Review

Small Molecule Immunosensing Using Surface Plasmon Resonance

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Abstract: Surface plasmon resonance (SPR) biosensors utilize refractive index changes to sensitively detect mass changes at noble metal sensor surface interfaces. As such, they have been extensively applied to immunoassays of large molecules, where their high mass and use of sandwich immunoassay formats can result in excellent sensitivity. Small molecule immunoassaying using SPR is more challenging. It requires antibodies or high-mass or noble metal labels to provide the required signal for ultrasensitive assays. Also, it can suffer from steric hindrance between the small antigen and large antibodies. However, new studies are increasingly meeting these and other challenges to offer highly sensitive small molecule immunoassay technologies through careful consideration of sensor interface design and signal enhancement. This review examines the application of SPR transduction technologies to small molecule immunoassays directed to different classes of small molecule antigens, including the steroid hormones, toxins, drugs and explosives residues. Also considered are the matrix effects resulting from measurement in chemically complex samples, the construction of stable sensor surfaces and the development of multiplexed assays capable of detecting several compounds at once. Assay design approaches are discussed and related to the sensitivities obtained.

Keywords: immunosensor; surface plasmon resonance (SPR); small molecule; steroid; toxin; conjugation
1. Introduction

Surface plasmon resonance (SPR) is an opto-electronic phenomenon that occurs when a photon of light is incident upon a noble metal surface such as gold or silver [1]. When the wavelength of the photon equals the resonance wavelength of the metal, then the photon couples with the surface and induces the electrons in the metal surface to move as a single electrical entity called a plasmon. This oscillation of electrons sets up an electromagnetic field that exponentially decays out from the metal surface, with significant electrical field strength typically occurring within 300 nm of the surface. When molecules with sufficient mass bind to the surface within the range of the electric field, they perturb the plasmon and change the resonance wavelength. When dealing with a fixed planar surface, this is seen as a shift in the resonance angle of the incoming photons. These shifts essentially reflect minute refractive index changes on the surface and so can be used to very sensitively detect the binding of mass to the surface, typically down to a change of less than $1 \times 10^{-6}$ refractive index units (RIU) for more sophisticated instruments (Figure 1). Refractive index is a ratio that changes from 1.0003 in air to 1.33 in water. Resonance units (RU) are often used to quantify refractive index changes in SPR biosensors, with 1 RU = $1 \times 10^{-6}$ RIU, and so are used as units describing SPR signal strength.

Figure 1. A schematic of the conventional Kretschmann optical configuration for SPR biosensing and the associated angle shift and sensorgram plot of resonance signal change with time [2]. Reprinted by permission from Macmillan Publishers Ltd: Nat. Rev. Drug Discov. 2002, 1, 515-528 copyright 2002; http://www.nature.com/nrd/index.html.

Immunoassays involve the binding of an antibody to its target antigen, and so the antibody, being a high mass (approximately 150 kDa) protein, can act as the signal generator in SPR transduction. Immunoassays therefore naturally lend themselves to SPR biosensing, and this is particularly the case
for large molecule antigens (Mr > 2 kDa) which can be assayed in sandwich immunoassay formats. A sandwich immunoassay format would involve immobilizing antibodies onto the sensor surface, utilizing chemistries that include functionalized self-assembled monolayers (SAMs) [3], polymer coatings [4] and proteins such as protein A that orient antibodies on sensor surfaces [5]. The high mass antigen can then bind the immobilized antibody directly, thus generating biosensor signal and this signal can be further enhanced by binding a secondary antibody that recognizes another epitope on the antigen (Figure 2A). The design of large molecule immunoassays using SPR is now a very mature field of research.

Small molecule antigens, however, pose challenges not encountered with large molecules. The foremost of these is that the antigen itself can not generate very much SPR signal, given its small mass. There are some reports of kinetics studies and assays using direct detection of small molecule targets but these generally suffer from low signal and poor sensitivity (Figure 2D). To obtain optimal assay sensitivity the antigen is therefore either labeled with a high mass label and used in competition with un-labeled sample antigen for binding to the surface in a competitive immunoassay, or the small molecule antigen is conjugated to the sensor surface and primary antibody is mixed with sample containing free antigen and the mixture is passed over the sensor surface (Figures 2B and C). In this case the mass is provided by the primary antibody and signal can be further enhanced by use of secondary antibodies either with or without conjugation to gold nanoparticles [4] (Figure 2B). In either case, the sensor signal is inversely proportional to the concentration of the antigen free in solution. Gold nanoparticles can provide signal enhancement both through their high mass and through cooperative plasmon enhancement by coupling between the localized plasmon field of the nanoparticle and the surface plasmon field of the gold sensor surface [6]. Small molecule assay formats require very careful design of the surface chemistry and the labeling employed so as to ensure optimal sensitivity.

Another critical concern with small molecule immunoassay using SPR is the potential for steric hindrance of the binding between antigen and antibody when there is either a large label proximal to the antigen or where the antigen is bound to the sensor surface. Careful use of appropriate linker chemistry can help mitigate these constraints and improve binding signal and sensitivity. The stability of the immunobiosensor surface to the harsh solutions used to regenerate and re-use the surfaces ready for another assay must also be considered, particularly for high-throughput applications. The use of high quality antibodies with high affinity towards the target compound are also important to achieving high SPR sensor signal and low limits of detection (LOD). If the antibody binds the analyte strongly then smaller concentrations of the analyte may be needed to inhibit antibody binding to the sensor surface, thus reducing LOD. Typically, antibodies with affinity constants of 1–10 × 10^9 L/mol are desirable. Another major consideration is the resistance of the surface to biofouling caused by non-specific binding from high mass contaminants in complex real-world samples. These can both disrupt antigen/antibody binding and deposit high mass on the surface, potentially distorting immunoassay signals.

This review will examine the general principles and parameters of small molecule immunoassays and how others have designed their immunosensor systems using SPR transduction. Specific classes of target small molecule antigens of particular interest are then investigated, namely steroids, toxins and food components, drugs and explosive residues, before some general comments are made on possible
future directions in small molecule SPR immnosensor research. A summary of some SPR immunoassays of small molecules is given in Table 1.

**Figure 2.** A schematic of some typical SPR immnosensor formats. **A.** Sandwich immunoassay for large molecules. **B.** Protein conjugate immobilized indirect inhibition immunoassay (can also link via self-assembled monolayers (SAMs) or carboxymethyl dextran polymers) with optional secondary antibody-gold nanoparticle labeling in a second step. **C.** Protein-labeled inhibition immunoassay. **D.** Direct small molecule immunoassay.

**2. General Principles and Immunosensor Assay Formats**

When developing small molecule immunobiosensors for use with SPR transduction, one must first consider the structure and assembly of the sensing surface. The small molecule antigen can be immobilized to the gold surface via the commonly used carboxymethyl dextran polymer layers. These layers are functionalized with carboxylic acid groups that allow covalent attachment of antigen or antigen derivatives containing an amino group through the formation of an amide bond. Steroid hormones have been conjugated using this technique in a convenient *in-situ* flow-through immobilization technique within the biosensor by attaching a linker at a point on the antigen distant from existing functional groups [4]. The resulting functionalized surfaces can withstand up to more than 1,100 binding and regeneration cycles [4]. They use hydrophilic oligoethylene glycol (OEG) linkers which can project the antigen into the aqueous fluid stream and allow for optimal antibody binding [4,7]. Binding is also optimized through careful selection of the position of conjugation on the steroid so that linkers are attached distant from existing functional groups [7].

Another key consideration is the method of signal generation. With antigen-immobilized formats, the primary antibody can provide the signal but studies using steroid antigens have also employed secondary antibodies to further enhance the signal strength with enhancements of 6–8 times the primary antibody binding signal [4] (Figure 3). In addition to this enhancement approach, gold nanoparticles have been used to further enhance signal by adding more bound mass and through cooperative plasmon coupling. Gold nanoparticles concentrate a high mass into a small volume. Their
surfaces allow convenient formation of coordinate bonds with thiol functional groups and so they can be easily conjugated to biomolecules as signal enhancement labels. The diameters of the nanoparticles can also be easily adjusted between 1–100 nm as required. The noble metal nanoparticles also undergo SPR themselves and so when they approach a gold SPR sensing surface their plasmon modes can couple with those of the surface and produce a large shift in refractive index and thus an enhancement of SPR signal much larger than that expected on the basis of high mass labeling alone [6]. This labeling approach has been widely applied to SPR sandwich immunoassays of large molecules but has only recently been extended to small molecules. When 25 nm gold nanoparticles conjugated to secondary antibodies were employed, enhancements of 13-fold were achieved for immunosensing of progesterone with a LOD of 8.6 pg/mL [4]. Protocols for the enhancement of small molecule SPR immunosensing have now been standardized and these approaches can provide a generic platform for small molecule immunosensing combining linker chemistry with nanoparticle enhancement [8]. The range of small molecule analytes to which gold nanoparticle enhancement has been applied is now growing and includes the steroid hormone metabolite estriol-16-glucuronide [9,10], the antibiotic chloramphenicol [11] and the neurotoxin ochratoxin A [12]. Consideration has also been given to the effects on SPR signal enhancement of the diameter of the nanoparticles [13] and their distance from the sensing surface [14].

**Figure 3.** A sensorgram of the primary monoclonal antibody (mAb) binding response to a progesterone-immobilized SPR sensor surface and enhancement of binding signal with secondary antibody followed by regeneration. Reprinted from reference [4], with permission from Elsevier. Copyright 2005.

Similar enhancement techniques can also be employed with other types of surfaces such as thiol SAMs terminating in carboxylic acid groups [3] (Figure 4). This approach has been used in combination with protein conjugates of the antigen to construct sensor surfaces for detection of progesterone [3], chloramphenicol [11] and ochratoxin [12]. In these cases gold nanoparticles of 10 nm, 40 nm and 40 nm, respectively, were employed. Whilst SAMs allow the binding events to take place closer to the gold surface where the field strength is higher, they are typically less stable under regeneration than dextran surfaces [3].
Figure 4. Standard curve plots for progesterone using a mixed SAM layer with protein-conjugate immobilization for SPR immunosensing using mAb only (●), secondary antibody enhancement (○) and 10 nm immunogold enhancement (▼). Reprinted from reference [3], with permission from Elsevier. Copyright 2007.

An alternative format for analyzing binding interactions in small molecule SPR immunosensing is to immobilize antibody and examine binding of small molecule protein conjugates. This approach has been used for progesterone when examining the effects of intermediate linker length on antibody binding strength, where a linker of 18-atoms was found to give high binding signal [15].

Most SPR immunoassays for small molecules are conducted in a microfluidic environment. Mathematical models of the effect of various system parameters such as flow rate, antibody concentration, and density of binding sites on the surface have been developed to help optimize sensitivity [16]. Such modeling approaches have been applied to concentration gradient immunoassay where space and time-dependent binding is analyzed in a two-dimensional imaging SPR configuration with introduction of sample and antibody through ports in a T-shaped sensor [17]. This approach involves parallel streams of fluid, one containing analyte and the other antibody and analyzes the steady state gradient set up by interdiffusion between the fluid streams through binding to the immobilized antigen phenytoin [18].

Multiplexing assays in array formats is of increasing interest with small molecules, given the success of protein arrays [19]. SPR imaging is a format that utilizes the localized plasmon fields of specially patterned metal surfaces to detect SPR shifts spatially in two-dimensions. It is widely used in the study of protein binding interactions [20]. A photo-crosslinked small molecule platform has been successfully applied to detection of receptor binding to estrogens using SPR imaging [21]. Screening protein-protein interactions in a microarray format using scattered light under SPR conditions is emerging as an imaging SPR format [22] and may be adapted in future to small molecule immunoassays.

Another key consideration is the continued functioning of the sensor in the presence of complex sample matrices such as serum [23] and human saliva. There is also on-going interest in making SPR small molecule sensor systems portable, and this is particularly useful in environmental measurements requiring on-site, real-time analysis. Benzo[a]pyrene and 2-hydroxybiphenyl have been detected using
an indirect competitive immunoassay where the SPR sensor system is confined to dimensions of 16 cm × 9 cm × 6 cm in a four flow-channel system [24].

Recently, interest has been building in alternatives to antibodies as binding agents in small molecule SPR formats. Aptamers (protein or oligonucleic acids that recognize specific target molecules) are being considered whereby binding between the aptamer and an immobilized partial complementary ss-DNA strand is disrupted by the presence of the small molecule target. This has been applied with gold nanoparticle labeling for detection of adenosine [25]. Such formats have shown detection performance down to 1 nM when used as an SPR biosensor for adenosine [26].

3. Steroids

Analysis of steroid hormones is of interest to scientists investigating human physiology, optimization of athletic training regimes, monitoring of reproductive cycles in humans and animals for artificial insemination and pregnancy planning, and in the diagnosis and treatment of hormonal disorders. SPR has been applied to the measurement of progesterone in milk for potential use in monitoring estrous cycles in cows [27]. This system immobilizes progesterone ex-situ by removing the carboxymethyl dextran coated gold sensing surface from the chip cover and dispensing onto it a solution of 3-carboxymethyloxime derivative outside the SPR instrument. It demonstrated a LOD of 3 ng/mL which was later improved to 0.4 ng/mL in cow’s milk [27,28]. This system uses significant dilution to overcome matrix effects [27,28], which can compromise the overall assay sensitivity. Estradiol, in both serum and seawater has been sensed using a combination of SPR and a coupled online in-tube solid-phase microextraction (SPME) system. This inhibition immunoassay gave an LOD of 170 pg/mL [29]. Estradiol has been measured in buffer-based assays using OEG covalent linker conjugation and LOD of 25 pg/mL were obtained when conjugation position is carefully considered [7]. Another report utilizes an estradiol-protein conjugate for immobilization and achieves inhibition at concentrations of about 0.6–30 ng/mL [30]. In serum, SPR has shown that sex hormone binding globulin can form aggregates that may impact on protein immunoassay performance [31]. Testosterone and estradiol have been derivatized with biotin and pre-incubated with streptavidin before immobilization on the sensor surface. Subsequently, antibody binding kinetics were examined and showed good correlation with quartz crystal microbalance (QCM) results [32].

The highly sensitive detection of the steroid hormones cortisol and testosterone in human saliva is of great interest to physiologists studying the effects of exercise on athletic performance. It can potentially provide a non-invasive means of monitoring hormonal fluctuations with time and could also be used to help diagnosis of hormonal disorders such as Addison’s disease and Cushing’s syndrome. Utilizing the covalent immobilization and OEG linker technology developed for progesterone [4], it has been possible to construct highly sensitive SPR immunoassays for both cortisol [33] and testosterone [34] (Figure 5). LOD of 49 pg/mL for cortisol and 15.4 pg/mL for testosterone have been reported in stripped human saliva matrix [33,34]. The typical physiological range is 0.1–10 ng/mL for salivary cortisol and 29–290 pg/mL for salivary testosterone [35]. The cortisol immnosensor uses secondary antibody enhancement of SPR signal [33], whilst the testosterone assay uses secondary antibody-nanogold enhancement to achieve greater sensitivity [34]. Non-specific binding has been reduced in the case of cortisol by the use of a surfactant in the antibody
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diluents, which removes the need for chemical extraction or extensive pre-treatment of saliva samples [33]. Other groups have proposed the use of a combination of membrane filtration and an H-filter [36] to remove mucins and other high mass components in saliva or using a flow-filter arrangement [37] but this can add time, expense and complexity to the biosensor system [36] and risk repeated fouling of membranes. A cortisol-protein conjugate immobilized format has demonstrated an LOD of 1 ng/mL, insufficient to cover the full physiological range for salivary cortisol [37]. Subsequent work attempting direct immunoassay has demonstrated poor sensitivity, with reported linear detection regions ≥13 ng/mL for saliva and ≥9 ng/mL in urine [38]. Cortisol has also been used as a target antigen in the development of SPR immunosensors utilizing nanohole array substrates [39]. Such a set-up has the advantage of simple optics (they only need a simple UV-visible spectrophotometer) giving the potential for portable units to be constructed. Detection of small molecules such as the steroids in environmental samples is of increasing importance [40] as understanding is gained of their effects on freshwater and marine ecosystems, and so SPR has also been applied to environmental measurement of 17β-estradiol in seawater [41].

**Figure 5.** Correlation plot for salivary cortisol samples analyzed by an SPR biosensor and by radioimmunoassay for 40 samples, from reference [33]. Reproduced by permission of The Royal Society of Chemistry (RSC).

4. Toxins and Food Safety and Composition

One of the key areas of interest in small molecule SPR immunosensing is the detection of toxins in foods and beverages to comply with increasingly strict requirements for food safety testing [42-44]. The key requirements are highly sensitive detection in complex matrices with high throughput. Also of interest is the detection of bioactives and additives in foods [42]. Much of the work in this area is focused on veterinary drugs, shellfish toxins and antibiotic residues, with less work devoted to mycotoxins. SPR is increasingly being explored as a transduction platform for such assays [45] along with other biosensor techniques, such as quartz crystal microbalance technology and electrochemical transduction [46]. Detection of toxins on-site using portable SPR is also beginning to be
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investigated [47]. The antibiotics neomycin and gentamycin have been measured in a parallel format using an imaging SPR system from IBIS Technologies, in a competitive assay with antibiotic immobilized through its amino groups to a carboxymethylated dextran. This resulted in detection ranges in the 1–50 ng/mL region [48]. The poor sensitivity is likely due to the absence of a linker between the antigens and the surface, or conjugation through an existing functional group.

Mycotoxin immunoassay using biosensors has attracted attention as a new technology for food safety screening [49]. Combined detection of T-2 and HT-2 toxins is possible using SPR by organic extraction from baby food, breakfast cereal and wheat. LOD of 25–26 µg/kg were obtained with CVs of 1.8–6.3% [45]. Fungal metabolites are often toxic, and an example is deoxynivalenol which can be conjugated to casein to coat an SPR biosensor surface for a competitive immunoassay [50]. These surfaces are reported to be stable for about 500 assay cycles but function in a high concentration range of 2.5–30 ng/mL [50]. Ochratoxin has been determined in cereals and wine using ochratoxin-OEG-ovalbumin protein conjugates on a SAM layer with 40 nm gold nanoparticle signal enhancement [12] (Figure 6). A radically different format for ochratoxin detection is based on hollow gold nano-sized balls coated in a dendritic thionine thin film for immobilization and sensing [51]. The technique was also tried in milk [51]. Staphylococcal enterotoxin can be detected in raw eggs using an antigen immobilized SPR competitive immunoassay which allows detection at 1–40 ng/mL from supernatant [52]. Aflatoxin B1 has been the subject of a protein conjugate-immobilized sensor format, with a polyclonal antibody and special regeneration conditions to overcome strong antibody binding [53]. In comparison, another study has favored using immobilized enzyme in an SPR format to detect aflatoxin [54].

Shellfish toxin detection is another major application that has been explored extensively using SPR immunosensing. Okadaic acid has been detected by organic extraction from mussels and scallops and demonstrated minimal matrix effects in a direct competitive assay format and with an LOD of 0.24 ng/mL in mussel extract, allowing detection below European regulatory limits of 160 µg/kg [55]. It has also been detected using an amine coupling method which has shown stability over more than 50 assay cycles, though the immobilization method used may suffer from steric hindrance of antibody binding [56]. Antibodies have been raised to domoic acid and immunoassays established in both competitive and displacement assay formats, with the competitive format giving an LOD of 3 ng/mL and correlating well to HPLC when applied to measurement in clams [57]. Another study has shown that domoic acid-immobilized surfaces can be stable for 800 assay cycles, with measurement complete within 10 min and with reported LOD in the ng/g region, below the EU action limit of 20 µg/g [58]. The binding interactions between domoic acid and polyclonal, monoclonal and recombinant antibodies have been examined using SPR formats with immobilized domoic acid [59]. A number of paralytic shellfish toxins have also been measured using SPR by employing a new saxitoxin polyclonal antibody and compared with ELISA and mass spectrometry techniques. The biosensor had a reported best LOD of 21.6 µg/100 g of shellfish compared to a European regulatory limit of 80 µg/100 g of shellfish [60]. Proteins of potential use in assays of saxitoxin have been screened using SPR by immobilizing the toxin via amine coupling onto the sensor surface [61].
SPR has also been used to probe the binding interactions between proteins and toxins and to examine such effects as the interaction between the ABC ring structure of ciguatoxin and its specific antibody, revealing that the bulkiness and aromatic nature of the antigen was critical to specific antibody recognition [62]. Another study has immobilized ciguatoxin using a 3-butene-1,2-diol side chain attached to the A-ring and then examined antibody binding interactions [63]. Tetrodotoxin has been measured using a modified OEG SAM surface structure, which achieved minimization of non-specific binding and an LOD of 300 pg/mL [64] (Figure 7). Using this approach the ratios of the functionalized and blocking SAM thiol chains could be optimized to provide adequate spacing between immobilized antigen molecules to allow stronger antibody binding [64]. 2,4-D can be detected by using a protein conjugate [65,66] to immobilize the antigen and an indirect competitive assay format. The latest report indicates an LOD of 100 pg/mL using multiple flow channels [65]. An emerging related method is the use of total internal reflection ellipsometry, which may provide an
alternative transduction approach for sensitive detection of small molecule toxins such as T-2 mycotoxin [67].

Determination of food composition using SPR immunosensing appears to have attracted much less attention than toxins detection, but investigators have measured compounds such as vitamin B-5, which was measured in a wide range of foods including reference samples, giving a LOD of 4.4 ng/mL using buffer extraction [68].

**Figure 7.** Immobilization of tetrodotoxin onto a mixed thiol SAM on an SPR gold surface.
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### 5. Drugs

Localized surface plasmon resonance (LSPR) has been used for the detection of the drug stanozolol using immobilized gold nanoparticles [69,70]. Here stanozolol-protein conjugate is electrostatically adsorbed onto the gold, and antibody binding to the surface is detected with an LOD of 0.7 ng/mL [70]. Often, detection of drug traces on surfaces is of interest, so a SensiQ® SPR biosensor has been applied to detection of methamphetamine traces on ceramic tiles down to 25 ng/100 cm² [71]. Using LSPR, it is possible to multiplex assays in an array and this has been applied to simultaneous detection of cocaine, ecstasy, heroin and amphetamine using a combination of antibodies and with antigen-protein conjugates immobilized on the array and allowing both SPR and ellipsometric sensorgram readouts [72]. The beta-agonist clenbuterol has been detected in urine and serum and the matrix effects involved have been investigated [73]. This study found that non-specific binding from urine was greatly influenced by salt concentration and pH and that ultracentrifugation was effective in combating non-specific binding for both urine and serum [73]. Another format detects chloramphenicol using a SAM/protein conjugate surface and gold signal enhancement [11]. Fenicol antibiotic residues in shrimp have been examined using organic extraction followed by detection of four different fenicols over two flow cells using a direct amine coupling to carboxymethylated dextran without linkers [74]. Chloramphenicol can be detected in milk using a BIACore™ inhibition format involving amine coupling of the antigen to the sensor surface and using an antibody recognizing an epitope distant from the amine group and thus helping to improve antibody binding and
sensitivity [75]. An LOD of 0.1 ng/mL is reported [75]. Warfarin has been examined by SPR, with the formation of a panel of warfarin-protein conjugates which were then used to raise a selection of anti-warfarin monoclonal antibodies [76]. These antibodies were then applied in inhibition assays of warfarin to determine the un-bound fraction in blood plasma ultrafiltrate [76]. This technique was compared with HPLC and gave a detection range of 4–250 ng/mL [76]. The sensor surfaces were produced by covalent amine coupling of 4’-aminowarfarin [76].

The study of binding interactions between antibodies and drug molecules is a rapidly emerging area where SPR’s biomolecular interaction analysis capability is being utilized to understand the kinetics of drug binding interactions. This field of fundamental biomedical research is too large to cover adequately in this review, but recent examples include the covalent immobilization of the drug panitumumab to a carboxymethylated BIACore™ surface by amine coupling to assess its clinical immunogenicity [77]. In another example, single-chain fragment variable (scFv) antibody fragments are identified that can recognize a heroin metabolite without recognizing morphine. The SPR biosensor was used to determine binding affinities using a biotin-PEG linker to conjugate the metabolite to the surface [78].

In addition to the detection of drugs in humans and animals, the determination of antibiotic residues in foods, and beverages such as milk, is also of significant interest as researchers look for alternatives to centralized LC-MS technology [79,68]. Immunosensors are seen as a potential way forward to faster and more affordable analysis [80]. β-Lactams have been measured in milk using a SPR surface where the analyte is immobilized via an immunogenic interaction between H1 and its complement antibody and gives detection for penicillin G as low as 1–2 μg/kg [81]. Benzimidazole carbamate residues have been detected in milk also, this time using a liquid extraction/partition technique and immobilized antigen-protein conjugate, giving an LOD of 2.7 μg/kg [82]. Also detecting in milk, an imaging SPR system was developed to detect five different antibiotic residues in 10-fold diluted milk, with performance comparable to conventional SPR designs [83]. Taking multiplexing even further, 13 fluoroquinolone antibiotics have been detected in poultry, fish and eggs [84,85] (Figure 8) giving detection for norfloxacin of below 0.5–1.5 ng/g [84]. Fluoroquinolone SPR has been compared with LC-MS techniques for identification of contaminated samples and has performed well in this regard and demonstrated time and sensitivity advantages over more conventional microbiological techniques [86]. Flumequine, another fluoroquinolone, has been measured in the blood serum and muscle of broiler chickens using immobilized antigen and detected concentrations of 15–800 ng/mL [87]. SPR techniques can also be used in concert with more conventional LC-MS techniques to analyze fluoroquinolones. The SPR biosensor can serve to screen the antibiotics of interest and identify relevant fractions for further investigation with LC-MS [88]. In the case of penicillin G detection in milk, enzymes have been applied in place of antibodies but still using SPR transduction [89].

The antibiotic cephalixin was detected in the range of 244–3,900 ng/mL in milk using an antigen-protein conjugate immobilized regenerable surface with a polyclonal antibody [90,91]. The high detectable concentration range in the assay could be attributed, at least in part, to the lack of a spacer chain between the cephalixin and the carrier protein and conjugation through an existing functional group on the drug analyte. Nicarbazin is a coccidiostat two-drug combination used as an additive to chicken feed to prevent disease outbreaks. One of these drugs, 4,4-dinitrocarbanilide is
detected in an SPR format that immobilizes a mimic of the drug on a carboxymethyl dextran surface for detection from liver and egg samples with LOD of 17–19 ng/g [92].

**Figure 8.** Assay standard curves for norfloxacin detection in chicken meat, egg and fish.

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Some progress has been made in the development of prototype systems capable of high-throughput detection. Bile analysis for sulfamethazine and sulfadiazine has been reported, where up to eight samples can be simultaneously detected with a throughput of up to 650 samples per day with low false negative and false positive rates [93]. This prototype technology was also applied to measurement of clenbuterol and ethinylestradiol in urine and to sulfamethazine, sulfadiazine and enrofloxacin in milk [94]. An earlier report detailed a sulfamethazine immunoassay using SPR in animal urine with LOD of 5 ng/mL for use as a drug residue screening test [95]. This format interestingly uses anti-idiotype antibodies immobilized on the sensor surface and detects sulfamethazine through the inhibition of monoclonal anti-sulfamethazine binding to the surface [95]. Sulfamethazine has also been detected in milk using covalent attachment of the antigen to a carboxymethyl dextran surface and trying two different antibodies, giving a best LOD of 1.7 μg/kg [96]. An earlier study reported detection down to <1 nM in milk [97]. Use of a secondary antibody to further enhance the signal in sulfamethazine immunoassay using SPR allowed reduction of the primary antibody concentration used [97].

BIAcore™ SPR detection has been applied to ivermectin determination in bovine liver using acetonitrile extractions and solid phase extraction (SPE) clean-up with LOD of 19.1 ng/g [98]. Streptomycin residue detection has been examined in an indirect inhibition assay format using protein conjugate immobilization and in a direct assay format. Both of these approaches led to poor detection performance, with IC$_{50}$ of 10–20 ng/mL [99]. 7-Hydroxycoumarin (umbelliferone) can be detected in human serum using SPR transduction where an antigen-protein conjugate is immobilized on a BIAcore™ sensor chip measuring in the range of 0.5–80 μg/mL [100]. The buffer composition was developed to try to minimize non-specific binding [100].
6. Explosives Residues

SPR is beginning to be applied to detection of explosives residues [101], forming a growth area in SPR immunosensing. 2,4,6-Trinitrotoluene (TNT) has been detected using an indirect competitive format where a TNT analogue is immobilized to the sensor surface in the form of a 2,4,6-trinitrophenyl-keyhole limpet hemocyanin conjugate [102]. Various monoclonal and polyclonal antibodies were tested for their binding strengths, and a lowest LOD of 2 pg/mL was reported when using a polyclonal antibody raised to the same conjugate [102], showing that careful matching of the antibody and surface chemistry was central to obtaining good detection performance. TNT has also been assayed using a dendrimer-modified SPR surface [103]. In this format, a thiol SAM combined with a poly(amidoamine) (PAMAM) dendrimer provides the support structure for attachment of dinitrotoluene-keyhole limpet hemocyanin conjugate which serves as the immobilized antigen for competitive immunoassay of TNT using a monoclonal antibody and giving an LOD of 110 pg/mL on a regenerable surface [103]. Another study of TNT SPR immunosensing uses a mixed thiol SAM where some of the thiols act as blocking agents and others are functionalized for immobilization of the antigen [104]. Three TNT analogues are used for immobilization and their efficacy in the immunoassay is assessed. The surfaces are reported to exhibit low non-specific binding [104]. Irreversible gas-phase detection of TNT has also been reported using a dry surface [105]. Hexahydro-1,3,5-trinitro-1,3,5-triazine (RDX) has been detected by an interesting SPR sensor construction that involves cross-linking gold nanoparticles using electropolymerized bisaniline cross-linking in the presence of Kemp’s acid, which yields an imprinted nanocomposite that demonstrated a high binding affinity for the explosive analyte [106]. The dielectric properties of the composite changes as π-donor –acceptor complexes are formed, which alters the localized surface plasmons of the nanoparticles, giving a reported LOD of 12 fM [106]. An OEG functionalized surface has been used to detect 2,4-dinitrotoluene with SPR immunosensing, whereby the analyte is immobilized onto an OEG SAM and an indirect competitive immunoassay is set up using a specially prepared polyclonal antibody [107]. An LOD of 20 pg/mL was obtained and binding response maintained for more than 30 assay cycles [107].

Table 1. A summary of some SPR immunoassays of small molecules, giving the target analyte, the sample medium, reported limit of detection (LOD) and method of enhancement (if used). Note that researchers often use different methods for calculating LOD.

| Target         | Medium                | Reported LOD | Enhancement          | Reference |
|----------------|-----------------------|--------------|----------------------|-----------|
| 2,4,6-Trinitrotoluene | Buffer                | 2 pg/mL      | none                 | [102]     |
| 2,4,6-Trinitrotoluene | Buffer                | 110 pg/mL    | none                 | [103]     |
| 2,4,6-Trinitrotoluene | Buffer                | 1,000 pg/mL  | none                 | [104]     |
| 2,4,6-Trinitrotoluene | Vapour                | 700 pg/mL    | none                 | [105]     |
| 2,4-D            | Buffer / river water  | 100 pg/mL    | none                 | [65]      |
| 2,4-D            | Buffer / river water  | 8 pg/mL      | antibody / protein   | [65]      |
| 2,4-Dinitrotoluene | Buffer                | 20 pg/mL     | none                 | [107]     |
| 2,4-Dinitrotoluene | Buffer                | 100 pg/mL    | none                 | [24]      |
| 2-Hydroxybiphenyl | Buffer                | 500,000 pg/mL| none                 | [100]     |
Table 1. Cont.

| Substance                        | Matrix             | Concentration | Signal          |
|----------------------------------|--------------------|---------------|-----------------|
| Aflatoxin B1                      | Buffer             | 3,000 pg/mL   | none            |
| Aflatoxin B1                      | Maize              | 970 pg/g      | none            |
| Amphetamine                      | Buffer             | 5,000 pg/mL   | none            |
| Benzimidazole carbamate           | Milk extract       | 2,700 pg/g    | none            |
| Benzo[a]pyrene                    | Buffer             | 100 pg/mL     | none            |
| Cephalexin                        | Buffer             | 4,880 pg/mL   | none            |
| Cephalexin                        | Milk               | 244,000 pg/mL | none            |
| Chloramphenicol                   | Shrimp extract     | 100 pg/mL     | none            |
| Chloramphenicol                   | Milk               | 100 pg/mL     | secondary antibody |
| Chloramphenicol                   | Buffer             | 0.00074 pg/mL | 40 nm gold      |
| Cocaine                          | Buffer             | 2500 pg/mL    | none            |
| Cortisol                          | Saliva             | 49 pg/mL      | secondary antibody |
| Cortisol                          | Buffer             | 360 pg/mL     | none            |
| Cortisol                          | Saliva             | 1,000 pg/mL   | none            |
| Cortisol                          | Saliva / Urine     | 2,000 pg/mL   | none            |
| Cortisone                         | Saliva / Urine     | 9,000 pg/mL   | none            |
| Deoxynivalenol                    | Wheat extract      | 2,500 pg/ml   | none            |
| Dihydrostreptamycin               | Milk               | 20,000 pg/mL  | none            |
| Domoic acid                       | Clam extract       | 3,000 pg/mL   | none            |
| Domoic acid                       | Shellfish          | ≥ 1000 pg/g   | none            |
| Ecstasy                           | Buffer             | 5,000 pg/mL   | none            |
| Estradiol                         | Serum/Seawater     | 170 pg/mL     | none            |
| Estradiol                         | Buffer             | 25 pg/mL      | secondary antibody |
| Estradiol                         | Buffer             | 600 pg/mL     | none            |
| Estradiol                         | Seawater           | 455 pg/mL     | none            |
| Estriol-16-glucuronide            | Buffer             | 14 pg/mL      | 15 nm gold      |
| Estriol-16-glucuronide            | Urine              | 16 pg/mL      | gold            |
| Florenicol                        | Shrimp extract     | 200 pg/mL     | none            |
| Florenicol amine                  | Shrimp extract     | 250,000 pg/mL | none            |
| Flumequine                        | Chicken serum      | 15,000 pg/mL  | none            |
| Flumequine                        | Chicken muscle     | 24,000 pg/g   | none            |
| Heroin                            | Buffer             | 500 pg/mL     | none            |
| Ivermectin                        | Bovine liver       | 19,100 pg/g   | none            |
| Methamphetamine                  | Buffer             | 9,000 pg/mL   | none            |
| Nicarbazin                        | Chicken Liver      | 17,000 pg/g   | none            |
| Nicarbazin                        | Egg                | 19,000 pg/g   | none            |
| Norfloxacin                       | Poultry meat       | 500 pg/g      | none            |
| Norfloxacin                       | Egg                | 1,000 pg/g    | none            |
| Norfloxacin                       | Fish               | 1,500 pg/g    | none            |
| Norfloxacin                       | Fish / Poultry/Egg extracts | 1,000-50,000 pg/g | none |
| Ochratoxin A                      | Wine / beverages   | 58-400 pg/mL  | 40 nm gold      |
| Ochratoxin A                      | Buffer             | 10 pg/mL      | none            |
| Okadaic Acid                      | Mussel extract     | 240 pg/mL     | protein label   |
| Okadaic Acid                      | Shellfish          | 20,000 pg/g   | none            |
| Penicillin G                      | Buffer / milk      | 1,000-2,000 pg/g | none         |
### Table 1. Cont.

| Analyte                        | Source                | Concentration | Label         | Ref. |
|--------------------------------|-----------------------|---------------|---------------|------|
| Penicillin G                   | Milk                  | 2,600 pg/g    | none          | [89] |
| Phenytoin                      | Buffer                | 1,900 pg/mL   | none          | [18] |
| Progesterone                   | Buffer                | 8.6 pg/mL     | 25 nm gold    | [4]  |
| Progesterone                   | Buffer                | 4.9 pg/mL     | 10 nm gold    | [3]  |
| Progesterone                   | Milk                  | 400-600 pg/mL | none          | [27] |
| Progesterone                   | Buffer                | 35-60 pg/mL   | none          | [27] |
| Progesterone                   | Milk                  | 3,000 pg/mL   | none          | [28] |
| Progesterone                   | Buffer                | 100 pg/mL     | protein label | [15] |
| Saxitoxin                      | Shellfish             | 216,000 pg/g  | none          | [60] |
| Stanozolol                     | Buffer                | 700 pg/mL     | colloidal gold sensor surface | [70] |
| Staphylococcal enterotoxin     | Egg supernatant       | 1,000 pg/mL   | none          | [52] |
| Streptomycin                   | Milk                  | 20,000 pg/mL  | none          | [99] |
| Sulfadiazine                   | Porcine Bile          | 28,000 pg/mL  | none          | [93] |
| Sulfamethazine                 | Porcine Bile          | 15,000 pg/mL  | none          | [93] |
| Sulfamethazine                 | Animal urine          | 5,000 pg/mL   | none          | [95] |
| Sulfamethazine                 | Milk                  | 1,700 pg/g    | none          | [96] |
| Sulfamethazine                 | Milk                  | <278 pg/mL    | secondary antibody | [97] |
| T-2/HT-2 toxins                | Cereals/baby food     | 25,000-26,000 pg/g | none | [45] |
| Testosterone                   | Saliva                | 15.4 pg/mL    | 25 nm gold    | [34] |
| Tetrodotoxin                   | Buffer                | 300 pg/mL     | none          | [64] |
| Thiamphenicol                  | Shrimp extract        | 500 pg/mL     | none          | [74] |
| Vitamin B-5                    | Food extracts         | 4,400 pg/mL   | none          | [68] |
| Warfarin                       | Plasma ultrafiltrate  | 4,000 pg/mL   | none          | [76] |

### 4. Conclusions

The use of SPR transduction for the immunosensory detection of important small molecule analytes has expanded rapidly over recent years, with particular interest in developing routine testing methods for toxins in foods, and drug residues in both foods and biological fluids. Its use for sensitive detection of steroid hormones and explosive residues has now also been explored. From the research covered in this review, it is evident that, provided a good quality antibody is used, those studies that have most carefully examined the structure of the sensing surface have achieved the best results in terms of both sensitivity and surface stability. Some studies still use direct amine coupling to sensor surfaces without use of intermediate linkers and so may sacrifice sensitivity through steric obstruction of antibody binding. When factors such as linker lengths, conjugation techniques, protein and gold nanoparticle enhancement and regeneration stability are considered and optimized then highly sensitive small molecule immunosensor systems can be developed that can detect in the pg/mL concentration region and overcome the challenges of small analyte size and steric obstruction of binding interactions. In future, it seems likely that the range of small molecule analyte targets examined will expand and that even more attention will be devoted to sensor surface construction, both in terms of the chemical immobilization layers and the underlying metal substrate, particularly given the promise of LSPR techniques for multi-analyte detection. It is hoped that in time SPR immunosensing of small molecules
will become routine in testing laboratories and that portable SPR transduction units will expand the range of applications of plasmonics where on-site and near real-time detection is crucial.

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