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Surface Plasmon Resonance Detection Using Antibody-Linked Magnetic Nanoparticles for Analyte Capture, Purification, Concentration, and Signal Amplification

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Rapid, sensitive, and accurate detection of analytes present in low concentrations in complex matrices is a critical challenge. One issue that affects many biosensor protocols is the number and nature of the interferences present in complex matrices such as plasma, urine, stool, and environmental samples, resulting in loss of sensitivity and specificity. We have developed a method for rapid purification, concentration, and detection of target analytes from complex matrices using antibody-coated superparamagnetic nanobeads (immunomagnetic beads, or IMBs). The surface plasmon resonance (SPR) detection signal from staphylococcal enterotoxin B (SEB) was dramatically increased when the IMBs were used as detection amplifiers. When SEB detection included a 10-fold concentration/purification IMB protocol, a substantial increase in detection sensitivity was observed. This procedure was used to successfully purify and concentrate SEB from serum and stool samples, then amplify the SPR detection signal. SEB at a concentration of 100 pg/mL was easily detected in both buffer and stool samples using this procedure. The IMB protocol also served to verify the analyte detection by using two different anti-SEB antibodies, mouse monoclonal antibodies attached to the magnetic nanobeads and rabbit polyclonal antibodies on the SPR sensor surface. Multiple detections of SEB in stool were performed using the same sensor surface by regenerating the sensor surfaces with a pH 2.2 buffer wash.

Surface plasmon resonance (SPR) biosensors, when derivatized with highly specific recognition elements, provide powerful tools for rapidly determining the presence and concentration of analytes in solution or suspension. Label-free detection and near-real-time analysis, coupled with the recent development of small, easy to use portable instruments make SPR biosensor systems excellent candidates for point-of-care detection devices, environmental monitoring systems, and for general laboratory instruments.†

Two major challenges in developing detection systems for clinical and environmental testing are overcoming interference from complex sample matrices and achieving the sensitivities required for detection or diagnosis. Samples from serum, saliva, and stool, as well as environmental samples such as lake or ocean water and soil contain many substances that can impede the binding of analyte to the SPR surface, or bind nonspecifically the SPR surface, interfering with the specific detection signal.

Several approaches have been used to enhance the sensitivity and reduce the background for analyte detection in complex solutions. They include selective enrichment of microorganisms prior to detection using polymerase chain reaction (PCR)-based protocols (reviewed by Benoit and Donahue in 2003);‡ immunomagnetic separation and concentration of analyte,§ and use of tangential filtration membrane barriers to exclude interfering molecules larger than the analyte of interest from the sample stream.¶ Each of these techniques has drawbacks such as the time required for selective enrichment, acidic elution steps (prior to detection) required for immunomagnetic separation, and the inability to remove interfering molecules from large analytes of interest using filtration protocols.

Because SPR-based assays rely on a change in refractive index near the sensor surface in response to a binding event, one means of enhancing SPR signals is to introduce secondary amplifying antibodies following the initial binding of target analyte to the SPR surface. Earlier, we described the utility of the secondary antibody verification/amplification method with SPR biosensors.¶ The use of an antibody specific for a different target epitope for the secondary amplification also provides verification of the analyte detection. The use of two complimentary antibodies for the detection protocol is similar to the enzyme-linked immunosorbant

assay (ELISA) method used in many current antibody-based detection systems.6

Dense particles linked to secondary antibodies have also been used to amplify the detection signal. Both colloidal gold nanoparticles7,8 and magnetic nanoparticles9 have been shown to increase the SPR signal when added as amplifiers. Colloidal magnetic particles have desirable properties for SPR detection because they can function as both a concentration/purification agent as well as an amplifier for detection. This accomplishes the two goals of increasing the sensitivity of the SPR assay (concentration and amplification by colloidal beads) as well as reducing the background interference (purification) simultaneously.

Staphylococcal enterotoxin B (SEB) (molecular weight 28.4 kDa) is one of several toxins produced by the bacterium Staphylococcus aureus and is a common cause of food poisoning outbreaks. Bacterial toxins such as SEB, which have a resistance to heat and enzymatic digestion, can cause intestinal illness in the absence of their bacterial progenitor.10 SEB is also considered a risk for use in bioterrorism due to its heat stability and high toxicity when aerosolized and inhaled.11 A sensitive, rapid assay for analyzing SEB in complex matrices would be useful.

We describe here assay protocols for immunomagnetic separation and concentration of SEB from analyte-spiked samples (buffer, stool, and serum) using small (50 nm) paramagnetic nanoparticles conjugated to monoclonal anti-SEB antibodies (immunomagnetic bead, IMBs). The nanoparticles also significantly amplify the SPR detection signal. Detection of picogram levels of SEB was achieved via this purification/concentration/amplification procedure.

The system used a semiautomated sample handling system, in which a single sample solution is delivered via syringe through an injection port into a 2 mL sample loop (Figure 1A). The sample then flows through a sample temperature conditioning channel, then sequentially over the surfaces of the eight three-channel sensor chips. The sample flow is followed by an automated wash step to return the sensor flow stream back to starting buffer. Data from each sensor channel is reported as a refractive index value once per second. The resolution of the instrument is approximately 1 \times 10^{-6} refractive index units (RIU). The numerical output from the system is a 10^6 multiple of the true value for ease of display and analysis, e.g., a 25 RIU value reported is actually 25 \times 10^{-6} RIU. The sensor housing has an adjustable temperature controller that maintains the temperature to within ±0.01 °C and was set to 25 °C for the experiments reported here. This

**EXPERIMENTAL SECTION**

**SPR Sensor System.** The portable 24-channel SPR biosensor system used for the experiments described here was constructed in our laboratory and was designed around the integrated three-channel Spreeta chips developed by Texas Instruments.12 The Spreeta SPR chips contain a light source and array detector integrated with the gold sensor surface (described in Naimushin and co-workers and Stevens et al.).12–14 The current system (Figure 1A) is an eight-chip, 24-channel system with an integrated digital signal processor (DSP). Readout and control functions are controlled with a touch-screen interface (Figure 1B). The system uses a semiautomated sample handling system, in which a single sample solution is delivered via syringe through an injection port into a 2 mL sample loop (Figure 1A). The sample then flows through a sample temperature conditioning channel, then sequentially over the surfaces of the eight three-channel sensor chips. The sample flow is followed by an automated wash step to return the sensor flow stream back to starting buffer. Data from each sensor channel is reported as a refractive index value once per second. The resolution of the instrument is approximately 1 \times 10^{-6} refractive index units (RIU). The numerical output from the system is a 10^6 multiple of the true value for ease of display and analysis, e.g., a 25 RIU value reported is actually 25 \times 10^{-6} RIU. The sensor housing has an adjustable temperature controller that maintains the temperature to within ±0.01 °C and was set to 25 °C for the experiments reported here.

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portable SPR system is housed in a 10 in. x 10 in. x 5 in. Pelican case and weighs less than 6 lb. The system operates from either line voltage (115 V) or a 12 V source such as a portable rechargeable battery.

The Spreeta sensor chip uses an 830 nm light-emitting diode (LED) light source, providing a probing distance of approximately 400 nm into the sample medium. Thus, a 50 nm particle coupled to antibody (15 nm, 150 kDa) and SEB antigen (3 nm, 28.4 kDa) is well within the range of detection by the Spreeta SPR sensor chip.

Sensor Surface Preparation. The Spreeta sensors were cleaned by bathing the gold sensor surfaces with a 10% nitric acid solution to remove organic materials. The surface was then rinsed with ddH2O (deionized, distilled H2O) and gently wiped with microscope lens paper wetted with 70% ethanol. Antibodies were attached to the cleaned gold sensor surface by direct physisorption. Twenty microliters of concentrated antibodies were attached to the cleaned gold sensor surface by direct physisorption. Twenty microliters of concentrated antibodies in Dulbecco’s phosphate-buffered saline (PBS) (Gibco no. 14200) was pipetted directly onto the horizontally oriented gold surface and allowed to incubate at room temperature for 2 h. The gold surface was then rinsed with PBS to remove unbound antibodies and inserted into the flow cell housing. All antibodies used were supplied by the Department of Defense Critical Reagents Program (CRP), (Frederick, MD). For the experiments described here, a single three-channel sensor was coated with polyclonal rabbit anti-SEB, while a second three-channel sensor chip was coated with polyclonal rabbit anti-Bacillus anthracis and served as a reference sensor. The remaining six three-channel sensors were not used, except when additional sensors were added for testing the regeneration of the sensor surface (Figure 7).

Preparation of Antibody-Coupled Nanoparticles (IMBs). Murine monoclonal anti-SEB antibodies were provided by the CRP. Biotinylation of the antibodies was carried out using EZ-Link sulfo-NHS–biotin (Pierce no. 21338, Rockford, IL) following the standard protocols provided by the manufacturer for biotiny-
Sample Materials. The stool sample used for this study was obtained from the University of Washington Medical Center (UWMC), under a University of Washington IRB approved project for assay development. The sample obtained was mostly liquid, and stool solids were allowed to settle overnight at 4°C before the sample was divided into 2 mL aliquots. This sample was considered a “blank” sample (containing no SEB) but was not analyzed for SEB content prior to the experiments described here. The unspiked control experiments in buffer and stool presented below indicate that little or no SEB was present in the stool sample prior to being spiked with purified SEB. Fetal bovine serum (FBS) was obtained from Gibco (no. 0437-028, Invitrogen Corporation, Carlsbad, CA). SEB (Sigma no. S-4881) was diluted into 2 mL of stool, serum, or buffer, and then added to an 18 mL nanobead mixture (2 mL of nanobeads in 16 mL of PBS-T) for a final volume of 20 mL.

Sample Processing. The control and SEB-spiked and solutions were incubated with IMBs at room temperature for 1 h. The sample processing steps are shown in Figure 3A. The control and SEB-spiked solutions were incubated with IMBs at room temperature for 1 h. The 20 mL samples were processed as shown in Figure 3A, with four wash steps of 250 µL each. The bead/analyte complex was eluted with 2 mL of PBS-T buffer for analysis by SPR.

Sensor System Operating Procedure. For each experiment, a 2 mL sample was injected into the holding loop and flowed through the system at 70 µL/min for 14 min; thus, a total of 1 mL of sample was introduced to the sensor before the flush step removed any remaining sample from the injection coil. The flow was then returned to the PBS-T starting buffer, which was used as the running buffer for all experiments. For the regeneration step, low-pH buffer [100 mM glycine (Fisher Scientific no. BP381-1), adjusted to pH 2.2 with 1 M HCl] was injected and flowed through the system for 10 min at a flow rate of 70 µL/min.
RESULTS AND DISCUSSION

Nanobeads Amplify the SPR Signal. Stepwise reagent addition demonstrated the level of amplification provided by the nanobeads. Figure 2A shows the sequential three-step protocol for detection of SEB and amplification of the detection signal: (1) the direct detection of free SEB in buffer solution, (2) amplification with 50 µg/mL biotinylated anti-SEB antibodies, and (3) a secondary amplification with streptavidin-nanobeads (200 µL stock µMACS beads in 2 mL total buffer) that bound to the captured biotinylated anti-SEB antibodies. Figure 2B shows a schematic of the three steps and the relative size of each particle. This multistep introduction of amplifiers demonstrated the dramatic amplification that the colloidal IMBs provide to the SPR sensing protocol.

Detection of SEB in Buffer and Stool Using the IMB Separation/Concentration Protocol. In subsequent experiments, monoclonal anti-SEB antibody coated nanobeads were mixed with solutions containing the target antigen (SEB) prior to processing and injection into the SPR system. Figure 3A illustrates the procedure used to concentrate and purify the samples. In Figure 3B a cartoon diagram of the interactions are shown. In addition to amplifying the signal, the magnetic properties of the beads were used to separate the analyte from the sample matrix (analyte affinity purification) and concentrate the sample prior to introduction to the sensor system.

Due to their small size, simply holding a magnet up to the solution is not adequate for immobilizing the nanoparticles. Magnetic nanoparticles require a column of larger packed magnetic beads (illustrated in steps 2 and 3 of Figure 3A) that, when placed in a magnetic field, capture the smaller nanoparticles. Removal of the magnet from the column allowed the beads to be eluted into a smaller volume of buffer prior to injection into the sensor system.

Figure 4 shows the results of three separate SEB detection experiments, two of which use the bead processing steps described in Figure 3. The largest signal (663 RIU, 58.2 RIU/min for the initial 5 min) was from a 1 ng/mL sample of SEB in buffer, processed according to the procedure in Figure 3. The intermediate signal (365 RIU, 30.6 RIU/min for the initial 5 min) was from a 1 ng/mL sample of SEB in stool processed according to the procedure in Figure 3, and the lowest signal (7 RIU, 0.5 RIU/min for the initial 5 min) was from a direct detection (no IMB processing or amplification) of a sample of 10 ng/mL SEB in buffer that was run for comparison. Unpurified stool and serum samples were not run through the instrument.

Figure 4 demonstrates the dramatic increase in sensitivity provided by the IMB protocol versus the detection of SEB alone. The 7 RIU signal from 10 ng/mL SEB without IMB processing (compressed in this figure) was significantly above background and was easily detected using the current system (with a background of approximately ±1 RIU). The 1 ng/mL samples processed with the IMB procedure provided a much larger signal upon binding of the analyte/bead complex. The concentration/amplification protocol rapidly purified analyte free from complex matrices and amplified the detection signal compared to the signal from the direct detection assay.

Detection Limits in Buffer and Stool Samples Using the IMB Protocol. Figure 5 illustrates the analysis of both buffer and stool samples containing four different concentrations of SEB (100, 200, 500, and 1000 pg/mL) using the procedure outlined in

![Figure 6](image-url) Detection of 1 ng/mL SEB in FBS by the procedure outlined in Figure 3A. A 2 mL sample of FBS was spiked with SEB and diluted to 20 mL with PBS-T buffer, with a final concentration of 1 ng/mL SEB. The control experiment was a repeat of the same procedure in FBS without SEB.

![Figure 7](image-url) (A) SPR sensorgram showing a low-pH wash to rapidly reset the sensor surface following a 1 ng/mL SEB detection in stool. (B) SEB detection in stool samples with zero (○) and one (□) regeneration of the surface demonstrating the functional regeneration of the sensor surface following SEB detection from the complex (stool) sample. The data for “zero” regeneration are the same as the stool detection data shown in Figure 5 and are shown for comparative purposes. Results for one regeneration at 100 (33 ± 7.4 RIU), 500 (156 ± 7.1 RIU), and 1000 (269 ± 19.8 RIU) pg/mL are shown. The control value (unspiked sample) was 15.5 RIU and was subtracted from the data shown.
Figure 3. Each data point represents the average of three separate sensor channels within the same experiment, with standard deviation error bars displayed. Verification of analyte identity was achieved simultaneously, since antibodies to one analyte epitope were used on the nanobead (murine monoclonal anti-SEB) and antibodies to multiple analyte epitopes were immobilized on the sensor surface (rabbit polyclonal anti-SEB). Reference channels derivatized with rabbit polyclonal anti-B. anthracis antibody were run simultaneously for each experiment and showed negligible nonspecific binding. Control experiments (samples run with no SEB present) were also performed, and this value was subtracted from the SEB-spiked result. The control value for unspiked buffer was 70 RIU, and the unspiked stool sample control was 17 RIU. The results shown had the reference channel and control values subtracted. Detection in stool involved a very complex matrix and did result in some loss of sensitivity. A comparison of direct detection of SEB in stool to SEB in buffer was not possible due to the potential for fouling of the fluidics system by particulates in the stool sample.

**SEB Detection in Fetal Bovine Serum.** Detection of SEB with the IMB procedure in a second complex matrix (FBS) was also demonstrated. A single concentration of SEB (1 ng/mL) diluted 1:10 in FBS solution was processed as shown in Figure 3. The results are shown in Figure 6. The lack of pre-existing anti-SEB antibodies in FBS eliminated potential complications from pre-existing anti-SEB antibodies often seen in adult sera. Sensitivity of detection of SEB in serum resulted in a signal that was comparable to that achieved in buffer.

**Regeneration of Sensor Surfaces.** Removal of bound SEB–nanoparticle complexes from the sensor surface was achieved with a low-pH buffer (100 mM glycine, pH 2.2) wash. Figure 7A shows the regeneration of the sensor following detection of a sample of 1 ng/mL SEB in stool. Figure 7B shows the detection of SEB at different concentrations with a low-pH regeneration wash step between each detection event. Regeneration appears to lower the signal slightly, but the signal remains robust for the concentrations tested. This figure demonstrates the potential for multiple uses of the same sensor surface for repeated assays, which can greatly reduce the cost and effort of repeated assays.

**CONCLUSIONS**

Toxins such as SEB are ideal candidates for detection by SPR because competing techniques such as PCR and microbial enrichment techniques require the presence of the toxin-producing organism for detection. This technique for detecting SEB with both increased sensitivity and lower background will enhance efforts aimed at improving food safety and rapid diagnosis of intoxication, as well as detecting the presence of SEB in the environment in the event of a deliberate release.

The assay sensitivities reported in this study should be adequate for detection of toxin in stool samples of affected individuals. Detection of staphylococcal enterotoxins in patient stool samples with a commercial ELISA with a reported sensitivity of 1 ng/mL (TECRA no. BP-211, Frenchs Forest, Australia, developed for use with food samples) and reverse passive latex agglutination (RPLA), (Oxoid no. TD0940, Cambridge, U.K., a kit for the detection of staphylococcal toxic shock syndrome toxin in culture filtrates) with a sensitivity of 2 ng/mL have been reported. These samples were correlated with isolates of methicillin-resistant *S. aureus* (MRSA) producing the specific enterotoxin detected by the ELISA or RPLA.

This IMB–SPR system should be adaptable for dramatically decreasing the time and expense of diagnosing the presence of many other enteric pathogens and/or their related toxins, for example, invasive fungal infections such as *Aspergillus* and *Cryptococcus* that are difficult to detect with current culture-based assays, and for detection of microbial-produced toxic proteins such as *Clostridium difficile* toxins (a common cause of nosocomial diarrhea with an increasing incidence of infection in hospitals worldwide). In the in the event of a Shiga toxin outbreak (the toxin produced by enterohemorrhagic *Escherichia coli*), rapid, sensitive detection of the toxin in stool is important for directing treatment because the bacterial progenitor requires significant time for isolation/demonstration. Environmental analyses, such as detection of domoic acid in clams or seawater, also require low-level detection as would detection of toxic proteins in the environment due to deliberate releases.

**Advantages of Nanoscale Paramagnetic Particles.** Larger (micrometer-sized) particles can also amplify the SPR signal; however, these particles rapidly sediment and are not easily mixed or directed to the sensor surface. Thus, analyte must either be eluted from the beads prior to analysis with the SPR biosensor, or complex magnetic systems must be used to direct the bead–analyte complex to the SPR surface (magnetic transport, reviewed by Gijs in 2004). Colloidal paramagnetic particles disperse evenly into solution with minimal mixing, do not settle out of solution like larger micrometer-sized paramagnetic particles, and diffuse rapidly to the sensor surface. These features of paramagnetic nanoparticles are important for automated applications and in-line processing.

The two-antibody assay described here is ideal for analytes with multiple unique epitopes, such as the protein detection described here. To use this procedure with small molecules the procedure must be modified. A small molecule without multiple determinants requires a competition assay (such as that described in Stevens et al.), which could be modified for use with nanobeads. This would add sensitivity and dynamic range to the assay and the ability to concentrate large volumes. Larger analytes (such as microorganisms and viruses) could also benefit from nanobead purification, increasing the sensitivity of detection.

The SPR platform used for the experiments reported here provides several important features, including portability, flexibility to analyze a broad range of analytes, high-throughput capability, ease of use, high sensitivity, and rapid confirmation of positive results that make it attractive for use in a broad range of applications in the medical laboratory and beyond. The portability of the SPR system, together with its multianalyte capabilities, also make it an ideal system for rapidly identifying biological agents that may be intentionally released and for systems to monitor and detect environmental toxins. We have previously reported adaptations of this system for detection of small molecules such as...
domoic acid in clam extracts\textsuperscript{14} and cortisol in saliva via a flow filtration system.\textsuperscript{4}

A significant advantage of the IMB protocol is the ease of concentrating analytes prior to introduction to SPR analysis. The experiments described here made use of a 10-fold concentration of SEB from stool and serum samples; however, much greater concentration ratios should be achievable for more dilute samples. With an added prefiltering step, 100- or even 1000-fold concentration should be achievable for dilute samples such as fresh or seawater.

The nanobead processing procedure is also amenable for automation. Control of the magnet and flow through the column can be integrated into the existing SPR fluidics system. The automated protocols can also include regeneration of the sensor surface and recycling of the assay materials, dramatically reducing the per-assay cost.

Future studies should include a more detailed examination of the reproducibility of the assay, including studies with spiked stool samples from multiple patients to determine if a generic concentration curve for stool samples can be generated, and studies from individuals with diarrhea from which enterotoxin-producing \textit{Staphylococcus} have been isolated. Although chip-to-chip variation was not examined systematically in the experiments described here, evidence that the variability is low is provided by data such as the regeneration graph in Figure 7B, where different sensors were used for two separate assays at each concentration of analyte.

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