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Enhanced Sensitivity of Two-Dimensional Bar-Codes Microfluidic Paper-Based Device (µPADs) By Gold Nanoparticles

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Abstract

Microfluidic paper-based devices ($\mu PADs$) has been studied extensively for rapid detection. However, the sensitivity of the detection are sacrificed for the rapid detection, which might be promote by plasmonic gold nanoparticles. In this paper, we demonstrated the $\mu PADs$ s $\mu PADs$ with Au nanoparticles integrated which could be used as a multi-metallic ion sensor for point-of-care testing. By combining the fast, accurate, convenient, and direct merits of a smartphone-based colorimetric readout with the designed $\mu PADs$ device, we could obtain relevant information such as the type of ion and relative concentrations. We optimized the information area, outline geometry, and reduction rate of $\mu PADs$ information areas. The metallic ion concentrations could be determined in the linear dynamic range of $1\sim20$ mg L^{-1} for Al^{3+} and $2\sim20$ mg L^{-1} for Cr^{6+} and Zn^{2+} . The proposed $\mu PADs$ sensor can achieve quantitative detection of metallic ion which was found to be highly sensitive for metallic ion detction against non-AuNPs. In addition, the prepared sensor was successfully applied to assay metallic ion in real environmental samples.

Keywords: Gold Nanoparticles; Paper-Based Devices; Multi-Metallic Ions Detection

Introduction

Point of care testing (POC) has a unique analytical detection advantages, which is widely used in infectious diseases [1,2], food safety [3], environmental monitoring [4]. For the first time presented by Whitesides research group in 2007 [5], paper as a substrate to construct microfluidic paper-based devices (μ PADs) has been studied extensively for rapid detection [6,7]. The fluid that driver by capillarity wick in two or three dimensions, which play a very important role in point of care (POC) diagnostic testing [8]. Colorimetric assay has experienced rapid development, which is expected to achieve each person's low cost for POC diagnosis testing [9]. Towards accurate quantification, several groups have tried various optical detector scanning10, cameras [11], smart phones [12] to analyse colorimetric detection. Although colorimetry can achieve quantitative analysis [13], this visual detection rely on the colour intensity or brightness on paper, which often bring a lot of uncertainty [14].

As the increasing of using smartphone based biosensor, potentially, the smartphone based detection technology provides an ideal platform for the rapid detection technology [15-17]. Meanwhile, the two-dimensional (2D) bar code is prevalent used in daily life in terms of payment, browsing the web and so on. Thus, in this approach, we utilized the complex motif of two-dimensional code to construct the hydrophobic and hydrophilic barriers, which can be scanned by smart phones. This new colorimetric analysis is kind of method using distance-based detection in paper [18]. In the distance-based detection principle, the analyte reacts with color agent and grows along the flowing stream line until all the analyte is depleted. Distance-based measurement for paper analytical devices has been successfully achieved in the horizontal direction of the quantitative analysis of heavy metals and glucose [15,19]. These distance detection methods have limitation in practical applications. First, in the horizontal direction distance method detection, color agent and analyte need more order of magnitude for a longer period of time to complete detection and the volatile liquid in the process will be faced. Second, more color products will be gathered in the fluid terminal that makes the distance detection in directly by naked eye observation appear larger error and difficult to identify.

There is growing explosion in the broadband light absorbers because of their importance in various application, such as surface-enhanced spectroscopy, solar energy conversion, photothermal therapy, and fabrication of novel optoelectronic devices. To improve the detection sensitivity, our new designed 2D bar code PADs with gold nanoparticles integrated which have the advantages to minimize the error which caused by the long distance on the horizontal channel. We proposed a convenience detection method that the result of a colorimetric detection could be obtained by scanning a 2D barcode via a smartphone. In this study, we also optimized the surface area, geometry, and reduction rate of $\mu PADs$ information areas for the real-time, effective, and accurate detection of metallic ions (Al³+, Cr²+, and Zn²+).

Experimental

All chemicals were used without any further purification and ultrapure water purified with a Mill-Q system (18.2 M Ω cm) from Merck Millipore (Darmstadt, Germany). Ammonium aluminium sulfate dodecahydrate, zinc nitrate hexahydrate, chrome azurol S, zincon disodium salt, 1,5–diphenylcarbazide (DPC, acetone, mercury(II) sulfate, iron (III) chloride anhydrous, copper (II) sulfate pentahydrate, sodium acetate, acetic acid glacial and manganese sulfate anhydrous were purchased from Sinopharm Chemical Reagent Company (Shanghai, China). Potassium dichromate was purchased from Guangzhou Jinhuada Chemical Reagent Company (Guangzhou, China). Lead nitrate and magnesium sulfate anhydrous were purchased from Aladdin (Shanghai, China). Nickel (II) nitrate was purchased from Enox (Jiangsu, China). Qualitative-grade filter paper was purchased from the Xinhua paper company (Hangzhou, China). Hydrophobic wax was used as barrier for all detection types. Adobe illustrator software (Crative Cloud 2014) was used for the design of the patterns. The μ PADss were patterned by a commercially available wax printer (FUJIXEROX Phaser 8560DN).

The overall design concept and operation flow chart is shown in Figure 1. The fabrication procedure involves using Adobe illustrator to design the information area and hydrophilic sample reservoirs, each analytical device were sized as $20 \times 20 \text{ mm}^2$. Then, the hydrophilic-hydrophobic contrast of patterns were printed on the appropriate sized filter paper using a commercial wax printer. The multiple information of the two-dimensional barcode that is composed of hydrophobic wax and hydrophilic reagents' reservoirs. Three white zone (Figure 1) which is the hydrophilic zone set as the testing area at the centred accommodate sample addition. We used commercial $\mu PADs$ software to scan and the information showed in the phone screen (Figure S1).

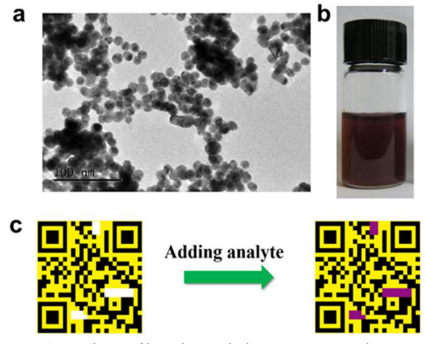


Figure 1: Operational concept of the two-dimensional code assay. Wax printing is simple, inexpensive and fast to fabricate the paper analytical devices. (a) TEM image of gold nanospheres with 15nm, the scal bar is 100nm; (b) The digital photo is a suspension of Au nanoparticles, the scal bar is 2cm (c) Schematic diagram and process flow of the $\mu PADss$ paper-based device

Results and Discussion

As shown in Figure 2, we provided six separated $\mu PADs$ with different amount of Au nanoparticles (AuNPs) suspension dropped on the three rectangle reservoirs, the capillary action allowed the analyte to migrate quickly in the formed microchannel. The amount of the au nanoparticles are measured with weight percent of the suspension from L0 to L5. Finally, all the $\mu PADs$ were separated sequentially for easy recognition which were scanned by using a smartphone. Compared to $\mu PADss$ with non-AuNPs of L6, the sensitivity of the metal ion detection could be found by the intensity of the colorimetric reactions.

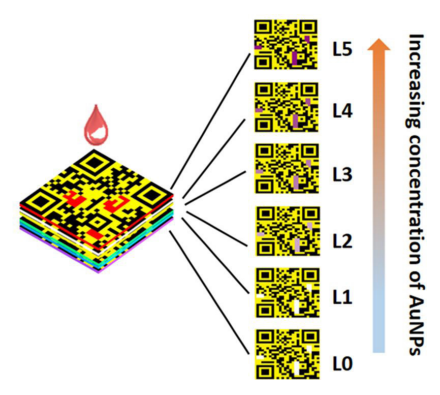


Figure 2: Six separated μ PADs device. The separated six layers were denoted as L0, L1, L2, L3, L4, L5 (L5=1.0%, L4=0.6%, L3=0.5%, L2=0.2%, L1=0.1%, L0=0) The μ PADs for the quantitative analysis was scanned by a smartphone

The recognition of μ PADs in balance with the size of the information area needs to consider. It is shown in Table S1 that the percentage of the information area with gold nanoparticles compared to the total surface of the μ PADs (%). We summarized the recognition capability of the smartphone depending on the different dimensions of the μ PADs. To rapidly identify the μ PADs information resulting from colorimetric reactions, the identification process was as follows. First, the μ PADs was erased by increasing information area ratio (> 5.3%, information area of the total surface of the μ PADs as shown in Table S1). Second, the hydrophilic blank area containing the colorimetric reagents and the analyses reacted in the reaction region and the resulting products gradually covered the information area and the μ PADs could be recognized by the smartphone. Finally, the sensitivity of the metal ions could be enhanced by the amount of AuNPs which could be recognized by the smartphone.

Wax printing appeared to be a simple and inexpensive method for the fabrication of $\mu PADs$ [19]. The process involves the penetration of the wax pattern into the paper sheet by heat to form the hydrophilic and hydrophobic areas of the PADs. The heating step could lead to the shrinkage of the $\mu PADs$ and could impede the recognition by the smartphone. In order to evaluate the influence of the heating step, we compared the reduction rate of $\mu PADs$ information areas presenting an outline geometry of 20 x 20 mm² and prepared with and without heating. Wax spread inside the porous filter paper due to the capillary action characterized by the Washburn's equation (Eq. 1) [20]:

with t as the heating time; η as the viscosity of the liquid; γ as the surface tension; L as the flow distance, and D as the average pore diameter of the porous material.

$$L = \sqrt{\frac{\gamma Dt}{4\eta}} \tag{1}$$

The results are depicted in the graphic shown in Figure 3. The average reduction rate of μ PADs information areas was 47.9% and the relative standard deviation value (RSD) for all measurements (n=5 measures for each different μ PADs information area) was 9.3%. When the information area was too small, the bar-code information could not be recognized because the hydrophilic region was blocked by the melted wax.

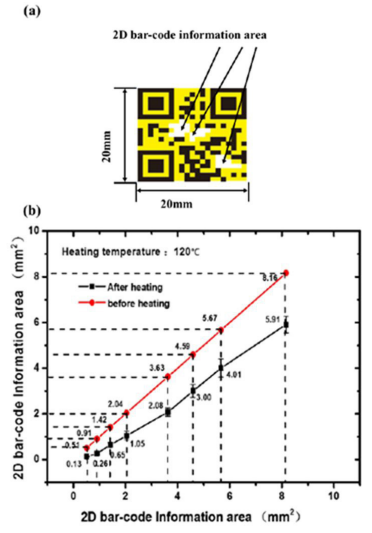


Figure 3: (a) μ PADs information area with an outline geometry of 20 x 20 mm². (b) The relation between the information area without and with heating process. The average reduction rate of μ PADs information areas was 47.9% (n=5)

To clarify the sensitive capability of the μ PADs, Figure 4 gives the resonances intensity various among different media or solution for L6 of μ PADs. It is interesting that the variation of the resonances for the peak is tending to approach as the value increasing of different metallic ion solution. To further investigate the μ PADs sensing capabilities of non-AuNPs and with AuNPs, the solution with different concentration was chosen and injected into 2D-bar codes based microfluidic channel (Table S2, ESI).

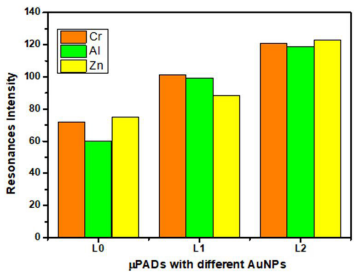
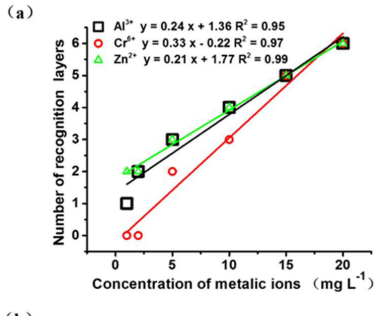


Figure 4: The resonant intensity various among different media or solution for Peak of $\mu PADs$

In order to evaluate the ability of the μ PADs to detect metal ions quantitatively, we used a distance-based detection in the vertical direction [21]. As the distance-based detection presented a nonlinear fluid velocity-dependent, the colorimetric products responded non-linearly to the variation of distance in the vertical direction [22,23]. We developed a specific layer deposition method for Figure 4. The resonant intensity various among different media or solution for Peak of μ PADs dropping the colorimetric reagents on the paper surface to counter this non-linear response. Once the information area was completely dried, a single aliquot of the analyte was dropped on the hydrophilic information area of the first layer for the complete analysis of the sample. The colorimetric reagent reacted with the metallic ions of the sample to form colorimetric products, which were also on the paper surface of the 2D-bar codes. The concentration of the colorimetric products distributed continuously along the top to the bottom layers via micro channels.

The colorimetric reagent accumulation gradually became less pronounced from top to bottom as the number of total layers increased. The results are depicted in Figure S2 and the relative standard deviation (RSD) for all measurements (n=5 measures for each different stacked layers) was of 5.0%. To maximize the detection limit, we considered the highest number of 2D-bar codes stacked layers available. The flow rate inside the microchannel would be close to zero (when the number of layers is greater than 8) and the colorimetric products would stop being generated according to the Lucas Washburn theory. Nevertheless, the recognition layers with colorimetric reagents could be enhanced if the colorimetric reagent dosage and sample rate increased. However, exceeding a certain threshold, the colorimetric reagents would destroy the hydrophilic μ PADs information area. The results indicated that the minimum path length (ca. 900 μ m) to be travelled by the fluid inside the device corresponded to the stacking of six layers. Therefore, the subsequent experiments were performed on 6-layers 2D-bar codes.



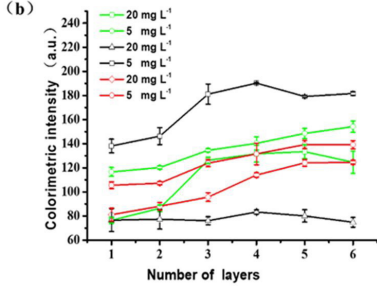


Figure 5: The recognition ranges of the 6 layer-device were 1 - 20 mg L^{-1} , 2-20 mg L^{-1} , 2-20 mg L^{-1} for Al^{3+} , Cr^{6+} and Zn^{2+} metallic ions, respectively. (b) The colorimetric intensities were used for the detection of Al^{3+} , Cr^{6+} and Zn^{2+} metallic ions and the concentrations were 5, 5, and 20 mg L^{-1} , respectively

In Figure 5a, μ PADs devices were tested using the same analyte range (1~20 mg L⁻¹) for all three respective metal ions (Al³+, Cr⁶+, and Zn²+). The experimental results indicated that the concentration of metallic ions could be quantified by counting the number of the layers recognized by the smartphone. The recognition linear dynamic range of the Al³+ metallic ions was 1-20 mg L¹, while the one of the Cr⁶+ and Zn²+ metallic ions were 2-20 mg L¹. The limits of detection of Al³+, Cr⁶+, and Zn²+ were 1, 2, and 2 mg L¹, respectively. These values are similar to the ones referred in the World Health Organization (WHO) guidelines [24,25]. Overall, the data clearly showed that the μ PADs detection of the metallic ions was possible with a smartphone (Figure 5a) without having to use a complex image analysis step for quantification. This proof of concept opens new avenues for the use of μ PADs paper-based devices in many fields such as the direct in-field detection of metallic ions by avoiding the need for sophisticated and expensive instrumentation located in a central laboratory that is commonly required for further quantitative analysis [26,27].

To demonstrate the reliability of the detection, we determined the relation between the colorimetric intensity and the recognition rate as summarized in Figure 5b. As the colorimetric reagents and metal ions reacted, colored products were formed in the hydrophilic information area. Due to the different recognition rates of each metallic ions, appropriate concentrations had to be chosen in order to analyze the colorimetric intensity of the sample. We decided to use a maximum concentration of 20 mg L⁻¹ and a minimum concentration of 5 mg L⁻¹ for the Al³⁺, Cr⁶⁺ and Zn²⁺ metallic ions to evaluate the number of layers that could be recognized by a smartphone (the RSD were 4.3%, 6.8%, and 3.2%, respectively, n=3). For the same metallic ion concentration, the colorimetric intensity increased together with the number of layers. And for the same number of layers in μ PADs devices, the colorimetric intensity increased as the metallic ion concentration increased. As mentioned above, the concentration of the colorimetric products could gradually decreased from the top to the bottom layers, reducing the number of layers that could be recognized by the smartphone. At a metallic ion concentration of 5 mg L⁻¹ and under optimal colorimetric intensity and recognition condition of the μ PADs in each layer, the measurement of the colorimetric intensity was above 140 (Al³⁺), 150 (Cr⁶⁺), and 140 (Zn²⁺), and the number of recognition layers were 4, 2, and 3, respectively. Paper based μ PADs detection of Al³⁺, Cr⁶⁺, and Zn²⁺ barely interfered by other ions, which also present in Figure S3.

Conclusion

In summary, we designed and fabricated $\mu PADs$ integrated with gold nanoparticles for multi-metallic ions real-time detection. We are able to simultaneously quantify the concentrations of three different metallic ions with high accuracy, and the limits of detection of Al^{3+} , Cr^{6+} , and Zn^{2+} were 1, 2, and 2 mg L^{-1} , respectively. We demonstrated a rapid method for the detection of metallic ions without the use of additional device, apart from the $\mu PADs$ and a smartphone. The high portability, applicability, and cost-effectiveness of the $\mu PADs$ detection approach would overcome the time and space limitations required in real-time diagnostic tests for environmental monitoring, food safety, and other fields.

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Electronic Supplementary Information (ESI)

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