Multichannel semiconductor-based electrodes for in vivo electrochemical and electrophysiological studies in rat CNS

Craig G. van Horne¹, Spencer Bement³, Barry J. Hoffer¹,² and Greg A. Gerhardt¹,²

Departments of ¹Pharmacology and ²Psychiatry, University of Colorado Health Sciences Center, Denver, CO 80262 (U.S.A.) and ³Department of Electrical Engineering and Computer Science, University of Michigan, Ann Arbor, MI 48109 (U.S.A.)

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Five-channel silicon-based microprobes were sputter-coated with carbon, coated with Nafion, and used for both in vivo electrochemical and single-unit electrophysiological recordings. High-speed electrochemical studies were performed in vitro and in vivo, which demonstrated that these multi-site probes were capable of monitoring the evoked overflow of monoamines in selected brain regions of the rat. In addition, action potentials from Purkinje cells in the rat cerebellum, identified electrophysiologically, were recorded from different sites on the same probe. Spontaneous firing rates could be monitored for up to 2 hours in order to investigate the effects of systemic administration of phencyclidine. These results provide preliminary evidence that solid-state multi-site probes can be utilized for both in vivo electrochemical and electrophysiological studies in the rat brain.
Fig. 1. A: illustration of the various types of multichannel (5 site) recording electrodes used in this study. Type 1 electrodes had shank widths of 200 or 250 μm with recording sites of 4000 and 8000 μm², while type 2 probes had shank widths of 150 and 180 μm with recording sites of 1000 and 2000, respectively. B: high-speed in vivo electrochemical recordings of potassium-evoked overflow of monoamines in the medial prefrontal cortex of the rat. The ejections of potassium are indicated by the arrows.

cial investigations while only the 1000, 2000, and 4000 μm² surface area electrodes were employed for both electrochemical and electrophysiological investigations (see below).

Male Sprague-Dawley Rats (250–300 g) were anesthetized with urethane (1.25 g/kg), intubated, and placed in a stereotaxic frame for both in vivo electrochemical and electrophysiological studies. Electrochemical recordings were performed in the rat striatum and medial prefrontal cortex as determined by stereotaxic electrode placement [17]. All in vivo electrochemical measurements were performed using a high-speed chronoamperometric recording system (IVEC-5; Medical Systems Corp.). The multichannel probes were characterized for sensitivity to dopamine, norepinephrine and serotonin in vitro, and their selectivities to these compounds versus ascorbic acid were characterized. The sensitivities and linear recording characteristics of all recording electrodes were determined by measuring responses to stock solutions of DA; releases were expressed quantitatively in terms of the DA calibration curves. Square-wave pulses of 0.0 to +0.55 V, with respect to a Ag/AgCl reference electrode, were applied to the working electrode for 0.20–1.0 s. The resulting oxidation and reduction currents were digitally integrated during the final 60–80% of the recording period; one digital count recorded was approximately equal to $1.7 \times 10^{-13}$ C. Releases of monoamines were elicited by local application of K⁺ (120 mM KCl and 2.5 mM CaCl₂; 100–250 nl ejected) from micropipettes having outer tip diameters of 5–10 μm [8]. These were attached to the electrochemical recording electrodes using wax with a tip separation of 275–350 μm. In addition, extracellular single-unit recordings of cerebellar Purkinje neurons in the anesthetized rat were obtained with the multi-site Nafion-coated probes and with conventional single-barrel glass micropipettes as previously described [14, 16].

The recording sites of all 4 probe designs, following dip-coating with Nafion, responded linearly to increasing concentrations of dopamine; the linear regression correlation coefficients of the calibration curves were greater than 0.997 (n=8 probes). In addition, some probes were also tested for their sensitivity to norepinephrine and serotonin, and were seen to respond linearly to these compounds as well. In addition, good selectivities for the detection of dopamine versus ascorbic acid (AA) were achieved. Typical selectivities for DA versus AA ranged from 100 to 700:1 and averaged $261 \pm 106$ (S.E.M., n=5) for the 8000 μm² area probes. However, the charging currents of these probes were often higher which resulted in the necessity to use lower gain settings on the current to voltage amplifiers. As compared to carbon epoxy or carbon fiber electrodes we have previously studied [7–9], the signal-to-noise characteristics were also lower. For example, our graphite-epoxy electrodes, which have a surface area close to 8000 μm² (geometric area based on a disk surface 100 μm in diameter), record signals of $5385 \pm 259$ (S.E.M., n=22) digital counts per micromolar change in DA (one digital account is approximately equal to $1.7 \times 10^{-13}$ C). This is
Fig. 2. A: electrophysiological recordings with a multichannel semiconductor electrode (2000 \( \mu \text{m}^2 \) surface area per site; width of electrode shank was 180 \( \mu \text{m} \)) and a conventional glass micropipette (bar = 10 \( \mu \text{m} \)). Two different cerebellar Purkinje neurons obtained from the lower two sites were recorded from the probe. B: demonstration of the ability of a multichannel electrode (4000 \( \mu \text{m}^2 \) recording area per site) to record the sustained activity of a cerebellar Purkinje neuron and cumulative dose-response curves from the systemic administration of PCP. The two insets show portions of the actual ratemeter tracings before and 5 min after the systemic administration of 13 mg/kg PCP.

in contrast to the 821 ± 80 (n = 5) counts per micromolar change in DA seen for the 8000 \( \mu \text{m}^2 \) recording sites of the semiconductor probes. In addition, the reduction current responses recorded by these probes were consistently lower than the signals recorded using carbon-based recording electrodes [9]. Typical reduction/oxidation current responses for DA using the 8000 \( \mu \text{m}^2 \) area probes averaged 0.31 ± 0.04 (S.E.M., n = 5) at a 5 Hz recording rate, as compared to 0.58 ± 0.02 (n = 22) recorded with standard carbon epoxy electrodes [9].

Individual electrochemical recordings, using multichannel (8000 or 4000 \( \mu \text{m}^2 \) recording sites) semiconductor-based probes, of monoamine overflow produced by local application of potassium were recorded in striatum and the medial prefrontal cortex of 2 anesthetized rats (Fig. 1B). The ‘Ox’ curves (upper traces) represent the high-speed oxidation current recordings while the ‘Red’ tracings (lower traces) are the reduction current responses. Signal 1 was recorded 1.0 mm from cortical surface while signal 2 was recorded at a depth of 2.0 mm. In one animal, the signals recorded from the prefrontal cortex averaged 2.2 ± 0.2 \( \mu \text{M} \) (n = 4) in amplitude; values were expressed in DA equivalents using calibration factors determined for each probe in vitro. These signals and oxidation/reduction ratios are quite similar to those recorded in this same region with conventional graphite epoxy electrodes [9].

Nafion-coated probes were also used for electrophysiological recording before or after the electrochemical studies (n = 4 animals). Insertion of the probe into the cerebellum far anterior or posterior in the folium, where Purkinje cells are aligned vertically, often yielded recordings from such neurons at two or more sites on the same probe (Fig. 2A); sites were selected using an electrode switching circuit connected to a single-channel recording amplifier. Action potentials from different neurons at two recording sites were seen simultaneously on a number of occasions (2 animals). Purkinje cells were identified by their firing rates (range = 6–30 Hz, n = 12 cells) in conjunction with their characteristic firing patterns composed of both simple and complex spikes [14]. The firing rates and waveform shapes recorded from the multichannel probes were similar to those recorded by a conventional glass micropipette in the same folium (Fig. 2A).

The ability of the probe to record sustained electrophysiological activity, and its alteration by pharmacological manipulation of presynaptic input, was investigated in 2 animals and such data are shown in Fig. 2B. In this experiment, activity from a cerebellar Purkinje neuron was continuously recorded for over two hours. During this time, phencyclidine (PCP) was administered parenterally in increasing doses, which caused dose-dependent decreases in the basal Purkinje cell firing rate; individual action potentials were periodically viewed to ensure that the effects of PCP were not due to local anesthetic effects or movement of the recording electrodes. In the cerebellum, PCP acts as an indirect noradrenergic agonist, probably through inhibition of the uptake carrier for norepinephrine [14].

This study provides the first documentation that Nafion-coated semiconductor-based multichannel
recording electrodes can be used to quantitatively measure monoamines both in vitro and in vivo using high-speed chronoamperometric recording techniques, and that the same multichannel probes can be used for single-unit electrophysiological recordings. Moreover, electrophysiology experiments that require long recording times, such as those analyzing drug dose–effect relationships, can be investigated with these probes.

Dip-coating of the multichannel probes with Nafion was seen to increase the selectivity of the recording sites for cationic neurotransmitters versus anionic interferents such as ascorbic acid. The selectivities of the individual probes for DA were similar to those seen using Nafion-coated graphite epoxy capillary or carbon fiber electrodes [7–9]. In addition, the Nafion coating did not appear to negatively influence the multichannel recording probes for single-unit electrophysiological measurements, as the recording characteristics of these sensors were seen to rival those seen with more standard glass micropipette electrodes. Interestingly, preliminary attempts to use the multichannel electrodes without the Nafion coating were relatively unsuccessful. Single Purkinje cells were not easily identified, and the signal-to-noise properties of the electrodes appeared to be significantly compromised. Thus, the Nafion coating may have a positive effect on the electrophysiological recording properties of the multichannel probes. In addition, the potential utilization of Nafion coatings to improve the electrophysiological recording properties of other metal recording electrodes should be investigated.

There are several additional technological advances that must be made before such probes will be truly useful to characterize the emergent properties of neural ensembles in behaving animals. First, the recording surfaces need to be improved for combined electrochemical and electrophysiological recordings. The type and thickness of carbon layer on the surface must be optimized, as well as the procedures for coating the probes with Nafion. Secondly, microdrives and wire-bonding techniques must be developed to allow implantation of such probes into awake behaving animals. Third, computer interfacing and hardware must be fabricated to allow simultaneous acquisition and storage of multiple single-unit electrophysiological and multichannel electrochemical signals. The cost of such hardware must be within the reach of individual laboratories. Finally, analytical schemes must be formulated and translated into real-time software, in order to quantitatively analyze such large volumes of data and to test hypotheses of neural network function. These complexities notwithstanding, the data in this paper suggest that semiconductor microprobes may play increasingly important roles as real-time biological sensors to analyze simultaneous pre-synaptic neurotransmitter dynamics and activity of multiple neurons.

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