Bienzymatic-based electrochemical DNA biosensors: a way to lower the detection limit of hybridization assays

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The use of the alkaline phosphatase (AP) as an enzyme label and the amplification of its analytical response with a diaphorase (DI) secondary enzyme were investigated in an electrochemical hybridization assay involving arrays of carbon screen-printed DNA biosensors for the sensitive quantification of an amplified 406-base pair human cytomegalovirus DNA sequence (HCMV DNA).

For this purpose, PCR-amplified biotinylated HCMV DNA targets were simultaneously bound to a monolayer of neutravidin irreversibly adsorbed on the surface of the electrodes and hybridized to complementary digoxigenin-labeled detection probes. The amount of hybrids immobilized on the electrode surface was labeled with an anti-digoxigenin AP conjugate and quantified electrochemically by measuring the activity of the AP label through the hydrolysis of the electroinactive p-aminophenylphosphate (PAPP) substrate into the p-aminophenol (PAP) product. The intensity of the cyclic voltammetric anodic peak current resulting from the oxidation of PAP into p-quinoneimine (PQI) was related to the number of viral amplified DNA targets present in the sample, and a detection limit of 10 pM was thus achieved. The electrochemical response of the AP label product was further enhanced by adding the diaphorase enzymatic amplifier in the solution. In the presence of the auxiliary enzyme DI, the PQI was reduced back to PAP and the resulting oxidized form of DI was finally regenerated in its reduced native state by its natural substrate, NADH. Such a bienzymatic amplification scheme enabled a 100-fold lowering of the HCMV DNA detection limit obtained with the monoenzymatic system.

Introduction

In the past decade, the detection of PCR-amplified DNA sequences through their hybridization has become an increasingly implemented method in the diagnosis of pathogenic organisms present in clinical, food and environmental samples. In this context, DNA biosensors coupling the inherent specificity of DNA recognition reactions with the sensitivity of transducers have been the subject of intense research activity due to their ability to provide the sequence-specific information in a more rapid and simplistic manner as compared to the traditional hybridization assays. Various transduction methods have been used to monitor nucleic acid binding events, including fluorescence,2 surface plasmon resonance,3 piezoelectric,4 and electrochemical techniques.5 Among them, electrochemistry has drawn great attention over other conventional methods for decentralized screening of infectious agents owing to its high sensitivity, low cost, rapid response, small dimensions, low power requirements, and compatibility with microfabrication technology. Different strategies for the detection of the DNA recognition event based on the intrinsic electroactivity of the nucleic acids,6 redox-active hybrid indicators,7 metal complexes,8 gold nanoparticles,9 or enzyme labels10 have been reported so far.

However, due to the intrinsic signal amplification provided by biocatalytic reactions, enzyme labels were involved in most of the amperometric detection schemes of DNA hybridization. The transduction of the enzyme activity into an electrochemical signal can result from either the catalytic conversion of a substrate to an electroactive product or a redox-mediated electrocatalytic transformation. This has been demonstrated with various enzyme labels such as horseradish peroxidase (HRP),10,11 alkaline phosphatase (AP),12 PQQ-dependent glucose dehydrogenase,13 bilirubine oxidase,14 glucose oxidase15 and esterase26 which allowed nano- to femtomolar detection limits of nucleic acids in solution to be achieved.

Recently, we investigated neutravidin-coated carbon screen-printed DNA sensors based on a mediated HRP label and showed that PCR-amplified human cytomegalovirus (HCMV) DNA fragments could be determined at the picomolar level.16 Molecular diagnosis of HCMV is commonly based on the PCR coupled with a detection method due to the low abundance of viral DNA in biological samples. Despite their undeniable efficiency, the PCR amplification techniques are expensive, time-consuming and not free from error (false positive). Hence, to provide an alternative for PCR methods while meeting the demands of clinical diagnostics for sensitivity and cost-effectiveness, the amperometric response of enzymatic DNA sensors must be undoubtedly further enhanced. A large variety of
strategies have been developed for amplifying enzyme-based amperometric responses including wiring the enzyme to an electron conducting polymer,\textsuperscript{16} the use of multiple enzymes per binding events,\textsuperscript{17} metallization catalyzed by an enzyme,\textsuperscript{18} recycling or accumulating the reaction product\textsuperscript{19} and the coupling of two enzyme labels through substrate or cosubstrate regeneration.\textsuperscript{19} Though this latter approach seems attractive for the development of highly sensitive hybridization assays, modest nanomolar detection limits of nucleic acids targets in solution have been reached to date with the glucose oxidase-HRP\textsuperscript{15} and glucose-6-phosphate dehydrogenase-diaphorase\textsuperscript{20} coupled systems. Recently, high amplification rates (>1000) of the amperometric responses of aminophenolic products generated either by AP\textsuperscript{20} or by β-galactosidase\textsuperscript{24} enzyme labels were obtained either by adding the Diaphorase (DI) from \textit{Bacillus stearothermophilus} in the solution or by co-immobilizing the DI enzyme on the electrode surface, respectively. In such biennial systems, the phenolic enzyme products can be oxidized at the electrode surface to give quinonimine derivatives according to a $-2e^- \rightarrow 2H^+$ reaction. In the presence of the second enzyme DI, the quinonimines are reduced back to aminophenols and the oxidized form of DI is finally regenerated in its reduced native state by its natural substrate, NADH.

The goal of the present work is to further extend the scope of this biennial electrochemical detection approach to the analysis of HCMV DNA hybridization. For such a purpose, we took advantage of our earlier optimized hybridization assay protocol involving easy-to-use disposable reproducible (relative standard deviation of 9\%) arrays of neutravidin-coated carbon screen-printed sensors.\textsuperscript{16} Briefly, the 406-base pair amplified HCMV DNA sequence targets were obtained by PCR amplification with biotinylated primers to produce biotinylated targets, which can be directly bound to the neutravidin-coated-electrodes, thus avoiding the use of an intermediary capture probe. The immobilization of the DNA targets and their hybridization with digoxigenin-labeled detection probes were performed in a single step and followed by alkaline phosphatase labeling with an anti-digoxigenin antibody conjugate (anti-Dig-AP). Alkaline phosphatase was selected as the primary enzyme label in association with the $p$-aminophenylphosphate (PAPP)/$p$-aminophenol (PAP) substrate/product couple. The first part of this paper is devoted to the study of the analytical performances of the biosensor for the detection of an amplified viral DNA sequence with the AP label. Then, we will show how the amperometric response of AP, and thus the sensitivity of the assay, can be further enhanced in the presence of DI enzyme.

**Experimental**

**Reagents and solutions**

The 5’-biotinylated primers (Bio-AC1 and Bio-AC2) used to PCR-amplify the 406-bp HCMV target and the 5’-digoxigenin-labeled HCMV target specific detection probes (Dig-AC3, Dig-AC4, Dig-B1) were a gift from Argene SA. All of these oligonucleotide sequences are the proprietary of Argene SA, as well as the Hybridowell™ kit whose reagents (hybridization and washing buffers, ETS2 negative control) were used in this work. Taq polymerase, Taq polymerase buffer and the four nucleotide bases (dNTPs) were purchased from Qiagen (France). A low DNA Mass Ladder\textsuperscript{4} for electrophoresis quantification was obtained from Invitrogen (France).

Lyophilized Neutravidin\textsuperscript{a} and lyophilized biotinylated alkaline phosphatase (Bio-AP) were purchased from Pierce (USA). Lyophilized Fab fragments from anti-digoxigenin antibody from sheep, conjugated with alkaline phosphatase (anti-Dig-AP) were obtained from Roche Diagnostics GmbH (Germany). Lyophilized Diaphorase (DI) from \textit{Bacillus stearothermophilus} (E.C. 1.6.99.-) was provided from Unitika (Japan). NADH, Bovine Serum Albumine (BSA), tris-(hydroxymethyl)aminomethane (Tris), magnesium chloride, ferrocenyl methanol, $p$-nitrophenylphosphate, and NaBH$_4$ were supplied by Sigma-Aldrich (France). Tween 20 was obtained from ProLabo (France). $p$-Aminophenylphosphate (PAPP) was obtained by chemical reduction of $p$-nitrophenylphosphate with NaBH$_4$ according to the protocol described in ref. 22.

Phosphate-buffered saline (PBS; 4.3 mM NaH$_2$PO$_4$, 15.1 mM Na$_2$HPO$_4$ and 50 mM NaCl; pH 7.4), Tris buffer (TB; 0.1 M Tris, 0.2 g L$^{-1}$ MgCl$_2$; pH 8.5) and all of the solutions were prepared with water purified by an Elgastat water system (Elga, France). Stock solutions of 2 mM PAPP and 8 mM NaBH$_4$ were daily prepared in PBS and stored at 4°C.

**Instrumentation and electrochemical measurements**

Cyclic voltammetry (CV) measurements were performed with an Autolab potentiostat (PGSTAT 12, Ecochemie) connected to a personal computer equipped with a GPES version 4.9 software. Disposable arrays of eight screen-printed carbon electrodes were prepared from a high impact polystyrene substrate (Sericol, Vaux-en-Velin, France) and a conductive carbon-based ink (Electrodag® PF 407A, Acheson Colloids) using a Presco screen-printing machine (USA). After a curing step, two insulating layers (Vinylfast 36–100, Argon) were printed over the array of eight electrodes, leaving uncovered sensing disk areas of 4.9 mm$^2$ and the electric contacts. The resulting ring-shaped layers around the working areas constituted the reservoirs for small-volume electrochemical biosensing cells. All the electrochemical measurements were carried out at room temperature in TB with a working volume of 20 µL and involved an Ag/AgCl wire reference and a platinum wire counter electrodes.

**Preparation of the biosensing surfaces**

Unless otherwise stated, all the incubations were performed at room temperature in a water-saturated atmosphere. A drop of 20 µL of a 0.5 mg mL$^{-1}$ saturating neutravidin solution in PBS was placed onto each working electrode surface and incubated for 2 h. The surface of each sensor was then carefully rinsed with PBS to remove the excess of neutravidin and the array of sensors was dipped in a 20 mL bath solution containing PBS with 0.1% (w/v) BSA (PBS-BSA) for 30 min. After another thorough wash in a PBS bath, the resulting biosensing platform was stored in PBS at 4°C until used.

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\textsuperscript{a} Lyophilized Neutravidin (2 mg/mL) was obtained from Invitrogen (France).
HCMV DNA hybridization assay procedure

Unless otherwise stated, all the incubations were performed at 37 °C in a water-saturated atmosphere. Each assay was performed onto the eight neutravidin-modified positions of the array using biotinylated target sequences according to our previously reported protocol.11c

Briefly, HCMV DNA was extracted from cell culture, amplified by PCR using biotinylated primers, and then quantified by agarose gel electrophoresis. After purification and quantification, the double-stranded 406-bp DNA samples were thermally denatured by heating for 20 min at 95 °C, diluted in the PCR negative control with concentrations ranging from 10^{-15} to 10^{-7} M, then cooled in an iced bath. Meanwhile, complementary probes tagged with a digoxigenin label (Dig-B1, Dig-AC3 and Dig-AC4) were prepared at a final concentration of 1 μM in the hybridization buffer and then mixed with the DNA sample (1:1 v/v). 20 μL droplets of the resulting mixture were applied onto the active surface of the biosensor array and incubated for 1 h. Each series of experiments included the analysis of a PCR negative control (containing all of the reagents, except DNA) and a noncomplementary DNA-amplified sequence (a human ETS2 DNA gene). After a washing cycle consisting of five rinses for 1 min in a 20 mL bath of fresh 1 x washing solution, the active surfaces of the array were covered with drops of 20 μL containing anti-Dig-AP (1/100 dilution in PBS-BSA containing 0.1% (v/v) Tween 20) and incubated at 37 °C in a water-saturated atmosphere for 30 min. A last washing cycle was then performed with PBS-BSA containing 0.1% (v/v) Tween 20 followed by PBS (five 20 mL baths for 1 min each with each buffer). Once carefully removed from the rinsing solution, the electrodes were stored in a 20 mL TB bath at 4 °C. The detection of the activity of the AP label and its amplification in the presence of DI were carried out according to the following two protocols. (1) 20 μL droplets of TB containing 1 mM PAPP were deposited onto the surface of each sensor. After a 20 min enzyme incubation period, the generated PAPP was determined by CV (v = 10 mV s^{-1}) and the resulting anodic peak current (i_1) at ~ +0.25 V vs. Ag/AgCl was taken as the analytical response. (2) After a thorough rinse with TB, the above AP detection procedure was then repeated with a 1 mM PAPP mixture containing 2 mM NADH and 50 nM Diaphorase and the electrodeoxidation current response (i_2) was measured on the CV curve (v = 10 mV s^{-1}) at +0.30 V vs. Ag/AgCl.

Results and discussion

AP-based electrochemical hybridization assay of HCMV DNA

The main procedure for the detection of amplified HCMV DNA is illustrated in Fig. 1A.

The immobilization of the biotinylated amplified 406-bp DNA fragments onto SPEs covered with a monolayer of neutravidin and their hybridization with complementary digoxigenin-labeled oligonucleotide probes have been performed as described in a previous work.11c The extent of hybrids formed was then determined by incubating an anti-digoxigenin AP conjugate where the optimal concentration was assessed in a series of preliminary experiments. Thereafter, as shown in Fig. 1B, the enzymatic reaction with the substrate PAPP deposited on the SPE surface was allowed to proceed and next the generated phenolic electroactive product was quantified by CV. The magnitude of the anodic peak current (i_1), which corresponds to the oxidation of PAPP into p-quinonimine (PQI) according to a (2e^- + 2H^+) reaction, is proportional to the amount of anti-Dig-AP anchored to the immobilized hybrids, and thus indirectly to the target DNA concentration initially present in the sample solution.

A main and general problem encountered with AP-based assays is the substrate blank current generated in the absence of AP. The purity of the PAPP was thus first examined and a background current of 15 ± 2.5 nA corresponding to the residual traces of PAP contained in a 1 mM TB solution of the synthesized PAPP was recorded. The minimal detectable current in this study was thus equal to 23 nA based on three times the standard deviation of the blank response. The sensitivity of the assay was investigated by varying the concentration of biotinylated amplified HCMV DNA products over the 10^{-15}–10^{-7} M range and the corresponding logarithmic standard plot (curve A) is shown in Fig. 2.

With the aim to further compare the mono- and the bi-enzymatic approaches, the current response i_1 was normalized to the blank response (sample which did not contain HCMV-amplified DNA fragments). The linearity range was extended over ca. 2 decades (from 0.1 to 20 nM HCMV DNA) and the signal saturated above 20 nM, owing to the limited amount of 406-bp HCMV DNA fragments immobilized onto the electrode surface i.e., one third of a packed monolayer according to our earlier estimation.11c A small baseline signal was recorded in the absence of HCMV DNA, thus indicating a very low nonspecific binding. The selectivity of the assay was also confirmed since a negligible nonspecific response was obtained when replacing the HCMV DNA target with a noncomplementary biotinylated amplified human ETS2 fragment. Finally, a detection limit of 10 pM (6 × 10^7 copies of HCMV-amplified DNA fragments per electrochemical cell) could be estimated using a signal-to-noise ratio.
of 3 (S/N = 3), which is 3 times lower than that previously obtained with the HRP label for the same assay format.\textsuperscript{11} These results are in good agreement with the kinetic parameters of each label since the AP has a 4-fold higher turnover value for PAPP\textsuperscript{13} than the HRP toward its osmium cosubstrate\textsuperscript{24} (k\textsubscript{cat AP} = 1150 s\textsuperscript{-1} and k\textsubscript{cat HRP} = 280 s\textsuperscript{-1}). Moreover, the HCMV DNA detection limit obtained in the present work competes favourably with other AP-based electrochemical DNA biosensors recently reported for the detection of PCR-amplified DNA products such as GMO-related sequences (1 nM or 6 x 10\textsuperscript{9} copies of a 195 bp region of the 35S promoter).\textsuperscript{12} Herpes simplex, Epstein-Barr and cytomegalovirus sequences of human viruses (2 nM or 7.3 x 10\textsuperscript{9} copies of viral DNA from a multiplexed PCR).\textsuperscript{12}

**Amplification of the AP electrochemical hybridization response with DI**

As sketched in Fig. 1C, the PAP generated by the immobilized AP label can be subsequently cycled in a redox reaction between PAP and PQI in the presence of the DI enzyme and its NADH substrate. The more practicable approach that consists in adding a DI concentration of 50 nM was selected since it provided both a measurable amplification rate. The calibration plots obtained with the mono- (curve A) and the bi- (curve B) enzymatic detection over the 10\textsuperscript{–15}–10\textsuperscript{−7} M HCMV DNA range are shown in Fig. 2. While both curves exhibited roughly the same signal saturation shape above 10 nM HCMV DNA, no linearity range could be accurately defined when working with the bi-enzymatic system. Though the nonlinear shape of curve B remained unexplained, the use of the DI enzyme amplifier significantly led to a wider working range (10\textsuperscript{–12}–10\textsuperscript{−8} M instead of 10\textsuperscript{–10}–10\textsuperscript{−8} M) and to a ~ 10-fold increase in sensitivity (calculated from the irreversibility of PAP at protein-covered screen-printed electrode surfaces—made of a random distribution of closely spaced insulating and conductive microscopic zones—combined with the contribution of the NADH electrochemical oxidation. At high DI concentrations, the progressive conversion of the pseudo plateau-shaped to a peak-shaped response is indicative of the passage from control by the kinetics of the enzymatic reaction to control by substrate diffusion. Since the sensor response was not obvious to define, similar experiments were conducted with the reversible ferrocenyl methanol (FeMeOH) cosubstrate to better assess the electrochemical oxidation of NADH. Since its contribution has occurred just after +0.3 V, the electrocatalytic current value measured at +0.3 V vs. Ag/AgCl was selected as the analytical response. The linear variation of i\textsubscript{2}—obtained under kinetic control—against the square root of the DI concentration (inset of Fig. 3) is in agreement with previously established results with the FeMeOH cosubstrate\textsuperscript{25} and clearly indicates that the PAP response, and thus the AP amplification rate, can be improved by raising the DI concentration in the solution.

Consequently, the concept of using a DI auxiliary enzyme to amplify the amperometric response of the AP label was evaluated for the determination of HCMV DNA sequences in the hybridization assay reported in the previous section. For this purpose, a DI concentration of 50 nM was selected since it provided both an acceptable value for the blank signal (i\textsubscript{2} = 100 nA) and a measurable amplification rate. The calibration plots obtained for the mono- (curve A) and the bi- (curve B) enzymatic detections over the 10\textsuperscript{−15}–10\textsuperscript{−7} M HCMV DNA range are shown in Fig. 2. While both curves exhibited roughly the same signal saturation shape above 10 nM HCMV DNA, no linearity range could be accurately defined when working with the bi-enzymatic system. Though the nonlinear shape of curve B remained unexplained, the use of the DI enzyme amplifier significantly led to a wider working range (10\textsuperscript{−12}–10\textsuperscript{−8} M instead of 10\textsuperscript{−10}–10\textsuperscript{−8} M) and to a ~ 10-fold increase in sensitivity (calculated from the irreversibility of PAP at protein-covered screen-printed electrode surfaces—made of a random distribution of closely spaced insulating and conductive microscopic zones—combined with the contribution of the NADH electrochemical oxidation. At high DI concentrations, the progressive conversion of the pseudo plateau-shaped to a peak-shaped response is indicative of the passage from control by the kinetics of the enzymatic reaction to control by substrate diffusion. Since the sensor response was not obvious to define, similar experiments were conducted with the reversible ferrocenyl methanol (FeMeOH) cosubstrate to better assess the electrochemical oxidation of NADH. Since its contribution has occurred just after +0.3 V, the electrocatalytic current value measured at +0.3 V vs. Ag/AgCl was selected as the analytical response. The linear variation of i\textsubscript{2}—obtained under kinetic control—against the square root of the DI concentration (inset of Fig. 3) is in agreement with previously established results with the FeMeOH cosubstrate and clearly indicates that the PAP response, and thus the AP amplification rate, can be improved by raising the DI concentration in the solution.

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comparison of the slope of the two calibration plots using linear scales). A detection limit as low as 100 fM could be estimated (6 × 10^5 copies of HCMV-amplified DNA fragments per electrochemical cell), thus yielding a 100-fold improvement in the detection limit of the AP-based HCMV DNA assay. To the best of our knowledge, the number of copies of DNA targets detected in the present work is roughly 160 times lower than those reported for other bienzymatic-based electrochemical hybridization assays.19

Conclusions

The convenient use of arrays of screen-printed DNA biosensors in association with the electrochemical detection of the AP enzyme label with the PAPP substrate led to the specific and sensitive detection of HCMV-amplified DNA. The obtained detection limit (6 × 10^5 copies of HCMV-amplified DNA fragments per electrochemical cell) is very competitive with other monoenzymatic electrochemical DNA sensors recently reported for the detection of PCR-amplified DNA products.12–A26

Especially, it has also been demonstrated that a greater level of sensitivity can be reached by amplifying the AP amperometric response of DNA biosensors with a DI auxiliary enzyme. This bienzymatic detection enabled a 100-fold lower HCMV DNA detection limit to be estimated. While the novel approach described in this work is promising, this detection limit is 700-fold higher than in our previous gold-amplified electrochemical transduction of oligonucleotide hybridization in polystyrene microwells with screen-printed microband electrodes (840 sequences per microwell).10 The length of the target (406 bp instead of 25 bp), the assay format (biosensor instead of microwell + microelectrode) are the main reasons to explain such differences. Hence, further improvements are still required to detect as low as hundreds of copies of target DNA per few tenths of microliters of sample with a bienzymatic system. This can be readily envisaged by co-immobilizing the DI enzyme on the electrode surface and/or using another aromatic monoester phosphate substrate with a very low level of residual traces of phenol.21

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