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Nanobody-based Immunosensor Detection Enhanced by Photocatalytic-Electrochemical Redox Cycling

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ABSTRACT: Detection of antigenic biomarkers present in trace amounts is of crucial importance for medical diagnosis. A parasitic disease, human toxocariasis, lacks an adequate diagnostic method despite its worldwide occurrence. The currently used serology tests may stay positive even years after a possibly unnoticed infection, whereas the direct detection of a re-infection, or a still active infection remains a diagnostic challenge due to the low concentration of circulating parasitic antigens. We report a time-efficient sandwich immunosensor using small recombinant single domain antibodies (nanobodies) derived from camelid heavy-chain antibodies specific to *Toxocara canis* antigens. An enhanced sensitivity to pg/mL level is achieved by using a redox cycle consisting of a photocatalytic oxidation and electrochemical reduction steps. The photocatalytic oxidation is achieved by a photosensitizer generating singlet oxygen ($^{1}O_{2}$) that, in turn, readily reacts with *p*-nitrophenol enzymatically produced in alkaline conditions. The photocatalor produces benzoquinone that is electrochemically reduced to hydroquinone generating an amperometric response. The light-driven process could be easily separated from the background, thus making amperometric detection more reliable. The proposed method for detection of the toxocariasis antigen marker shows superior performances compared to other detection schemes with the same nanobodies and outperforms by at least two orders of magnitude the assays based on regular antibodies, thus suggesting new opportunities for electrochemical immunoassays of challenging low levels of antigens.

Immunoassays have become one of the most useful techniques in biomedical sciences with countless applications in diagnostics, pharmacokinetics and bioequivalence studies due to the straightforward concept and substitution of radioactive labels by enzymes (leading to ELISA).^{1,2} The analytical performance of immunoassays largely relies on the affinity of antibodies that usually include either polyclonal antibodies (pAbs) obtained from animal immunization or hybridomaderived monoclonal antibodies (mAbs). The latter have the advantage of comprising a homogeneous population of antibodies resulting in more consistent and reproducible results. However, conventional antibodies are composed of heavy and light chains that severely impede their production in bacterial systems. Moreover, the lack of a systematic validation of commercial mAbs resulted in the lack of reproducibility in experiments performed with supposedly the same mAbs and well-known pAbs and mAbs obtained from different antibody resellers.^{3,4} These factors have led to the exploration of alternatives to mAbs with single-chain antibodies that can be successfully expressed in *Escherichia coli* in high yields and with minimum batch-to-batch variations.

The discovery of single-domain antibodies in camelids paved the way for a more straightforward bioengineering, reproducible production of small single domain antibodies, so called nanobodies (Nbs).⁵ Unlike conventional pAbs and mAbs, Nbs derive from the variable domain of heavy-chain antibodies (which are devoid of light chains) and consist of a single polypeptide of ~15 kDa. Thus, Nbs retain full capacity to bind to cognate antigens with the same or even higher affinity compared to mAbs due to an extra-long finger-like shaped complementarity-determining region (CDR3). In addition, Nbs can be reproducibly obtained in high yields in prokaryotic and eukaryotic systems. Nbs can be also used as modular entities in protein engineering and expressed as multimeric fusions, or in combination with enzymes or site-specific tags.

Many Nb-based analytical detection systems have been developed, focusing in particular on medical diagnostics.⁶ Conversely, Nb-based electrochemical immunosensors remain substantially less established and include the detection of procalcitonin,⁷ apolipoprotein-A1,⁸ cystatin C,⁹ aflatoxin B₁,¹⁰ pyrethroid insecticides¹¹ and the toxic microalgae *Alexandrium minutum*¹². The high sensitivity of the electrochemical techniques combined with the high affinity of Nbs affords the detection of analytes at unprecedentedly low levels.

We have recently developed an electrochemical method based on Nbs, HRP-label and magnetic beads for diagnosis of the active form of human toxocariasis (HT), a parasitic disease caused by the accidental ingestion of *Toxocara canis* eggs.^{13,14} Although this is the most sensitive technique to date, the found concentrations of specific antigens in real samples are scattered close to the LOD,¹⁵ warranting further improvement of the method aimed at lowering the LOD.



Figure 1. (A) Conventional detection scheme based on alkaline phosphatase (ALP) labeled antibodies (IgG) with optical or electrochemical readouts. (B) Advanced ELISA based on previously reported Nbs with oriented immobilization. (C) New photoamperometric detection scheme based on the advanced Nbs and the photosensitized oxidation-electrochemical reduction redox cycling (shown in bold).

The detection of active HT infections remains a diagnostic challenge due to the very low level of circulating parasitic antigens (T. canis excretory-secretory antigens, TES) present in the serum during the course of the disease. Furthermore, the only standardized diagnostic tool for HT is serology that detects specific anti-TES IgG by ELISA and western blot.¹⁶ These approaches have the notorious disadvantage of remaining positive even years after infection due to the persisting seropositivity. Therefore, a positive test must be carefully interpreted together with clinical and epidemiological data.¹⁷ In this context we note that alkaline phosphatase (ALP) is one of the most popular enzymatic labels in immunoassays that can be detected either optically or electrochemically (Fig. 1A). Discovery of nanobodies opened new opportunities for bioengineering of advanced immunoreagents (Fig. 1B), while development of electrochemical immunosensors addressed improvements in the reading step. In contrast to optical assays (i.e., ELISA), electrochemical immunosensors retain the ALP

label at the electrode surface. This results in a high steadystate concentration of the enzymatic product at the electrode surface shortly after adding the reagents without the need of a reaction development step. To improve assay performances, the ALP enzymatic substrate and measuring buffer were optimized.¹⁸⁻²⁰ In particular, phosphorylated ascorbic acids¹⁹ and ferrocene derivatives¹⁸ were suggested as alternatives to phosphorylated phenols to avoid electrode fouling. Moreover, enzymatic recycling systems^{21,22} can additionally amplify the ALP assay.²³⁻²⁷ However, a mismatch in the optimal pH for ALP and the recycling enzymes, as well as the generally poor reproducibility and stability of enzymes during and after sensor manufacturing processes limit the practical applicability of such schemes.

We report here the application of our recently introduced concept for detection of phenols using a chemically robust photocatalyst, a robust, fully fluorinated zinc phthalocyanine with bulky perfluoroalkyl peripheral groups.²⁸ In contrast to enzymatic recycling systems, the photocatalyst is fully compatible with the ALP assay conditions. Moreover, the amperometric detection of *p*-nitrophenol (pNP) is conducted at a comparatively low potential due to the photocatalytic oxidation-electrochemical reduction redox cycling (Fig. 1C), which is driven by light and can be switched off on demand to additionally measure the baseline signal. This approach was adopted for the detection of TES to enhance the sensitivity and robustness of the electrochemical immunosensor based on ALP-labeled Nbs.

EXPERIMENTAL SECTION

Preparation of TES, capture and detection nanobodies. Production and purification of TES antigens were carried out as previously reported.²⁹ The concentration of TES in the stock solution was determined from absorbance at 280 nm (OD280 of 1 corresponds to 1 mg/mL) in line with our previous works.¹³⁻¹⁵ Library construction, selection, production and purification of Nbs were done according to Conrath et al.³⁰ and described in detail previously.¹³ Capture Nb (Nb 2TCE49) was prepared as bivalent-monospecific construct in vivo biotinylated by co-expressing pACYC184 plasmid (Avidity, USA) containing pBirACm enzyme. Detection Nb (Nb 1TCE39), which recognizes a different, non-overlapping epitope of TES compared to capture Nb, was expressed from pHEN6c vector containing only a His-Tag. After purification, detection Nb was chemically coupled with ALP using a commercial kit (ab102850, Abcam, UK).

Electrode preparation. The supported photosensitizer (3 wt% $F_{64}PcZn$ on TiO₂ Aeroxide P25) was prepared as described previously.²⁸ A 3 µl of 10 mg/mL suspension of the supported photosensitizer in MQ water was drop-casted on carbon screen printed electrodes (SPE, ItalSens IS-C, Palm-Sens). After drying, the modified electrodes were kept in the absence of light and humidity.

Photoelectrochemical detection of pNP. A pNP (>99%, Fluka) stock solution of 50 mM was prepared in absolute EtOH and added in 0.1 M glycine pH 9.6 buffer containing 1 mM ZnCl₂ and 1 mM MgCl₂. A drop of 90 μ l of pNP solution of variable concentration was placed on the modified SPE electrode and the position of the working electrode was aligned with the laser used for illuminations, a red 660 nm diode laser (Roithner Lasertechnik, Austria) with an output power of 30 mW and spot diameter of 3 mm.

The electrochemical measurements were conducted using a μ Autolab III (Metrohm Autolab, the Netherlands) instrument. The photocurrent was obtained by subtraction of the dark background current recorded just before an illumination. The Ag pseudo-reference electrode has a potential of *ca.* +0.12 V *vs.* SCE in the used glycine buffer.

Determination of ALP activity. The ALP activity was determined spectrophotometrically at 25°C using an AvaSpec-2048L spectrophotometer (Avantes, the Netherlands) in the 0.1 M glycine pH 9.6 buffer containing 1 mM MgCl₂ and ZnCl₂. For the measurements, ALP from calf intestine (ab102850, Abcam, UK) was injected in the measuring buffer containing 1 mM pNPP (> 98%, MP Biomedicals, USA). Then reaction progress was monitored at 410 nm ($\varepsilon_{pNP} = 1.82 \cdot 10^4 \text{ M}^{-1} \cdot \text{cm}^{-1}$) for 5 min. The slope of the initial linear segment was plotted as the function of ALP concentration (in the range 10 – 500 ng/mL). The linear graph corresponded to a specific activity of 46.4 ± 2.5 U/mg (µmol pNP per min per mg ALP).

Photoamperometric detection of ALP. The measurements of the ALP were conducted at room temperature (22°C) in the glycine buffer. ALP was injected in a drop of 1 mM pNPP, which was previously placed on the modified electrode. The photocurrent was measured at continuous or periodic illumination. The slope of the time-dependent growth of the photocurrent was determined, similarly to the optical assay procedure. The slope was plotted as a function of ALP concentration. Using the known sensitivity of the electrodes to pNP, the obtained slopes can be converted into the specific activity of ALP, which, in this way, is obtained independently of the spectrophotometric method.

Photoamperometric TES detection. The detection format used in the present work (Fig. 1C) was adopted from our previous work on electrochemical detection of TES using streptavidin precoated magnetic beads and the same Nbs.13-15 However, in the present work Nb-ALP was used instead of Nb-HRP as detection Nb. In brief, Streptavidin precoated magnetic beads (Dynabeads[™] M-280, ThermoFisher Scientific) were incubated with capture Nb 2TCE49 (2 µg per 1 mg beads) in PBS containing 0.05% Tween 20 (PBST20 buffer) for 1 h on a rotary shaker. Next, the beads were washed 5× with PBST20 and re-suspended in PBST20 to obtain a 5 mg/mL suspension. Detection Nbs were diluted in PBS to get 500 ng/mL. Then, 10 μ l of the modified beads and 10 μ l of diluted detection Nbs were added in each microtube containing 0.5 mL of TES solution, vortexed and placed on a rotary shaker for 30 min. Next, the beads were washed 2× with PBST20 and, if not used immediately, left in PBS without vortexing. Approximately 5 min before the measurements, PBS was replaced with the measuring buffer (0.1 M glycine pH 9.6 containing 1 mM ZnCl₂ and 1 mM MgCl₂) to ensure activation of ALP by Mg²⁺ and Zn^{2+} cations. Just before measurements, the beads were concentrated and transferred in a small amount of measuring buffer $(7-10 \ \mu l)$ on the working electrode covered with a 90 μl drop of measuring buffer containing 0.5 mM pNPP. Photoamperometric measurements were conducted at a fixed potential of -0.32 V using the red light diode laser aligned with the position of the working electrode. For comparison, conventional square wave voltammetry (SWV; amplitude, 0.05 V; frequency, 20 Hz) was used to detect ALP on unmodified SPEs without illumination. All measurements were conducted at room temperature and at least in triplicate.

Detection of TES in serum. Commercially available serum from human male AB plasma (Sigma-Aldrich) was diluted 1/10 with PBS and spiked with TES. Pooled serum of nine highly positive samples (OD in anti-TES-IgG assay > 1.7, from a previous study on the seroprevalence of T. canis in children from Cuba.³¹ Informed consent of the parents or legal guardians of the children was obtained.) containing endogenous immunoglobulins with specificity to TES was used as a control. The serum was diluted 1/10 with PBS and spiked with TES or subjected to the immune-complex dissociation (ICD) procedure, similar to the protocol developed by Rodriguez-Caballero et al.³² Briefly, the diluted serum was mixed with the same volume of 100% glycerol and double volume of 0.5 M EDTA (pH 7.4). Then, the mixture was boiled for 6 min and centrifuged at 19,000×g for 10 min. The supernatant was separated and spiked with TES for photoamperometric detection.

Sandwich ELISA. Sandwich ELISA was performed according to our previously reported optimized protocol^{13,14} but, here, with Nb coupled to ALP. The Nb-ALP conjugate was used in different concentrations, *viz.* 20, 50 or 100 ng/mL with only a minor effect on the result. The conjugate was incubated for 1 h at room temperature with shaking.

In the last step, the wells were washed and the fresh substrate buffer (0.1 M glycine pH 9.6 containing 1 mM MgCl₂, 1 mM ZnCl₂ and 5 mM pNPP) was added to the wells. The reaction developed at 37°C in the dark with shaking. Readings of absorbance at 450 nm ($\varepsilon_{pNP} = 3.8 \cdot 10^3 \text{ M}^{-1} \cdot \text{cm}^{-1}$) were taken after 15, 30, 60 and 90 min. All measurements were done in triplicate, except for blanks which were measured in 6 separate readings. The limit of detection (LOD) was calculated as 3× the standard deviations of blanks divided by the sensitivity in the low concentration range.

RESULTS AND DISCUSSION

pNP photoelectrochemical detection. The amperometric detection of pNP was implemented by employing photosensitized oxidations triggered by the highly active photosensitizer, fully fluorinated phthalocyanine F₆₄PcZn which is capable to form singlet oxygen (¹O₂) under red light illumination.^{28,33} The photosensitized oxidation of pNP yields an oxidized product, benzoquinone (BQ),³⁴ which is electroactive and can be electrochemically transformed into a reduced product, hydroguinone (HQ), giving a cathodic current useful in electroanalysis.²⁸ Since HQ can also undergo the photosensitized oxidation producing BQ, the process forms a redox cycle (Fig. 1C, part of the scheme in bold), thus enhancing the steady-state amperometric responses. The cathodic current from the electrochemical reduction of BQ into HQ is higher when the photooxidation takes place in the vicinity of the electrode surface, which increases the local near-electrode concentration of BQ.

To demonstrate the possibility of photoamperometric detection of pNP in conditions suitable for analysis with ALPlabeled immunoreagents, we used glycine buffer pH 9.6, which is the same buffer as used in the ELISA based on the same immunoreagents. The applied potential is an important parameter of the suggested amperometric detection scheme and affects both the sensitivity to phenols and background currents in blank buffers.³⁵ Fig. 2 illustrates the effect of applied potential on photoamperometric detection of 50 μ M pNP in comparison to the blank buffer. pNP shows a typical wavelike profile with a half-wave potential at -0.27 V and saturation at around -0.40 V (see also Fig. S1 for blank-subtracted data). However, photocurrents in blank buffer also become noticeable at -0.40 V, which required the use of -0.32 V, where sensitivity reaches 80% of the value at -0.45 V, but the blank photocurrent is negligible (Fig. 2, inset).

Figure 2. Light-chopped linear sweep voltammetry of the modified SPE electrode in glycine buffer pH 9.6 in the absence and presence of 50 μ M pNP. Scan rate, 1 mV/s. The inset depicts chronoamperometric behavior at the fixed potential of -0.32 V.



Figure 3. (A) Photoamperometric detection of pNP in glycine buffer pH 9.6 at -0.3 V using red 659 nm laser pointer. (B) Calibration curves constructed for the peak current and steady-state currents (both cathodic) in photoamperometric detection and the near steady-state current (anodic) obtained in chronoamperometric detection at +0.85 V. (C) Zoom in the linear region. Error bars represent SD (n = 3).

Interestingly, minor anodic photocurrents were observed in the blank buffer at potentials more positive than -0.32 V, most likely due to the oxidation of superoxide ions formed via photoinduced electron-transfer reactions involving O2 and side reactions of ¹O₂ with components of the buffer and the modified electrode. A comparatively low formal potential of $E^{\circ}(O_2/O_2) = -0.265 \text{ V} \text{ vs. SCE}^{36}$ (ca. -0.38 V vs. the pseudoreference Ag electrode used), as determined by cyclic voltammetry in 0.1 M NaOH, corresponds well to the region where the anodic photocurrent appears. The cathodic photocurrents in the blank buffer at potentials more negative than -0.32 V were attributed to the reduction of O_2^- , H_2O_2 and likely ¹O₂ generated in the proximity of the electrode (¹O₂ has a limited average diffusion length due to its short lifetime. Less than 0.1% of produced ¹O₂ will diffuse 250 nm in water even in the absence of any quenchers).^{37,38} Nevertheless, the blank photocurrents are rather minor and can be minimized by the careful selection of the working potential. Eventually, the potential of -0.32 ± 0.01 V provided the lowest blank and was used in subsequent work.

The effects of variable pNP concentration. Fig. 3A shows chronoamperograms recorded at -0.32 V in solutions containing 0.01 - 1.0 mM pNP. After switching the light on, a cathodic photocurrent appears and reaches steady-state values within 15 s (5 s in the low concentration range). The pNP photocurrent is linearly proportional to the pNP concentration till 100 μ M, while leveling off at higher values. In contrast to the steady-state current, the peak current (which is observed for

high concentrations shortly after the light is on) keeps growing with the concentration till 500-750 μ M expanding the practically useful concentration range (Fig. 3B). The leveling off at around 500 μ M results from the limited solubility of oxygen (up to 260 μ M) in air-saturated water. The sensitivity in the low concentration range was -0.032 A/M (-0.45 A/M/cm²) with LOD around 0.5 μ M pNP (3xSD).

To estimate the enhancement of the sensitivity due to the redox cycling mechanism, regular chronoamperometric curves were obtained for oxidation of pNP using a sufficiently large potential, +0.85 V vs. the used pseudo Ag reference SPE (+0.97 V vs. SCE), which is 0.15 V more positive than the pNP oxidation peak potential obtained by SWV using the same conditions. A near steady-state current was observed by the time point 180 s. At this reading time, the sensitivity was only 0.09 A/M/cm² with the LOD around 5 μ M (3×SD). Measuring a non-steady-state current at an earlier time, when the current decays steeply, increases the sensitivity (e.g., the sensitivity is twice higher at the reading time of 10 s) but does not improve the actual LOD due to the increased variation in blanks. Thus, the redox cycling enhances the steady-state currents about fivefold compared to the regular amperometric detection while further improving detectability due to intrinsic baseline correction when switching the light on and off. The latter effect resembles the advantage of voltammetry techniques given that baseline correction allows the separation of a net peak current from a baseline current.

Detection of ALP activity correlated with optical assays. ALP dephosphorylates the phenolic group in pNPP releasing pNP, which is of high practical interest in ELISA based on ALP labeled immunoreagents. Moreover, pNPP is electrochemically inactive and thus can be used in electrochemical assays based on the same principle. We additionally discovered that pNPP, in contrast to pNP, does not result in any photocurrent. Thus, the ALP activity can be quantified photoelectrochemically with the setup described here.

Fig. 4A shows the behavior of the current upon the addition of different ALP concentrations under continuous illumination. The current rises in time with a slope proportional to the concentration of ALP, similar to conventional optical assays. To further improve the detectability of pNP and ALP, the light-chopped amperometry was conducted (i.e., with periodic illumination instead of continuous illumination, Fig. 4B). This causes the dependence of the ALP concentration slope to be linear over a wider ALP concentration range (Fig. 4C) with a value of $-1.60 \text{ nA/min/(ng_{ALP}/mL)}$. Taking into account the sensitivity of the electrode to pNP (-0.032 A/M) as obtained above, the slope gives the specific activity of $50 \pm 2 \mu \text{mol/min/mg}_{ALP}$ (µmol pNP formed per min per mg of ALP). Measurements by the conventional optical assay at 25°C in the same buffer resulted in the specific activity of $46.4 \pm 2.5 \mu \text{mol/min/mg}_{ALP}$ that matched well the activity obtained by the electrochemical assay. In these conditions, the LOD for ALP determination by the photoamperometric method and the conventional optical method was almost the same and estimated as 0.2 ng/mL or 0.01 U/L ALP.



Figure 4. Photoamperometric detection of ALP activity in solution under continuous (A) and periodic (B) illumination. (C) Dependence of the slope on the amount of ALP. Background, 0.1 M glycine buffer pH 9.6 containing 1 mM MgCl₂ and 1 mM ZnCl₂. The numbers in the captions of panels (A) and (B) are the amount of ALP in ng/ml.



Figure 5. Effect of measuring time (A), concentration of Nb-ALP conjugate (B) and amount of beads (C) on the detection of 1 ng/mL TES antigens in comparison to blanks.

It is well known that small amounts of Mg^{2+} and Zn^{2+} are necessary for the catalysis of ALP (Mg^{2+} stabilizes the enzyme structure and Zn^{2+} participates in the catalysis). Thus, 1 mM $MgCl_2$ and 1 mM ZnCl₂ were present in the glycine substrate buffer as recommended by the generic ELISA protocol. In the absence of Mg^{2+} and Zn^{2+} or in the presence of only Mg^{2+} , about 25% residual activity was measured photoelectrochemically and 10% was measured optically. This comparatively large drop in activity is explained by the preparation of the initial solutions in PBS buffer, which binds divalent cations. The presence of Mg^{2+} and Zn^{2+} in glycine measuring buffer aims to recover and promote the activity. Noteworthy, some ELISA protocols require only Mg^{2+} in the measuring buffer, which is due to replacement of PBS dilution buffer by Tris dilution buffer to avoid washing of tightly bound Zn^{2+} .

Detection of TES antigens. As anticipated, the sensitivity of the detection step was high starting with the first illumination at 50 s (Fig 5A). The photocurrent responses were usually maximal at the second illumination at 130 s. However, at least 3 consecutive illuminations were conducted in all cases to ensure reproducibility of the data. The concentration of the Nb-ALP conjugate was varied from 2 to 50 ng/mL (Fig. 5B). The responses increased steeply from 2 to 10 ng/mL and lev-

eled off at 25 ng/mL. However, a slight increase of the background was observed at 25 and 50 ng/mL, likely due to unspecific adsorption of the Nb-ALP conjugate on the beads. Thus, 10 ng/mL was taken as the optimal concentration for the calibration curve.

Finally, the amount of magnetic beads used in a single measurement was varied (Fig. 5C). The amount of 50 μ g was optimal in the used setup, which corresponded to our previous work with HRP-labeled Nbs.¹⁴ This amount of beads is visually just enough to cover the complete working electrode, thus, involving most of the available sensor surface in the detection. A thicker layer of beads, which are dark brown may dim the light for the underlying layer of the photosensitizer. The increase from 50 to 100 μ g of beads per electrode indeed hampers the response, decreasing it twofold. An estimation based

on the bead diameter (2.8 μ m), bead density (1.6 g/cm³) and the geometric electrode area (0.071 cm²), as provided by the producers, suggests that 50 μ g of beads per electrode corresponds to the coverage of 0.7 mg/cm² or 7 μ m (2 – 3 monolayers) at random close packing. Nevertheless, the distribution of beads over the surface is not fully uniform. The layer is slightly thicker just above the center of the magnet, i.e. in the middle of the electrode. The use of 50 μ g gives a good compromise between the amount of Nb-ALP on the beads leading to higher sensitivity and dimming the light that reaches the photosensitizer leading to lower sensitivity. In general, the optimal amount of beads is proportional to the electrode area and, thus, can be predicted for electrodes of other diameters by normalization per their electrode area.



Figure 6. Calibration curves for detection of TES antigens obtained by photoamperometric (A, B) and conventional SWV methods (C, D). Numbers in the legends (A,C) denote TES concentration. The insets (B, D) show the actual experimental curves for TES concentration near the LODs. The curves in the inset B are aligned to the same dark current before the illumination. The SWV curves in the inset D are given after subtraction of an average blank followed by a linear baseline correction (using linear fits between 0.2 and 0.5 V).

Calibration curves for TES antigens. Fig. 6A and B illustrate photoamperometric detection of TES in a wide concentration range in comparison with regular SWV technique (Fig. 6C and D) and ELISA with the same reagents (Fig. 7). In all cases, TES was well-detectable at pg/mL levels, which is important for applications since the diagnostically relevant TES concentration is expected to be below 1 ng/mL and, taking into account the necessity of sample preparation, may rather be in the low pg/mL level.^{13,14} In this view, photoamperometric detection is advantageous with an LOD of 1 pg/mL (Table 1) followed by SWV (LOD = 24 pg/mL) and ELISA (LOD = 35 pg/mL).

The other calibration curve parameters also drastically differ for the three methods (Table 1). The usable concentration range was the largest for ELISA (0.05 - 100 ng/mL) with EC₅₀ (half-maximal effective concentration) of 20 ng/mL, which is 7× and 200× higher compared to SWV (EC₅₀ = 3 ng/mL) and photoamperometry (EC₅₀ = 0.1 ng/mL), respectively. This difference, however, should not be attributed to a change in the affinity of the immunoreagents, but to a limitation by the sensors' maximal measurable concentration of pNP (Fig. 3).

It is also interesting to compare sensitivities among the assays with the same Nbs, expressed as slopes of linear correlations in the low concentration range (Table 1). The slope was $5\times$ higher for the photoamperometric sensor with Nb-ALP compared to the amperometric sensor with Nb-HRP.¹⁴ Yet, the slopes for the ELISAs with Nb-ALP and Nb-HRP were the same (around 0.10 a.u./(ng/mL), while also having the identical maximal absorbance of *ca.* 3 a.u. at the highest concentration. This suggests that the performance of Nb-ALP and Nb-HRP is about the same, whereas the enhancement in the sensitivity for the photoamperometric sensor results from the advanced detection scheme. Noteworthy, the LOD was 2× better for ELISA with Nb-ALP compared to the ELISA with Nb-HRP due to only better reproducibility in blanks in the case of Nb-ALP.



Figure 7. Detection of TES antigens by ELISA with the same immunoreagents as used in the electrochemical methods. The inset shows the actual response for a TES concentration closest to the LOD in comparison to the balnk.

SWV allows subtracting a background current to obtain the specific peak current similar to photoamperometry. This is typically done by extrapolating a baseline to the peak position, which is, however, problematic for pNP due to a steep nonlinear background at the peak base. In this work, the shape was partially corrected by subtraction of an average blank curve followed by linear extrapolation of the baseline to obtain the data as in Fig. 6D inset, from which the current at the peak position can be read. This allows objective readings of responses and automatization in data processing, but the eventual LOD is rather high (20x higher compared to photo amperometry, whereas the slope is only $3.6 \times$ lower). If the background correction in SWV could be further improved, for example, by taking an alternative ALP substrate that is oxidized at a lower potential (e.g., *p*-aminophenylphosphate) the performance of the voltammetric sensor would be noticeably improved as well.

It is also possible to compare the slopes of the optical ELISA and the electrochemical sensors by converting the slopes to the same units. First, the a.u. of absorbance can be converted into pNP concentration using the extinction coefficient (3.8-10⁴ M^{-1} ·cm⁻¹) and the optical path (0.3 cm for 100 µl solution in a well of the used ELISA plate). The concentration can be turned into the photocurrent using the sensitivity of the photo amperometric method to pNP (-0.032 A/M as measured earlier in this work). The sensitivity of the ELISA thus corresponds to an equivalent of 3 μ A/(ng/mL), which is 10× lower than the sensitivity of the photoamperometric sensor to TES. Obviously, a longer development time can give a proportionally higher sensitivity in the ELISA at low concentrations with, however, unavoidable saturation of the reading values at high concentrations. In other words, the development time of 15 h (10× standard time) would give a sensitivity comparable to that in the photoamperometric method. In contrast, the high sensitivity of the photoamperometric sensor is reached instantly due to the production of pNP right on the sensor surface

Table 1. Comparison of methods for TES antigens detection.

leading to a high local concentration of pNP. This effect is enhanced by the redox-cycling process and improved baseline correction due to the modulation of the photoreactivity by light.

Detectability of TES in serum. The need of lowering the detectability of TES antigens is met by its detection in spiked blank serum (10× diluted) and serum highly positive for anti-TES-IgG (10× diluted) in comparison to PBS, all in the same experimental conditions (Fig. 8). Not surprisingly, serum affected the detectability of TES, suppressing the sensitivity twofold compared to PBS. More importantly, serum containing large amounts of endogenous antibodies against TES completely suppressed the responses at 10 and 30 pg/ml TES levels. This means that endogenous antibodies block the epitopes recognized by at least one of two Nbs used (the capture and detection Nbs recognize different epitopes). The ICD procedure is generally applied as a pretreatment step to remove endogenous IgG hindering detectability of antigens. After ICD, TES can be indeed detected with reasonably good sensitivity, which is, however, about twice lower than in blank serum and almost five times lower than in PBS (Fig. 8). Notably, TES was not found in non-spiked anti-TES-IgG positive serum despite high seropositivity. In fact, high seropositivity does not necessarily correlate with the level of TES and may even occur a long time after past infection when TES antigens already left the circulation.^{32,39} Nevertheless, the results show that ICD is compatible with our detection method and does not lead to any increase in the background or false negative as the case for highly-positive serum before ICD.



Figure 8. Detection of TES spiked in PBS, blank serum and serum highly positive for anti-TES-IgG before and after the immune-complex dissociation (ICD) treatment. In each case the serum was diluted 1/10 with PBS.

Taking into account the decrease in sensitivity, the sample dilution (4x) in the ICD procedure and possible additional sample dilution due to limited sample volume, a margin of sensitivity by a factor of 100 is expected to ensure the detection of TES in real samples, which corresponds to our previous study on detection of TES using HRP-labeled Nbs leading to an LOD in serum after IDC of 0.9 ng/ml.¹⁵ In the present work the sensitivity has been improved by a factor of 10, allowing detection at 20 – 100 pg/ml depending on the dilution factor.

Method Immunoreagent Slope (in the low concentration rate	Usable concentra- Apparent EC ₅₀ LOD e) tion range (ng/mL) (ng/mL) (ng/mL)	Ref- er- ence
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Photoamperometric	Nb-ALP	31 µA/(ng/mL)	0.003 – 1	0.1	0.001	tw
Voltammetric (SWV)	Nb-ALP	8.6 µA/(ng/mL)	0.03 –10	3	0.024	tw
Amperometric	Nb-HRP	5.8 µA/(ng/mL)	0.01 - 10	1.5	0.006	14
ELISA	Nb-ALP	0.10 a.u./(ng/mL)	0.05 - 100	20	0.035	tw
ELISA	Nb-HRP	0.11 a.u./(ng/mL)	0.3 - 200	20	0.090	14
ELISA	mAb/pAb/IgG-HRP	-	0.1 - 1000	10	0.440	32,40
ELISA	mAb/pAb/IgG-HRP	-	4 - 1000	60	4 ^a	41
ELISA	mAb/pAb/IgG-HRP	-	5 - 100	-	5 ^b	39
ELISA	pAb-HRP	-	78 - 40000	-	78°	42
ELISA	mAb-HRP	-	20 - 25000	-	20 ^c	43,44

^aMeasured in PBST + 3% BSA; ^bmeasured in PBST + 1% BSA; ^cmeasured in serum; tw = this work.

Importantly, although the photoamperometric method does not quantify TES concentrations above 1 ng/mL in PBS, it detects TES down to a few pg/mL – a concentration level that is inaccessible using previously reported ELISAs with conventional polyclonal and monoclonal antibodies (Table 1). These low levels are considerably more relevant for the diagnostic of the disease due to low amounts of TES present in the serum of patients with active infections. In general, the detection of biomarkers and pathogens at trace levels is of high interest in point-of-care and point-of-need diagnostics. Detection at comparatively high levels is routine with ELISA and conventional lateral flow assays. Novel sensors address the need to detect antigens in the low concentration range without the use of high-end laboratory equipment.

Taking into account, the lack of adequate laboratory diagnostic methods and specific clinical picture for covert toxocariasis, the developed setup can be offered for routine laboratory detection of TES. The sample throughput can be improved by the use of an electrochemical multiplexer and precisely positioned LEDs to provide sufficient and reproducible light flux to a set of sensors operating in parallel. Furter development of a point-of-care device, however, is limited by the need for ICD. This issue can be addressed by selecting Nbs that recognize epitopes unavailable for endogenous IgG. Such cryptic epitopes, in theory, can be reached by Nbs due to their unique paratope architecture with a preferential recognition of concave epitopes.⁴⁵ Moreover, production of the disposable photoamperometric sensors requires only low-cost chemo- and thermostable photosensitizers and low-cost procedures such as drop-casting and screen printing leading to little additional costs.

In a broader context, the reported strategy could be adjusted for detection of other parasitic excretory-secretory antigens and other challenging biomarkers or pathogens, including viral antigens (as a better alternative to serological analysis). The importance of the development of highly sensitive methods for the detection of an infectious agent can be exemplified with the rapid tests to identify the SARS-CoV-2 antigens in nasopharyngeal swabs. In contrast to serological tests or RT-PCR, the direct detection of antigens allows the identification of patients exactly during the period where they actively transmit the disease.⁴⁶ However, sensitivity remains an important issue to address. We think that the enhancement established by the photocatalytic-electrochemical redox cycling strategy might extend the applicability of immunoassays and point-of-care devices for detection of antigenic biomarkers present at trace levels.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website.

Light-chopped linear sweep voltammogram of pNP after the subtraction of a blank measurement (PDF)

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All authors have approved the final version of the manuscript.

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