Nitric Oxide Detection with Glassy Carbon Electrodes Coated with Charge-different Polymer Films

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Abstract: Trace amounts of nitric oxide (NO) have been determined in aqueous phosphate buffer solutions (pH=7.4) by using a glassy carbon electrode coated with three charge-different polymer films. The glassy carbon electrode was coated first with negatively charged Nafion film containing tetrakis(pentafluorophenyl)porphyrin iron(III) chloride (Fe(III)TPFPP) as the NO oxidation catalyst, and then with positively charged poly(acrylamide-co-diallyldimethylammonium chloride) (PADDA) and with neutral poly(dimethylsiloxane) (silicone) at the outermost layer. This polymer-coated electrode showed an excellent selectivity towards NO against possible concomitants in blood such as nitrite, ascorbic acid, uric acid, and dopamine. All current ratios between each concomitant and NO at the cyclic voltammogram was in 10⁻³ ~ 10⁻⁴. This type of electrode showed a detection limit of 80 nM for NO. It was speculated from the electrochemical study in methanol that high-valent oxo-iron(IV) of Fe(TPFPP) participated in the catalytic oxidation of NO.

Keywords: polymer-coated electrode, nitric oxide detection, tetrakis(pentafluoro-phenylporphyrin) iron(III) chloride.

Introduction

Nitric oxide (NO) plays significant roles in the microbiology, physiology and chemistry of an environment. Since NO itself became to be known as an endothelial-derived relaxing factor [1,2], it has been clarified that NO
worked as a neurotransmitter in the brain [3], and as an antitumor agent [4]. The determination of NO in biological media is of great importance but it is complicated by the short life due to the high reactivity of NO. Several methods have been implemented for measuring NO [5]. Among these methods, the electrochemical one is the most suitable for in-situ NO detection. Direct electrochemical reduction or oxidation of NO can be achieved on bare metals or carbon materials [6,7]. However, many substances, which disturb the detection of NO, are present in vivo. Coating of the electrodes with a Nafion membrane prevents side reaction of nitrite oxidation, but can not prevent the interference by positively charged substance in blood, for example, dopamine. In order to improve the selective detection of NO, several polymer films have been coated onto base electrodes, and are summarized in Bedioui and Villeneuve’s review [8]. In these polymer films, WPI membrane [9] has been shown to be effective for selective NO detection, but the nature of the membrane is not indicated. Single layer membrane on electrodes [10,11] has been claimed to be effective for selective detection of NO, we tried to prepare an electrode coated with three charge-different polymer films and checked its performance for the selective detection of NO. Since Malinski and Taha [12] first clarified that electropolymerized Ni(II)porphyrin was effective for the high-sensitive detection of nitric oxide, many metallloporphyrins [13-20] had been investigated to clarify their functions in the catalysis. Present authors also studied on various iron porphyrins [18] and found that tetrakis(pentafluorophenylporphyrin) iron(III) chloride (Fe( TPFPP)Cl) was an excellent catalyst for nitric oxide oxidation. We utilized this porphyrin as the oxidative detection of nitric oxide in this work.

**Experimental**

**Chemicals**

As the polymer for electrode coating, negatively charged Nafion alcoholic solution (Aldrich, 5 wt%), positively charged poly(acrylamide-co-diallyldimethylammonium chloride) (PADD A) aqueous solution (Aldrich, 10 wt%), and neutral poly(dimethylsiloxane) in xylene (Toray-Dow, KE-9140, 60 wt%) were used. The latter two solutions were diluted to 5 wt% and 10 wt% with the original solvent, respectively.

Nitric oxide (NO) (Nippon Sanso) diluted with Ar (5% NO) was bubbled into the electrochemical cell for 20 min. The concentration of NO in solutions was estimated to be 87.5 µM [5]. The NO solutions with lower concentrations than 87.5 µM were prepared by adding a constant amount of the above solution into 10 ml phosphate buffer in a glove box filled Ar. Ar gas for purging dissolved oxygen in electrolytic solution and in 5% KOH for purification line of NO was of ultra-pure (Nippon Sanso, 99.9999 %). This gas was passed for 30 min before passing NO.

**Preparation of Electrode**

A glassy carbon rod with 3 mm φ embedded in a phenol resin pipe with 6 mm φ was used as the electrode. The electrode was cleaned by polishing with 0.06 µm alumina and then washing in an ultrasonic bath. The surface was at first coated with a known volume of Nafion solution (5 wt%) as to be 16 µm after drying, and
next with an aqueous solution of PADDA (5 wt%), and finally with a xylene solution of silicone (10 wt%), as shown in Fig. 1. This coating order was selected to avoid mixing of each two polymers. In the estimation of film thickness, volume of polymer solution, coating area, and the density of polymer in solid state were taken into account. The density of polymer has been assumed to be 1.0 for Nafion, 1.2 for PADDA, and 1.0 for silicone, respectively, on the basis of each technical data. Fe(TPFPp) as the NO oxidation catalyst was originally dissolved in the Nafion solution, and the concentration was adjusted to 10 nmol per dried Nafion 1 mg [18].

![Figure 1](image.jpg)

**Figure 1.** Structure of a glassy carbon electrode coated with charge-different polymer films.

**Electrochemical Measurements**

Electrochemical measurements were carried out in a one-compartment cell with three electrodes by using an electrochemical analyzer (BAS, model 100B/W). A glassy carbon disk (0.071 cm$^2$) coated with polymer films, an Ag|AgCl in 3M NaCl (all potential values are given against this electrode), and Pt plate were used as working, reference, and counter electrode, respectively. Cyclic voltammograms of water-insoluble Fe(TPFPp)Cl was done in 0.1 M methanol solution of tetrabutylammonium perchlorate (TBAP). A Ag|Ag$^+$ was used as the reference electrode in this solution. The potential was 0.20 V against the above Ag|AgCl electrode. Effect of hydroxide on the cyclic voltammograms of Fe(TPFPp)Cl was examined by adding small amount of tetrabutylammonium hydroxide aqueous solution.

**Results and Discussion**

**Effect of the Thickness of Polymer Film**

We determined the most suitable film thickness of three charge-different polymers as following. Suitable film thickness of Nafion for NO detection has already been studied in Ref. [18], and is found to be 16 µm. This thickness was utilized in the present study. Then, we checked the second layer of positively charged PADDA
by changing the thickness under the fixed thicknesses of Nafion (16 µm) and silicone (15 µm). Suitable film thickness of PADD A was determined by the electrochemical reaction of dopamine as a possible interfering substance in vivo.

**Figure 2.** Cyclic voltammograms of 1 mM dopamine in 50 mM PBS (pH=7.4) at a bare GC (a) and at a GC|Nafion (16 µm)|PADD A (b, 3 µm; c, 5 µm)|Silicone (15 µm) electrode. Scan rate, 20 mV s⁻¹.

**Figure 3.** Cyclic voltammograms of 87.5 µM NO in 50 mM PBS (pH=7.4) at a GC|Nafion(16 µm) (a), and at a GC|Nafion (16 µm)|PADD A (5 µm)|silicone (b, 15 µm; c, 23 µm; d, 46 µm) electrode. Scan rate, 20 mV s⁻¹.
Figure 2 shows the result. PADDA of 3 µm thickness (Fig. 2a) could not inhibit the penetration of dopamine, but 5 µm thickness (Fig. 2b) were able to completely inhibit its penetration. The thickness more than 5 µm showed almost the same cyclic voltammogram as that of 5 µm.

On the other hand, thin silicone film is necessary for high sensitive detection of NO. However, the film thickness below 15 µm showed many cracks on the film. So, an effect of the thickness on NO detection was studied by preparing following electrode: GC | Nafion (16 µm) | PADDA (5 µm) | silicone with different thickness. The result is shown in Fig. 3. Increasing in the thickness gave decreased response to the NO oxidation. As the result, we selected 15 µm for the silicone film at the outermost layer.

Selective Detection of NO

By using the above polymer film-coated GC electrode, it was checked the selectivity of NO for nitrite, dopamine, ascorbic acid, and uric acid as possible concomitants in vivo. The cyclic voltammograms of 1 mM each concomitant on bare GC, on GC|Nafion (16 µm), and on GC|Nafion (16 µm) |PADDA (5 µm) |silicone (15 µm) electrode were measured. In this experiment, negatively charged nitrite, ascorbic acid, and uric acid were completely deactivated on (b) Nafion-coated GC electrode. On the other hand, positively charged dopamine was not deactivated at a GC|Nafion electrode, in which the oxidation peak potential in the cyclic voltammogram was shifted from 0.2 V to 0.5 V (vs. Ag/AgCl) by the Nafion coating, but the peak current and the shape of wave were almost the same as those at a bare GC. However, dopamine was almost deactivated on the triply polymer film-coated GC electrode, as has already been shown in Fig. 2c.

Performance of the GC|Nafion|PADDA|silicone electrode was summarized in Table 1. Each oxidation peak current has been evaluated by subtracting the base current without the substance. An effect of four possible concomitants on NO detection was found to be almost negligible at the present triply polymer-coated electrode.

Table 1. Oxidation peak currents of NO and interfering compounds observed at a bare and at a triply polymer film-coated GC electrode and their ratio.

| Compound                  | Oxidation Peak Current at Different Electrodes / µA | Peak Current Ratio between (a) and (b) |
|---------------------------|----------------------------------------------------|---------------------------------------|
|                           | (a) Bare GC (b)Polymer Film-coated GC              |                                       |
| NO (87.5 µM)              | 2.49 *                                              | 2.32                                 |
| Nitrite (1 mM)            | 32.41                                               | 2.60×10⁻³                            |
| Ascorbic Acid (1 mM)      | 12.12                                               | 2.70×10⁻³                            |
| Uric Acid (1 mM)          | 16.52                                               | 2.70×10⁻³                            |
| Dopamine (1 mM)           | 14.71                                               | 1.03×10⁻²                            |

* Nafion (16 µm) was coated.
Amperometric Detection of Nitric Oxide

By using the triply polymer-coated GC electrode, amperometric detection of nitric oxide in deoxygenated phosphate buffer with pH=7.4 was carried out. Aliquots of phosphate buffer saturated with 5% NO (87.5 µM) was successively injected into the cell containing 10 ml phosphate buffer to attain 87.5 nM ~ 878 nM. The potential of the working electrode was maintained at 0.9 V (vs. Ag|AgCl). Figure 4 shows typical amperograms measured at triply polymer film-coated GC electrode with successive addition of NO.

**Figure 4.** Typical amperograms measured at a GC|Nafion (16 µm)|PADDA (5 µm)|silicone (15 µm) electrode with successive injection of NO.

**Figure 5.** Detection curve of NO obtained by amperometry at a GC|Nafion (16 µm)|PADDA (5 µm)|silicone (15 µm) electrode.
These amperograms clearly show an increase in oxidation currents with NO injections. The relation between the increment of current after injection and that of NO concentration was roughly linear in the range of concentrations investigated, as shown in Figure 5. The limit of NO detection at a GC electrode coated only with 16 µm Nafion (Fe-TPFPP) film was 18 nM [18]. Although the limit became high by the triple layer polymer coating, NO of 80 nM was detected by the present electrode. Time elapsed before peak current after NO injection in Fig. 4, so-called response time, was roughly 15 s. This time also became longer, compared with 1 ~ 2 s at a GC | Nafion (16 µm) electrode [18]. A higher detection limit of NO at the present electrode seems to result from longer response time through increased film thickness. It took about 30 min for the measurement of amperograms in Fig. 4. In the second and third cycles after exchanging the electrolytic solution, we observed almost the same amperograms as those in the first one. Stability and reproducibility of NO detection by this electrode were fairly good, but we did not check long-term stability.

Catalysis by Fe(TPFPP) for NO Oxidation

Various iron porphyrins have been clarified to show catalytic activity for nitric oxide oxidation [18]. Especially, water-soluble iron porphyrin, iron(III) meso-tetrakis(N-methylpyridinium-4-yl) (Fe(TMPyP)), showed a selective oxidation of NO against nitrite. This selectivity has been ascribed to different catalyses by different high-valent iron complexes, namely oxo-iron(IV) for NO oxidation to nitrite and oxo-iron(IV) π-cation radical for nitrite oxidation to nitrate [21].

![Figure 6. Cyclic voltammograms of 4x10⁻⁴ M Fe(TPFPP)Cl at a GC electrode in 0.1 M TBAP methanol solution (a) without and (b) with 4x10⁻⁴ M tetrabutylammonium hydroxide. Scan rate, 100 mV s⁻¹.](image-url)
It is not so easy to clarify the catalysis of water-insoluble Fe(TPFPP) in Nafion film being insulator. So, it was examined to estimate the catalysis by dissolving Fe(TPFPP) into methanol. Figure 6 shows the cyclic voltammograms of Fe(TPFPP) in methanol with and without tetrabutylammonium hydroxide (Bu₄NOH). In pure methanol (Fig. 6a), the cyclic voltammogram was simple. It showed one Fe³⁺/Fe²⁺ redox at −0.05 V (vs. Ag|Ag⁺) and the porphyrin ring oxidation above 1.05 V (vs. Ag|Ag⁺). On the other hand, the cyclic voltammogram in the presence of hydroxide (Fig. 6b) was complicated. It showed Fe³⁺/Fe²⁺ redox at more negative potential, compared with that in pure methanol. This suggested that electron-donating species (hydroxide) ligated to the iron site. In addition to this negative shift of the redox potential, the cyclic voltammogram showed a new distorted redox at 0.9 V before the ring oxidation. This characteristic was very close to that observed at water-soluble Fe(TMPyP) [22]. Formation of oxo-iron(IV) (TPFPP) was estimated before the ring oxidation.

When NO was introduced into the solution of Fe(TPFPP) containing Bu₄NOH, the Fe³⁺/Fe²⁺ redox was disappeared and two oxidation peaks appeared as can be seen in Fig. 7. In analogy with Fe(TMPyP), the disappearance of the redox can be explained by the reductive nitrosylation, namely the formation of Fe(II)(NO) complex. The first oxidation peak corresponding to 1-electron oxidation at 0.55 V [23] can be ascribed to the oxidation of Fe(II)(NO) to Fe(III) + NO. The second large oxidation peak is due to oxidation of free NO. Agreement of the onset potentials of NO oxidation (Fig. 7a) and oxo-iron(IV) formation (Fig. 7b), and also an increase in the NO oxidation current by the presence of Fe(TPFPP) (Fig. 7a and Fig. 7c) suggest EC catalytic cycle as in the case of Fe(TMPyP) in aqueous solution [22].

![Figure 7](image_url)

**Figure 7.** Cyclic voltammograms of 4×10⁻⁴ M Fe(TPFPP)Cl at a GC electrode in 0.1 M TBAP methanol solution (a) saturated with 5% NO and (b) without NO. Curves (c) and (d) are obtained in 0.1 M TBAP methanol solution saturated with 5% NO and without NO, respectively. Scan rate, 100 mV s⁻¹.
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\[(\text{HO})\text{Fe(III)} - e^- \rightarrow \text{O=Fe(IV)} + \text{H}^+\]
\[\text{O=Fe(IV)} + \text{NO} + \text{OH}^- \rightarrow (\text{HO})\text{Fe(III)} + \text{NO}_2^-\]

The distribution of Fe(TPFPP) in Nafion film, which was doped by dissolving into Nafion solution, is not so simple. A 2/3 of water-insoluble H2TPP has been found to distribute in hydrophilic ion channels and the other 1/3 in hydrophobic region of Nafion film [24]. If Fe(TPFPP) was distributed in hydrophilic region, it is surrounded by water molecules and may form hydroxo-complex, (HO)Fe(III).

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