Biomolecule detection via target mediated nanoparticle aggregation and dielectrophoretic impedance measurement

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A new biosensing system is described that is based on the aggregation of nanoparticles by a target biological molecule and dielectrophoretic impedance measurement of these aggregates. The aggregation process was verified within a microchannel via fluorescence microscopy, demonstrating that this process can be used in a real time sensor application. Positive dielectrophoresis is employed to capture the nanoparticle aggregates at the edge of thin film electrodes, where their presence is detected either by optical imaging via fluorescence microscopy or by measuring the change in electrical impedance between adjacent electrodes. The electrical detection mechanism demonstrates the potential for this method as a micro total analysis system (μTAS).

Introduction

The analysis of biological molecules has become increasingly important over the past few years, due in part to the potential threat that deadly toxins, viruses, and bacteria pose as warfare agents. Consequently, the science and engineering of detection, identification and quantification of small quantities of target biomolecules are rapidly expanding fields of endeavor.

In this paper, a sensing mechanism for biomolecules that is based on nanoparticle aggregation is presented. This method is demonstrated to be capable of detecting nanomolar concentrations of a target biomolecule by the dielectrophoretic impedance measurement method. The positive dielectrophoretic force is utilized to capture suspended nanoparticle aggregates onto an interdigitated, electrode array. As a proof of principle, the aggregation of biotinylated nanoparticles in the presence of avidin and their collection by dielectrophoresis (DEP) within a microchannel are demonstrated as the basis of an automated, biomolecular detection microsystem.

Principal mechanisms

Dielectrophoresis (DEP) is the movement of polarizable particles in non-uniform, alternating electric fields.1 The time-dependent DEP force for a spherical particle suspended in an electrolyte is given by

\[
<F_{\text{DEP}}> = 2\pi a^3 \varepsilon_m \Re[f_{CM}] |\Delta \varepsilon_{\text{real}}|^2
\]

(1)

where \(E_{\text{rms}}\) is the root mean square (rms) value of the electric field, \(a\) is the radius of the particle and \(\varepsilon_m\) is the real part of the complex permittivity of the suspending medium; \(f_{CM}\) is the Clausius–Mossotti factor and is given by

\[
f_{CM} = \frac{\varepsilon_p - \varepsilon_m}{\varepsilon_p + 2\varepsilon_m}
\]

(2)

where \(\varepsilon_p\) and \(\varepsilon_m\) are the complex permittivities of the particle and the medium, respectively. For a dielectric material, the complex permittivity is,

\[
\varepsilon*(\omega) = \varepsilon - j\frac{\sigma}{\omega}
\]

(3)

where \(j = \sqrt{-1}\), \(\omega\) is the angular frequency of the electric field, \(\varepsilon\) is the permittivity and \(\sigma\) is the conductivity.

From eqn. (1) to (3), it is readily apparent that the DEP force is a function of particle size, the electric field gradient, the frequency of the applied signal, and the dielectric properties of the materials. The positive DEP force, which collects the nanoaggregates at the electrode edges, is achieved when the real part of \(f_{CM}\) is positive; while the negative DEP, which pushes the nanoaggregates away from the electrode edges, occurs if \(f_{CM}\) is negative.

In this paper, positive DEP forces are used to selectively capture and concentrate nanoparticle aggregates containing a target biomolecule, over individual nanoparticles. This is possible because the aggregates only form in the presence of the target and because of the dependence of the force magnitude on particle radius cubed. Alternatively, by preparing nanoparticles of a different composition, negative DEP can be achieved and used to levitate and transport aggregates to a collection/detection site.

Fluorescent nanoparticles are attractive alternatives to organic dyes as biological labels due to their stability, resistance to photobleaching and tunability of photoemission. Additionally, their surfaces can be conjugated with biologically active small molecules, DNA, and proteins with specific binding properties as well as polymers that are inert to biological adhesion.5–8 As part of a detection scheme, they provide a robust platform for chemical/biological information and signal detection.

In prior work, it was demonstrated that nanoparticles with surfaces that were modified with a ligand for proteins (biotin) would aggregate in the presence of the complementary protein (avidin). This process emulates the well known Latex11 agglutination test,9 used in a number of commercially available...
immunoassays. The nanoparticles (Fig. 1, top) were composed of a 40 nm diameter CdS@SiO$_2$ inorganic core, surface-grafted with water solubilizing poly(ethylene glycol) (PEG) chains via a monosiloxane linkage. The PEG chain ends carried a biotin group (Fig. 1, bottom left). The kinetics of aggregate growth rate were studied using dynamic light scattering (DLS), and the rate of aggregation was shown to depend upon the nanoparticle surface biotin content and the initial concentration of avidin. Under optimal conditions, micron sized aggregates formed within several minutes (Fig. 1, bottom right), promoted by avidin concentrations down to nanomolar levels.

The dielectrophoretic impedance measurement (DEPIM) method was reported by Suehiro for bacteria detection. This method utilizes positive dielectrophoresis to attract polarized particles to the edges of an interdigitated electrode array, resulting in an end-to-end alignment of particles that is similar to a pearl chain. If the pearl chains bridge the electrode gap, the admittance between the electrodes increases. By monitoring the change in impedance, the bacteria can be detected and its population can be evaluated by the magnitude of the change. In this paper, the DEPIM method is employed to detect the presence of collected nanoparticle aggregates.

Materials and methods

Materials

All materials were purchased from commercially available sources and used without further purification. Depending upon the detection method, some of the chain ends of the surface grafted PEGs were conjugated with functional groups in addition to biotin. Due to the low quantum yields of silica coated CdS nanoparticles prepared using the reverse micelle method, a fluorescent dye, 5-fluorescein isothiocyanate (FITC), was attached to some of the chain ends to enhance the fluorescence intensity of the particles. This procedure has been described previously. For another set of particles, some of the PEG chain ends were conjugated with ferrocene carboxylic acid in order to impart redox activity to the particles. The N-hydroxysuccinimide ester of carboxyferrocene was prepared and allowed to react with some of the amine-terminated PEGs on the particle surface to yield ferrocene carboxamide chain ends. The reaction was allowed to proceed for 24 h, and the particles were then isolated by five centrifuge/decant/resuspend cycles with CH$_3$OH/hexane/THF (1 : 1 : 3).

CV of ferrocene-containing nanoparticles

Electrochemical measurements were performed using a PAR model 263 potentiostat/galvanostat using a gold film working electrode, a silver wire counter electrode and 0.1 M (nBu$_4$N$^+$)(PF$_6$$^-)$ supporting electrolyte in dry acetonitrile solution. Potentials were recorded vs a standard calomel reference electrode (SCE). The solution was purged with N$_2$ for 10 min prior to recording the voltammogram. Functionalized nanoparticles were deposited onto the gold electrode to produce a thin film. Free ferrocene was added to the solution of each experiment as an internal standard.

Fabrication of DEP electrodes

A standard, lift-off process was used to pattern the metal electrodes, as described below. The electrode area was photolithographically defined on the surface of standard, borosilicate glass microscope slides. Then, a 150 Å chrome adhesion layer was deposited, followed by the deposition of a 150 nm thick platinum layer by sequential e-beam evaporation in a single pump-down. The resulting length and width of the electrodes and the gap between two adjacent electrode edges were 1.8 mm, 50 μm and 5 μm, respectively.

Fabrication of a DEP microchannel

A convenient way to construct sealed and optically clear chambers for dielectrophoretic applications has been developed using SU-8, the details of which are to be published elsewhere. Briefly, the process starts by photolithographically defining an open channel, 50 to 100 μm wide, in a 20 μm thick layer of SU-8 directly above the electrode array, then mechanically drilling 1 mm diameter, inlet and outlet ports into the glass slide, and finally bonding a thin (100 μm) glass cover on top, using a thin, intermediate layer of prebaked SU-8 (about 3 μm). After exposure through a mask, the unexposed, intermediate SU-8 is developed from the glass surface within the chamber, leaving a clean glass surface for optimal transparency. Tubing and electric wiring are then connected for supplying solution and electric signal, respectively.

DEPIM equipment and test setup

The dielectrophoresis channel and electrodes were connected to peripheral measurement, imaging and actuation instruments.
was spiked. The formation of the redox wave at 668 mV increased after the solution of the electrolyte solution. In this third experiment, the intensity of the functional groups. Confirmation that the new signal is higher than that of ferrocene due to the electron withdrawing nature of carboxylate derivatives of ferrocene is known to be higher.

The CV of the particle film containing the ferrocene carboxamide chain ends showed two redox waves at 448 and 668 mV. The oxidation potential of carboxylate derivatives of ferrocene is known to be higher than that of ferrocene due to the electron withdrawing nature of the functional groups. Confirmation that the new signal at the higher oxidation potential corresponded to that of the conjugated ferrocene derivative came from a repetition of the experiment with ferrocene carboxylic acid added to the electrolyte solution. In this third experiment, the intensity of the redox wave at 668 mV increased after the solution was spiked.

Results and discussion

CV of redox active nanoparticles

Confirmation of the presence of ferrocene groups on the nanoparticle surface via elemental analysis or UV/Vis spectroscopy failed due to the small number of ferrocene groups per particle and light scattering from suspensions of the particles. However, cyclic voltammetry (CV) proved to be effective. A film of derivatized nanoparticles was deposited onto a gold film working electrode, and the film was submerged into an electrolyte solution containing ferrocene as an internal standard. Fig. 3 shows the results of three experiments. In the control experiment, the electrode film contained particles without carboxamide chain ends. The corresponding voltammogram only showed a redox wave corresponding to the ferrocene internal standard at 448 mV. The CV of the particle film containing the ferrocene carboxamide chain ends showed two redox waves at 448 and 668 mV. The oxidation potential of carboxylate derivatives of ferrocene is known to be higher than that of ferrocene due to the electron withdrawing nature of the functional groups. Confirmation that the new signal at the higher oxidation potential corresponded to that of the conjugated ferrocene derivative came from a repetition of the experiment with ferrocene carboxylic acid added to the electrolyte solution. In this third experiment, the intensity of the redox wave at 668 mV increased after the solution was spiked.

Microchannel experiments

When suspensions of biotinylated nanoparticles ([nanoparticle]_0 = 0.15 mg mL⁻¹) were mixed with solutions of avidin ([avidin]_0 = 10 to 1500 nM), in a mL sample volume, large nano- and micron-sized aggregates resulted, as verified by SEM images of dried samples (Fig. 1). In order to demonstrate and verify that the aggregation process would also take place within the <100 nL volume of the microchannel, fluorescence microscopy was utilized. Although individual nanoparticles are smaller than the limit of resolution of the microscope camera and are therefore unresolvable, the large aggregates are clearly visible due to the high concentration of the FITC within those structures. Also, the microchannel confines the nanoparticles to a relatively small region and volume, which greatly simplified locating the aggregates within the small field of view of the microscope at 400 × magnification.

Before each fluorescence imaging experiment, the microchannel and reservoirs were cleaned and checked in the microscope to ensure the absence of fluorescence from contaminants or other artifacts. First, premixed solutions of avidin (100 nM) and the nanoparticles were added to the microchannel. After several minutes, bright fluorescent spots, of several micron radii, appeared, randomly dispersed within the microchannel. In order to verify that the appearance of the bright fluorescent spots was caused by avidin promoting the formation of aggregates, a laminated flow experiment was carried out. Two syringes and a hand driven syringe pump were attached to the inlet reservoir of the microchannel. One syringe contained biotinylated nanoparticles dispersed in water while the other contained an aqueous solution of avidin. The two solutions were introduced simultaneously into the microchannel, forming laminating flow columns as shown in Fig. 4,
The microchannel was continuously imaged during the experiment using the fluorescence microscope. The flow was stopped, allowing the two solutions to mix at their interface by diffusion. After approximately ten minutes, bright, luminescent spots appeared along the center of the microchannel. A photomicrograph of the luminescent spots inside the microchannel is shown in Fig. 4, bottom.

After demonstrating aggregation within a microchannel, a series of experiments were performed to demonstrate collection of the aggregates with DEP. Preliminary DEP experiments used solutions of pre-formed nanoaggregates. The solutions were injected into the microchannel at a flowrate of 40 \( \mu \text{L min}^{-1} \) using a syringe pump while a DEP signal of 5 volts ac at 1 MHz was applied. After six minutes, aggregates were observed to collect at the electrodes via fluorescence microscopy (Fig. 5A and 5B).

While the ligand/protein system of biotin and avidin provide excellent proof of principle, more noteworthy analytes are envisioned. For example, the conjugation of antibodies and proteins to the nanoparticle would allow for aggregation to be induced by either proteins, antibodies, or small molecules. Therefore, a series of systems were explored to exploit the versatility of this sensing mechanism.

First, aggregation of biotinylated CdS@SiO\(_2\) nanoparticles was induced with anti-Biotin IgG antibodies to demonstrate the ability to use antibodies. Next, avidin and anti-Biotin antibodies were independently conjugated to SiO\(_2\) nanoparticles. A water-soluble bis-biotin compound was prepared.
to serve as a small molecule analyte. Upon introduction of the small molecule analyte to a suspension of SiO$_2$ particles conjugated with either avidin or anti-Biotin IgG antibodies, aggregation was induced. Although each system has not yet been tested in the DEPIM apparatus, it can be concluded that this method provides sufficient versatility to be applied to a variety of analytes.

A final point to be addressed is the ultimate sensitivity of this device. Preliminary DEPIM experiments employed analyte concentrations that were determined from dynamic light scattering (DLS) experiments to yield ideal aggregation conditions. In testing the ability for other analytes to induce aggregation, DLS results indicated that the analyte concentration needed to induce aggregation varied from system to system, ranging from as high as 300 nM to as low as 800 pM. Therefore, further testing is necessary in order to optimally apply this technique to different analytes.

In addition, the range of analyte concentration which can induce aggregation can be expanded. Initial DLS experiments that used 40 nm particles resulted in an analyte concentration range from 50 to 200 nM. By using formulations of particles of various sizes within the aggregation process, the analyte concentration range for DLS experiments was expanded to 10–500 nM.

One problem encountered with the preliminary system was the precipitation of large aggregates from solution. Therefore, PEG linkers of large molecular weight were prepared to improve the dispersability of larger aggregates. The kinetic growth mechanism was not altered, which allows for direct implementation of this improved system.

Conclusions

Selective, biomolecule detection by the target mediated aggregation of nanoparticles was demonstrated using a model system of biotinylated nanoparticles and avidin. Formation of the aggregates within a microchannel of volume less than 100 nL was conclusively demonstrated in a laminated flow experiment. Aggregates were dielectrophoretically captured between thin film electrodes residing inside a microchannel and detected by both fluorescence microscopy and a change in interelectrode impedance. Further study is required to determine the detection limit capability of this system. Based on DLS analysis of different analyte systems and the utilization of nanoparticles of various sizes and longer PEG linkers, the formation of aggregates at picomolar concentrations is deemed plausible. It remains to be determined how few aggregates need to be formed in order to detect their presence by DEPIM. Smaller electrode spacing would require fewer aggregates to be formed for detection, and would also lower the detection limit. Finally, work is currently being completed to prepare nanoparticles with higher quantum yield photoemission than CdS@SiO$_2$ nanoparticles, i.e. water-soluble CdSe.

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References