCG, 10 ng mL\(^{-1}\)), BSA (1 µg mL\(^{-1}\)), ascorbic acid (AA, 1 µg mL\(^{-1}\)), dopamine (DA, 1 µg mL\(^{-1}\)) and L-lysine (LL, 1 µg mL\(^{-1}\)). The interference degree was evaluated by comparing the ECL intensity of a mixture of AFP and interfering substance with that of AFP alone. As can be seen from Table 1, less than 5% variation of log(EI) responding to 10 ng mL\(^{-1}\) AFP was observed in the presence of different interferents (Figure 5), which demonstrated a good selectivity of the developed ECL immunosensor for AFP detection.

Figure 5. Selectivity analysis of the ECL immunosensor in the presence of different interferents. The concentrations of the interferents were: CEA (10 ng mL\(^{-1}\)), HIgG (1 µg mL\(^{-1}\)), CA19-9 (10 ng mL\(^{-1}\)), HCG (10 ng mL\(^{-1}\)), BSA (1 µg mL\(^{-1}\)), DA (1 µg mL\(^{-1}\)), L-lysine (1 µg mL\(^{-1}\)).

3.6. Determination of AFP in Human Serum Samples

In order to investigate the possible application of this immunosensor in clinical analysis, recovery experiments were performed by standard addition methods in human serum. The experimental results were shown in Table 2 and the recovery was in the range from 95% to 110%, which indicated that the developed sensor might be applied for the determination of AFP in human serum for routine clinical diagnosis.

| Sample number | Added (/ng mL\(^{-1}\)) | Found (/ng mL\(^{-1}\)) | Recovery (/%) |
|---------------|-------------------------|-------------------------|--------------|
| 1             | 0.50                    | 0.51 ± 0.02             | 102.0        |
| 2             | 1.00                    | 0.97 ± 0.02             | 97.0         |
| 3             | 5.00                    | 5.28 ± 0.2              | 105.6        |
| 4             | 10.0                    | 9.8 ± 0.3               | 98.0         |
| 5             | 50.0                    | 47.6 ± 2.3              | 95.2         |

\(^1\) Mean value ± SD of three measurements.

3.7. Regeneration

The magnetic probe (SBAR) can be modified and removed from its surface by magnetic field added on the flat bottom of SPCEs. All these steps can renew the electrode surface and make the electrode
reusable. As Figure 6 showed cyclic voltammetry (CV) curves of different electrodes (a, bare SPCEs; b, the immunocomplexes were immobilized on the working electrode of SPCEs surface; c, after renewing of SPCEs.) in 1 mM K$_3$Fe(CN)$_6$ (0.1 M KCl), well-defined CVs, characteristic of diffusion-limited redox processes, are observed at the bare SPCEs [Figure 6(a)]. The peak currents decreased [Figure 6(b)] after SBAR sandwich-type immunocomplexes were immobilized on the working electrode of SPCEs surface because the immunocomplexes would block the electron transfer. After removing the magnet and rinsing the sensor with PBST, the currents was further increased [Figure 6(c)] almost black to bare SPCEs, due to the bulk of immunocomplexes were rinsed out.

**Figure 6.** Cyclic voltammograms performed in 1 mM Fe(CN)$_6^{3−}$ solution (0.1 M KCl) at 100 mV s$^{-1}$ for (a) bare; (b) SBAR sandwich-type immunocomplexes; (c) removing the magnet and rinsing with PBST.

4. Conclusions

In this paper, a magnetic probe (SBAR) consisting of streptavidin-coated magnetic particles (SCMP) and RuL@SiO$_2$-Au~RuL-Ab2 complexes is used to detect AFP as a model protein. SCMP were used as supporting material for the preparation of the sandwich-type immunocomplexes. A magnetic separation step was then used to isolate the complexes from the unbound components, considerably reducing incubation and washing times. Furthermore, the SBAR sandwich-type immunocomplexes can be modified and removed from its surface by magnetic field added on the flat bottom of SPCEs, therefore, the electrodes, unlike traditional electrochemical immunosensors, do not require complex modification and cleaning steps. In addition, the RuL@SiO$_2$-Au~RuL-Ab2 composite exhibited dual amplification since SiO$_2$ could load large amount of reporter molecules (RuL) and gold particles could provide large active surface to load more reporter molecules (RuL-Ab2). Thus the immunosensor can simultaneously realize separation, enrichment and determination, with high sensitivity, which would be valuable for clinical immunoassay for AFP in human serum.
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