

An Analog FSCV Device Combined with a Phase-Locked Loop for in Vivo Detection of Dopamine

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Abstract—Objective: In this work, our goal was to develop an easy-to-implement fast-scan cyclic voltammetry (FSCV) device which can be reliably employed for in vivo detection of dopamine. **Methods:** Readily available analog integrated circuits (ICs) were used to generate monopolar triangular voltammetric waveforms with adjustable holding and peak potentials. The timing of the voltammetric scans were controlled by a phase-locked loop implemented around a standard microcontroller so that the power line frequency noise, which generally complicates the detection of dopamine through the recordings, could be effortlessly eliminated by background-subtraction. **Results:** The practicality of the system was tested in vitro using a dopamine solution and in vivo using a freely moving rat chronically implanted with a carbon fiber microelectrode in the ventromedial striatum (VMS). The system consistently detected dopamine in the solution and dopamine released in the VMS in response to reward presentation. **Conclusion:** We demonstrated a low-cost and easily reproducible solution specifically designed for in vivo measurements of dopamine from behaving subjects. **Significance:** Based on utilization of analog ICs, the triangular waveform generated in the present system is linear in contrast to the conventional, digital-to-analog converter-based systems, which fundamentally approximates a staircase form to a linear ramp. We believe the present technique can be applied for the measurement of other electrochemically active analytes requiring a triangular waveform and elimination of power line frequency noise.

Index Terms—Dopamine, electrochemistry, fast-scan cyclic voltammetry, neuromodulator, neuroscience, phase-locked loop.

I. INTRODUCTION

RESEARCH into motor control, goal-directed behavior and reinforcement learning is significantly focused on the

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characterization of phasic changes in the activity of the midbrain dopaminergic neurons and the modulations of dopamine concentration in the structures innervated by dopaminergic axons [1]–[3]. Electrophysiology enables recording the activity of dopaminergic neurons with high temporal and spatial resolution [4], [5]. In addition to electrophysiology, fast-scan cyclic voltammetry (FSCV), as another powerful technique, offers the capability of monitoring the phasic dopamine release in the striatal structures with high temporal resolution and spatial acuity [6]–[9].

FSCV, as an electroanalytical technique, is used to measure the concentration of electrochemically active molecules in the vicinity of an electrode [10]. It applies electrical potential to the electrode and induces oxidation and reduction of the analyte at the surface of the electrode. These reactions lead to faradaic current flow between the electrode and the analyte. This current is directly proportional to the concentration of the analyte in the medium [11]. In the case of in vivo detection of dopamine via FSCV, carbon fiber microelectrodes are conventionally used due to their small size, biocompatibility and inertness during application of voltage scans [12]–[14]. It is implanted into the target tissue and a triangular voltammetric waveform is applied to the microelectrode versus a Ag/AgCl reference electrode [15]. Typically, the applied potential is held at -0.4 V versus the reference electrode. During the scans it is linearly ramped to a maximum voltage of 1.3 V at a scan rate of 400 V/s and returned to the holding potential (-0.4 V) with the same scan rate [16]–[19]. The voltammetric scan is, therefore, completed within 8.5 ms. The faradaic currents occurring during the scans are then used to detect the concentration of dopamine. In order to extract the faradaic currents through the scans, cyclic voltammograms (CVs), which present the overall current passing through electrode versus applied voltage, are collected for a baseline period and these CVs are then subtracted from the ones acquired in following scans. The remaining current in the subsequent scans after subtraction corresponds to faradaic current, in other words background-subtracted current, which represents the change in the concentration of dopamine in the vicinity of the implanted microelectrode [20], [21]. Therefore, FSCV allows monitoring the phasic changes in dopamine concentration relative to a baseline reference point [22]–[24].

The conventional approach for detection of dopamine (and

other electrochemically active analytes) via FSCV has been based on employing an operational amplifier (opamp) in I/E (current-to-voltage) converter mode: The working electrode is directly connected to the inverting input of the operational amplifier and the voltammetric waveform is applied to its non-inverting input. In this configuration, the voltage of the working electrode follows the waveform voltage and the resulting current passing through the electrode is converted to voltage at the output of the operational amplifier [25]. The voltage output of the operational amplifier is recorded using a PC or another digital system and the current passing through the electrode is extracted from the recordings in a post hoc fashion to create CVs.

Since the faradaic currents generated by redox reactions of physiological concentrations of dopamine are in the orders of nanoamperes [26]–[28], the development of a reliable and low-noise FSCV system which supports *in vivo* detection of dopamine is a non-trivial problem. There are commercially available professional devices such as WaveNeuro Potentiostat System and WaveNow Wireless Potentiostat System (Pine Research Instrumentation, NC, USA) and research devices in the literature which allows dopamine measurement using FSCV technique [29]–[31]. In these FSCV devices, the voltammetric waveform is generated using a digital-to-analog converter (DAC). Instead of using an integrated FSCV device, the other solution for detection of dopamine is to create a FSCV setup where the triangular waveform is produced by a data acquisition card controlled by a PC and applied to the working electrode through a I/E converter [32], [33]. In such setting, data acquisition card also employs a DAC fundamentally for the generation of the triangular waveform. In order to create a waveform with a high scan rate (e.g. 400 V/s) and close to a true triangle, use of a DAC with a high resolution and a high update rate is necessary. In addition, powerful microprocessors and advanced embedded system programming skills are required to control these integrated circuits (ICs). These requirements significantly increases the development costs of the equipments used in FSCV.

In the present article, we propose a novel FSCV device which generates triangular voltammetric waveforms without using DAC and is specifically designed for facilitating *in vivo* detection of dopamine. In lieu of employing a DAC, the present device implements a customized analog circuit which converts pulses generated by a standard microcontroller into triangular waves. Triangular waves are followed by a DC holding potential and repeated every 100 ms for online monitoring of dopamine transients as in conventional DAC-based systems. The present design approach decreases the development and implementation costs of FSCV systems which use triangular scans, and allows easy implementation in a standard neuroscience research laboratory by utilizing readily available electronic components. The present design based on analog circuit components also enables generation of true triangular waves with linear, continuous rising and falling ramps in contrast to the existing systems, which basically generate a staircase waveform due to use of DAC. In addition,

the holding and peak voltages, and the scan rate for the anodic and cathodic sweep can be easily adjusted using the present device. Here we describe the structure of the proposed FSCV device and validate the practicality of the device by *in vivo* measurements acquired from a freely moving rat chronically implanted with a carbon fiber microelectrode. Last but not least, we demonstrate how to interface the device with a data acquisition card and a commutator for continuous recordings from behaving subjects.

II. METHODS

A. Structure of the FSCV Device

The FSCV device presented here was designed to enable measurement of dopamine from freely moving subjects. **Fig. 1** illustrates the FSCV hardware and its interfacing with an experimental setup, and **Fig. 2** demonstrates the circuit diagram of the hardware. The FSCV hardware is composed of four modules: 1) waveform generator 2) phase-locked loop (PLL), 3) headstage and 4) amplifier. In this section, we describe the design and implementation of these modules.

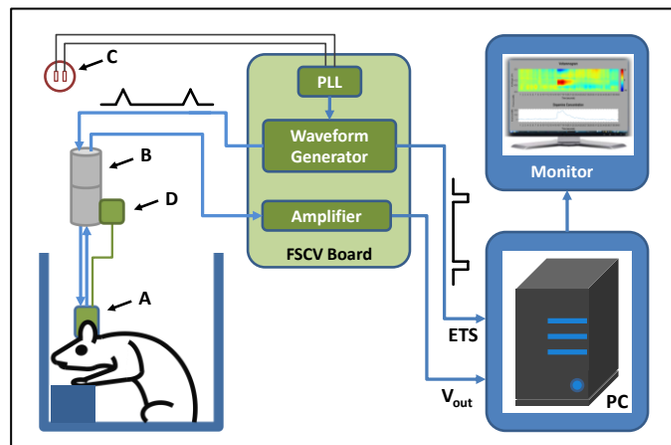


Fig. 1. Block diagram of the FSCV hardware and the experimental setup. The headstage (A) implementing a I/E converter is fixed on rat's head and receives triangular signals from the waveform generator through a commutator (B) used to keep the headstage cables untangled during recordings. The output of the headstage is fed to the amplifier through the commutator and the voltammetric signals (V_{out}) are recorded using a PC equipped with a data acquisition card. A phase-locked loop (PLL) is used to match the phase of the generated triangular waveform to the phase of mains (C). The pulses indicating the starting times of the scans are used as the external trigger signal (ETS) to initiate the data acquisition every 100 ms. Batteries (D) mounted onto the commutator directly supplies power for the headstage.

1) Waveform generator

The waveform generator module of the FSCV hardware is used to produce the triangular voltammetric waveform which is applied to a working microelectrode versus a reference electrode for detection of dopamine. **Fig. 2B** shows the circuit diagram of the waveform generator. The waveform generator basically consists of analog circuit elements and converts pulses with a width of 4.25 ms into triangular waves with a width of 8.5 ms using an integrator circuit. The first stage of the waveform generator is a summing amplifier and is used to offset the pulse signal fed to the second, integration stage. The

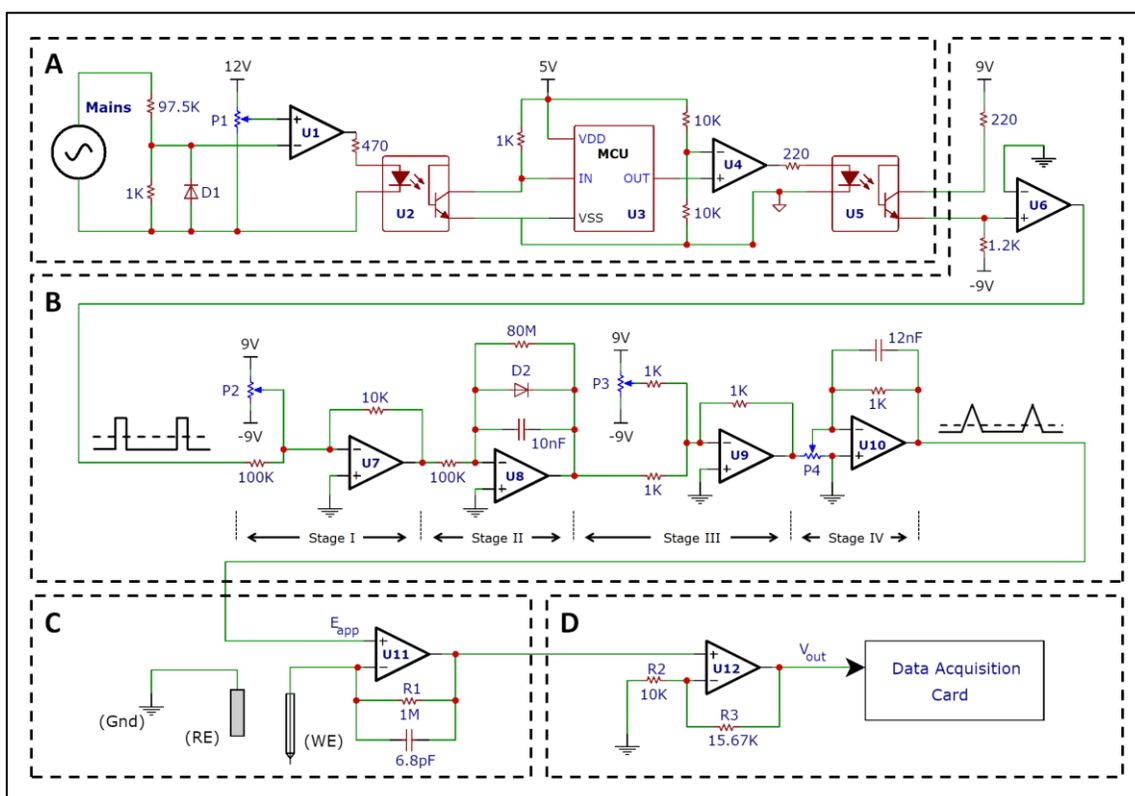


Fig. 2. Circuit diagram for the FSCV hardware. (A) Phase-locked loop (PLL) module. MCU: microcontroller unit. (B) Waveform generator module. (C) Headstage module. RE: reference electrode, WE: working electrode, Gnd: ground. (D) Amplifier module.

offsetting process at this stage using potentiometer P2 is performed to create symmetry between the maximum and minimum voltages of the pulse signal at the input of the integration stage. This symmetry in the pulse signal is required for obtaining symmetry between the rising and falling slopes of the triangular waveform at the output of the integration stage. The offset and amplitude of triangular waveform generated by the integration stage is then adjusted by potentiometers P3 and P4, respectively, in the third and fourth stages. Diode D2 (1N4005) is utilized within the integration stage to prevent the operational amplifier U8 from saturation and create a rectified, monopolar triangular waveform. The operational amplifiers used for U7 and U8 are TL07x series (Texas Instruments, TX, USA) because of their high slew rate (13 V/ μ s) and low input bias current (65 pA), which are required for offsetting the pulse signal without distorting its shape at the first stage and limiting the DC gain of U8 at the integration stage with a large-value feedback resistor (i.e. 80 M Ω), respectively [34]. The operational amplifiers used for U8 and U9 are also TL07x series due to their easy availability and low-cost. The voltage comparator used for U6 is LM311 (Texas Instruments) because of its fast response time (i.e. 165 ns) and wide supply voltage range.

2) Phase-locked loop (PLL)

Capacitive coupling between the power line (mains) and the tissue is a significant source of noise, which hampers the detection of the small faradaic currents resulting from the redox reactions of physiological levels of dopamine. This type of noise can be eliminated by subtracting the background

current if the phases of the power line signal and the applied voltammetric waveform precisely match each other [35]. For monitoring the power line frequency and controlling the timing of the voltammetric waveform, we implemented the PLL module for the FSCV device presented here (see Fig. 2A). In the PLL module, the power line signal is divided by approximately 100 using 1 W resistors, half-wave rectified by a diode (D1, 1N4007) and converted into pulses using a comparator (U1). A microcontroller (U3) monitors these pulses through an optocoupler (U2) and outputs phase-locked pulses with a width of 4.25 ms and a frequency of 10 Hz. The width of the phase-locked pulses is precisely controlled using the timer of the microcontroller. The duty cycle of the pulses at the input of the microcontroller is 65%, set by potentiometer P1 in the PLL module. Mains is isolated from the microcontroller using optocoupler U2 for electrical safety. Another optocoupler (U5) is used between the PLL and waveform generator modules for isolating their power supplies from each other. The optocouplers used for U2 and U5 is 4N35 (Vishay Semiconductors, PA, USA) and the microcontroller used for U3 is PIC18F4520 (Microchip Technology, AZ, USA).

3) Headstage and amplifier

The headstage (see Fig. 1) is used to apply the triangular waveform to the working microelectrode versus the Ag/AgCl reference electrode. Fig. 2C illustrates the circuit diagram of the headstage, implementing a I/E converter using an operational amplifier [25]. TLC227x series (Texas Instruments) operational amplifier is used for U11 due to their

low input bias current (1 pA), low input current noise (0.6 fA/ $\sqrt{\text{Hz}}$) and availability in small size with quad-channel versions (e.g. TLC2274IPW), which can be advantageous for future applications requiring a higher number of recording microelectrodes. The power for headstage is directly supplied from separate batteries (see **Fig. 1**) instead of providing it through the commutator so that any significant noise in the supply voltage of the headstage that can be caused by the rotation of the commutator is avoided. Lithium-ion batteries (3.7 V) are used to supply voltage to the headstage due to their light-weight. In the headstage, an operational amplifier with a high input impedance is used (i.e. TLC227x series) to prevent any large distortion in the voltammetric waveform received from the FSCV board through the commutator (see **Fig. 1** and **Fig. 2**). Three channels of a 32-channel commutator (Plexon Inc., TX, USA) are used in the recording. One channel is used for the connection between the ground and the reference electrode, another one is used for transmitting voltammetric waveform to the headstage and the other is used for transmitting the output of the headstage to the amplifier module (see **Fig. 2C**).

The amplifier module (**Fig. 2D**) is used to adjust the maximum amplitude of the signal at the input of the data acquisition card in the PC so that its full voltage range can be used while avoiding saturation. Typically, we set the gain of the amplifier to 2.567 for chronically implanted carbon fiber microelectrodes as shown in **Fig. 2D**. Using the present configuration, the current from the working microelectrode (i) could be calculated by measuring the voltage at the output of the FSCV (V_{out}) device as follows:

$$i = \frac{V_{out}R_2}{R_1(R_2 + R_3)} - \frac{E_{app}}{R_1} \quad (1)$$

where E_{app} is the voltage applied to the working electrode or the non-inverting input of the opamp in the headstage (U11), R_1 is the feedback resistor in the headstage, R_2 and R_3 are the resistors to adjust the gain of the amplifier module.

4) Data acquisition

The output of the voltammetry hardware (V_{out}) is recorded using a PC equipped with a data acquisition (DAQ) card (**Fig. 1**). The DAQ card (National Instruments, PCI-6040E, TX, USA), which supports 12-bit resolution, is configured to perform analog-to-digital conversion (ADC) through one channel with a sampling rate of 200 kHz. LabVIEW software (National Instruments) is used to control the DAQ device. In order to control the initiation time of the recordings and match them to the voltammetric scans, the phase-locked pulses used for generation of the triangular waveform are also employed for externally triggering the DAQ card (see **Fig. 1** and **Fig. S1**). The DAQ device is configured to acquire 3000 samples for each triggering pulse. 1700 samples, which correspond to the voltammetric scan time (8.5 ms), are used to construct CVs.

B. Electrode Fabrication and Implantation

A carbon fiber microelectrode as the working electrode and a Ag/AgCl electrode as the reference electrode were

implanted into the rat brain for testing the practicality of the FSCV device presented here. In this section, we describe the methods we followed for the fabrication of these electrodes and their implantation into the rat brain.

1) Microelectrode and Reference Electrode Fabrication

Chronically implantable microelectrodes and reference electrodes were fabricated as previously described [6], [36]. A single carbon fiber with a diameter of 7 μm (Goodfellow Cambridge Ltd. #C005722, UK) was aspirated into a polyimide coated fused silica capillary (Polymicro Technologies #TSP020090, AZ, USA) with an outer diameter of 90 μm . One end of the capillary was sealed using a two-component epoxy (Epoxy Technology Inc. EPO-TEK 301, MA, USA). The epoxy was applied after becoming viscous so that a smooth, bulb-shaped microelectrode tip could be formed to reduce the tissue damage during implantation. The other end of the capillary was fixed onto a copper wire with a diameter of 1 mm using epoxy and silver paint (Loctite Co. #3863, Germany) was applied to provide the electrical connection between the carbon fiber and the wire. The silver paint was coated using a thin layer of dental acrylic subsequent to drying to further secure the electrical connection between the wire and the carbon fiber. Finally, the carbon fiber protruding from the other end of the capillary was cut to a length of 150-200 μm to prepare the recording tip of the microelectrode.

Reference (Ag/AgCl) electrodes were fabricated via electrolysis. A silver wire with a diameter of 0.2 mm was used to make the reference electrode. The positive terminal of a 5V-DC power supply with 500 mA current limitation was connected to the silver wire and the negative terminal to a stainless steel wire. Both wires were immersed into a hydrochloric acid solution (36%) and the current was applied for approximately 2-3 minutes until the bubbles disappeared, indicating that the electrolysis process was complete.

2) Voltammetry Surgery

All animal procedures presented in this paper were approved by and conducted in accordance with the regulations of the Istanbul Medipol University Ethics Committee on Animal Maintenance and Experimentation (Protocol Number: 38828770-604.01.01-E.1815, Date of Approval: August 31st, 2015). One Wistar male rat weighing 420 g was chronically implanted with a carbon fiber microelectrode in the ventromedial striatum (VMS) in the left hemisphere (anteroposterior +1.5 mm, mediolateral -2.1 mm, dorsoventral 6.4 mm) and a Ag/AgCl reference electrode in the contralateral hemisphere (anteroposterior +1.5 mm, mediolateral +2.1 mm, dorsoventral 3.0 mm) [37]. Implantation of the electrodes was performed under anesthesia achieved with an i.p. injection (1.5 cc/kg) of a mixture of ketamine and xylazine (100 mg/kg and 12 mg/kg in saline, respectively). Maintenance of anesthesia was achieved with isoflurane gas as needed. Craniotomy was created bilaterally and the dura mater was removed over the implantation sites carefully while avoiding damaging blood vessels and the pia

mater [38]. The carbon fiber microelectrode was lowered into the target position slowly (100 $\mu\text{m}/\text{min}$) after reaching a depth of 6 mm. The positions of the working and reference electrodes were fixed using dental acrylic. A female SubMiniature version A (SMA) connector was fixed onto the dental acrylic on the head of the rat. The copper wire of the carbon fiber microelectrode was soldered to the inner pin and the reference wire to the outer pin of the SMA connector. SMA connectors were preferred for the connection between the headstage and the implanted electrodes due their screw-type coupling mechanism providing the reliability in the electrical connection (see **Fig. S2**). The rat was given 1 month to recover from the surgery before initiating the voltammetric recording [13], [32].

III. RESULTS

The practicality of the FSCV device presented here was tested by performing measurements of dopamine concentration both in vitro and in vivo. For the in vitro measurements, we recorded the background current from a saline solution and performed another recording from the same solution after addition of dopamine. The dopamine concentration of the latter solution was 1 μM . **Fig. 3A** demonstrates the color plot of the background-subtracted voltammetric signal acquired from this solution. The corresponding background-subtracted cyclic voltammogram for a single scan is shown in **Fig. 3D**.

Secondly, the FSCV device was examined for in vivo detection of dopamine using a rat chronically implanted with a carbon fiber microelectrode in the ventromedial striatum (see **Voltammetry Surgery**). Prior to the initiation of the recording, the rat was placed into the experimental setup represented in **Fig. 1** and tethered to the FSCV device for 1 hour for stabilization of the background current. **Fig. 3B** displays the background-subtracted voltammetric signal acquired from the rat in response to presentation of reward (i.e. 20 μl of 30% sucrose solution) using a pipette. The changes in redox currents as a function of the applied voltage (E_{app}) are illustrated in the plot by color codes. The reward presentation time is indicated by black triangular marker at the top of the plot. The background-subtracted cyclic voltammogram corresponding to the single scan pointed by the gray triangular marker is presented in **Fig. 3E**. The output of the FSCV device (V_{out} in **Fig. 2**) for a single scan during the present recording is demonstrated in **Fig. 3F**.

Finally, we tested the noise in the whole system. In order to evaluate the noise in the system, we connected a resistor between the inverting input of the operation amplifier in the headstage (U11) and ground (see **Fig. 2**), and performed recording using the data acquisition system presented here (see **Data acquisition**). The resistor here was employed to simulate the current passing through the working microelectrode. The value of the resistor was 1.2 $\text{M}\Omega$ to obtain a peak background current (i.e. 1.026 μA) comparable to the one recorded in vivo (i.e. 1.066 μA , see **Fig. 3F** and **Eq. (1)**). **Fig. 3C** demonstrates the background-subtracted current pass-

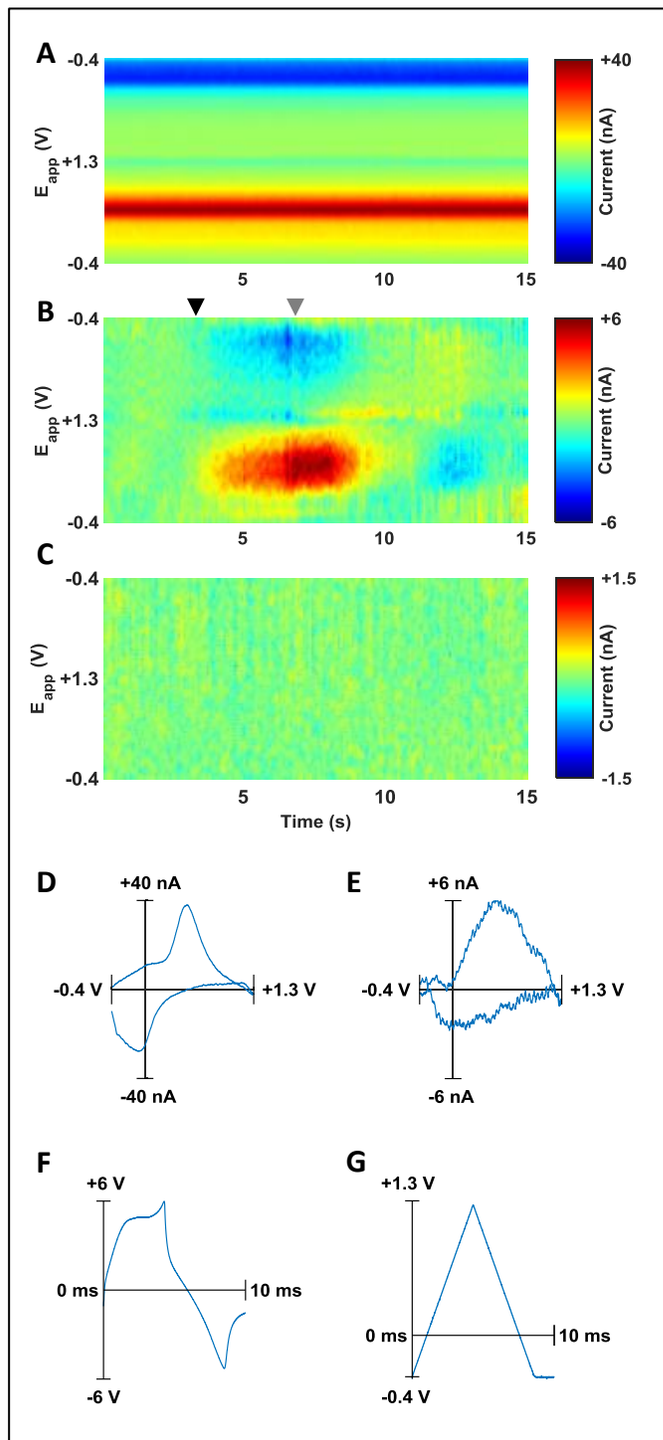


Fig. 3. Dopamine detection using the present FSCV device. (A-C) Background-subtracted voltammetric signal acquired from 1 μM dopamine solution (A), the ventromedial striatum of an awake rat in response to reward presentation (B), and 1.2 $\text{M}\Omega$ resistor simulating the working microelectrode for evaluating the noise in the FSCV device (C). (D and E) Background-subtracted CVs corresponding to the plots shown in (A) and (B), respectively. (F) The signal at the output of the FSCV device (V_{out}) for a single scan in (B). (G) The triangular waveform generated by the present FSCV device. All plots except for (G) were filtered by centered moving averaging with a window size of 61 to suppress high frequency noise without phase shifting.

ing through the resistor as a function of the applied voltage. The background CV was calculated by averaging all scans included in the present plot. The overall standard deviation of

the noise across this recording was 2.913 nA without any filtering and 0.138 nA with centered moving average filtering with a window size of 61, which was also used in the plot in **Fig. 3B**. The overall standard deviation of the noise in **Fig. 3B** within the first three seconds, where the background-subtracted CVs were stabilized, was 2.836 nA without any filtering and 0.313 nA with centered moving average filtering with a window size of 61. **Fig. 3G** demonstrates the triangular waveform generated by the present FSCV device. The present triangular waveform was recorded by the data acquisition system described here (see **Data acquisition**).

IV. DISCUSSION

In the present work, our goal was to develop a practical, low-cost and easy-to-implement FSCV device for in vivo measurement of dopamine concentration. The solution proposed here was implemented using readily available and low-cost electronic components, and allowed measuring the changes in dopamine concentration in vivo (see **Fig. 3**). The faradaic currents caused by the oxidation and reduction of dopamine were clearly observable through the background-subtracted CVs. The present device generated smooth triangular voltammetric waveforms whose timings were effectively controlled by the phase of power line (mains) signal via the PLL circuit described here. Thanks to the present PLL circuit, we could effectively eliminate the interference caused by the mains (see **Fig. 3A-C**) and obtained a low-noise system. The dopamine signal could be detected in vivo without using a faraday cage or electromagnetic shielding.

The triangular waveform generator module smoothly converted pulses into monopolar triangle waves each followed by a DC holding potential. The diode in the integrator circuit effectively enabled generation of a monopolarity in the waveform. Overall, the present design approach based on use of readily available analog circuit components allowed the generation of a linear ramping signal without any discontinuity, which is unavoidably introduced by DACs, generally used for producing the voltammetric waveform in FSCV systems. In conventional systems, a DAC converter with a high resolution and update rate is preferred to improve the similarity of the generated voltammetric waveform to a triangular waveform. However, DACs with such specifications can be costly and require powerful microprocessors for control. In this work, we removed these requirements with a low-cost and practical solution. The microcontroller simply produced pulses with a specified width using timer interrupts and the waveform generator module, implemented using readily available analog ICs, converted these pulses into triangular waveform with adjustable holding and peak potentials.

The potentiometers involved in the waveform generator module enabled setting the scan rate and waveform amplitude in a user-friendly and easy manner. It is known that there can be a 200 mV shift in the reference potential at the Ag/AgCl electrodes after implantation and during longitudinal recordings [39], [40]. This shift can be identified by observing

the faradaic peaks in the background currents and background-subtracted currents, and it can be compensated by offsetting the applied waveform by +200mV. We did not observe such shift in our in vivo test, however we believe the generated waveform can be easily offset using P3 potentiometer (**Fig. 2B**) if such compensation is required. When a change in the width of the rising ramp of triangular waveform is needed for different applications, the width of the pulse generated by the microcontroller can be adjusted by updating its program. The width of the falling ramp can be simply controlled by potentiometer P2 (**Fig. 2B**) without updating the program of the microcontroller. Testing the applicability of the present waveform generator for detection of other electrochemically active molecules was out of the scope of this work. We believe the present system can also allow detection of the ones requiring a triangular waveform since it supports setting the holding and peak potentials [41], [42].

We optically isolated the PLL module from the waveform generator module in order to allow replacement of the microcontroller used here with other ones and prevent any noise interference which might be caused by the microcontroller operation and its clock circuitry. This design also allows replacement of the present PLL module with other simple pulse generators (e.g. function generators) when any phase-locking and elimination of mains noise is not necessary. Using such readily available signal generators, one can create the pulses with required width and frequency without using a microcontroller. Since there is optical isolation at the input of the triangular waveform generator module, the elimination of noise interference from these devices is guaranteed.

In the present system, we utilized a commutator to allow the rat freely move in the cage while it was tethered to the FSCV board. We connected the headstage batteries directly to the headstage instead of providing the supply voltage from the FSCV board through the commutator (see **Fig. 1**). It has been an effective solution to avoid the voltage drifts in the headstage which could appear if the headstage power were supplied through the commutator. The present system ensured stabilization of the background current within 1 hour after connecting the headstage to the implanted electrodes. Our results based on the in vivo recording (see **Fig. 3B**) demonstrated that the stability in background current could be effectively achieved using this configuration.

In the present system, we used a data acquisition card to record the output of the FSCV hardware with a resolution of 12-bit and a sampling rate of 200 KHz. Based on these sampling rate and resolution, we empirically found that a centered moving average filter with a windows size of 61 allowed us to visualize dopamine signal practically. We preferred using a centered moving average filter instead of a digital low-pass filter to avoid any phase shift-related distortion in the visualized background-subtracted CV. We believe it is also possible to use a lower sampling rate (e.g. 100 KHz) in data acquisition while reliably detecting the faradaic currents from dopamine. The digital filter used should be updated according to the specifications of the data acquisition equipment. An analog low-pass filter can also be

implemented into the amplifier module of the present device instead of performance of digital filtering.

V. CONCLUSION

We described a low-cost and easy-to-implement analog triangular waveform generator for FSCV and validated its practicality for in vivo and in vitro detection of dopamine. We demonstrated its operation by developing a low-cost FSCV device, and interfacing the device with chronically implanted electrodes and a data acquisition system. The present device based on use of readily available analog circuit elements facilitates the implementation of a FSCV system in a standard neuroscience research laboratory. After the present system is established in a laboratory setting, the user can easily change the holding and the peak potentials of the voltammetric waveform as needed using the potentiometers included in the system. The present system generates a triangular voltammetric waveform with linear ramps in contrast to conventional systems generating the voltammetric waveform using a DAC, which technically has a limited resolution and update rate. We believe the present approach for voltammetric waveform generation can also be used in the applications targeting detection of other electrochemically active molecules requiring a triangular voltage scan.

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REFERENCES

- [1] M. Howe, P. L. Tierney, S. G. Sandberg, P. E. M. Phillips, and A. M. Graybiel, "Prolonged dopamine signalling in striatum signals proximity and value of distant rewards.," *Nature*, Aug. 2013, doi: 10.1038/nature12475.
- [2] X. Jin and R. M. Costa, "Start/stop signals emerge in nigrostriatal circuits during sequence learning.," *Nature*, vol. 466, no. 7305, pp. 457–462, 2010, doi: 10.1038/nature09263.
- [3] J. N. Reynolds, B. I. Hyland, and J. R. Wickens, "A cellular mechanism of reward-related learning.," *Nature*, vol. 413, no. 6851, pp. 67–70, Sep. 2001, doi: 10.1038/35092560.
- [4] W. Schultz, P. Dayan, and P. R. Montague, "A neural substrate of prediction and reward.," *Science*, vol. 275, no. 5306, pp. 1593–1599, Mar. 1997, doi: 10.1126/science.275.5306.1593.
- [5] P. N. Tobler, C. D. Fiorillo, and W. Schultz, "Adaptive coding of reward value by dopamine neurons.," *Science*, vol. 307, no. 5715, pp. 1642–5, Mar. 2005, doi: 10.1126/science.1105370.
- [6] P. E. M. Phillips, D. L. Robinson, G. D. Stuber, R. M. Carelli, and R. M. Wightman, "Real-Time Measurements of Phasic Changes in Extracellular Dopamine Concentration in Freely Moving Rats by Fast-Scan Cyclic Voltammetry.," in *Drugs of Abuse. Methods In Molecular Medicine*, vol. 79, J. Q. Wang, Ed. New Jersey: Humana Press, 2003, pp. 443–464.
- [7] D. L. Robinson, B. J. Venton, M. L. A. V Heien, and R. M. Wightman, "Detecting subsecond dopamine release with fast-scan cyclic voltammetry in vivo.," *Clin. Chem.*, vol. 49, no. 10, pp. 1763–1773, 2003, doi: 10.1373/49.10.1763.
- [8] I. Willuhn *et al.*, "Phasic Dopamine Release in the Nucleus Accumbens in Response to Pro-Social 50 kHz Ultrasonic Vocalizations in Rats.," *J. Neurosci.*, vol. 34, no. 32, pp. 10616–10623, 2014, doi: 10.1523/JNEUROSCI.1060-14.2014.
- [9] M. Klanker, L. Fellingner, M. Feenstra, I. Willuhn, and D. Denys, "Regionally distinct phasic dopamine release patterns in the striatum during reversal learning.," *Neuroscience*, no. May, 2016, doi: 10.1016/j.neuroscience.2016.05.011.
- [10] M. M. Arnold, L. M. Burgeno, and P. E. M. Phillips, "Fast-Scan Cyclic Voltammetry in Behaving Animals.," in *Basic Electrophysiological Methods*, E. Covey and M. Carter, Eds. Oxford University Press, 2015.
- [11] B. J. Venton and Q. Cao, "Fundamentals of fast-scan cyclic voltammetry for dopamine detection.," *Analyst*, vol. 145, no. 4, pp. 1158–1168, 2020, doi: 10.1039/c9an01586h.
- [12] R. M. Wightman, "Microvoltammetric Electrodes.," *Anal. Chem.*, vol. 53, no. 9, pp. 1125–1134, Aug. 1981, doi: 10.1021/ac00232a791.
- [13] N. T. Rodeberg, S. G. Sandberg, J. A. Johnson, P. E. M. Phillips, and R. M. Wightman, "Hitchhiker's Guide to Voltammetry: Acute and Chronic Electrodes for in Vivo Fast-Scan Cyclic Voltammetry.," *ACS Chem. Neurosci.*, vol. 8, no. 2, pp. 221–234, Feb. 2017, doi: 10.1021/acscchemneuro.6b00393.
- [14] R. B. Keithley *et al.*, "Higher sensitivity dopamine measurements with faster-scan cyclic voltammetry.," *Anal. Chem.*, vol. 83, no. 9, pp. 3563–3571, 2011, doi: 10.1021/ac200143v.
- [15] R. M. Wightman, L. J. May, and A. C. Michael, "Detection of Dopamine Dynamics in the Brain.," *Anal. Chem.*, vol. 60, no. 13, pp. 769–793, Jul. 1988, doi: 10.1021/ac00164a718.
- [16] M. F. Roitman, R. A. Wheeler, R. M. Wightman, and R. M. Carelli, "Real-time chemical responses in the nucleus accumbens differentiate rewarding and aversive stimuli.," *Nat. Neurosci.*, vol. 11, no. 12, pp. 1376–1377, 2008, doi: 10.1038/nn.2219.
- [17] J. J. Day, M. F. Roitman, R. M. Wightman, and R. M. Carelli, "Associative learning mediates dynamic shifts in dopamine signaling in the nucleus accumbens.," *Nat. Neurosci.*, vol. 10, no. 8, pp. 1020–1028, 2007, doi: 10.1038/nn1923.
- [18] M. L. a V Heien *et al.*, "Real-time measurement of dopamine fluctuations after cocaine in the brain of behaving rats.," *Proc. Natl. Acad. Sci. U. S. A.*, vol. 102, no. 29, pp. 10023–10028, 2005, doi: 10.1073/pnas.0504657102.
- [19] P. Takmakov *et al.*, "Carbon microelectrodes with a renewable surface.," *Anal. Chem.*, vol. 82, no. 5, pp. 2020–2028, 2010, doi: 10.1021/ac902753x.
- [20] J. O. Howell, W. G. Kuhr, R. E. Enzman, and R. Mark Wightman, "Background subtraction for rapid scan voltammetry.," *J. Electroanal. Chem. Interfacial Electrochem.*, vol. 209, no. 1, pp. 77–90, Sep. 1986, doi: 10.1016/0022-0728(86)80187-5.
- [21] M. DeWaele *et al.*, "A baseline drift detrending technique for fast scan cyclic voltammetry.," *Analyst*, vol. 142, no. 22, pp. 4317–4321, 2017, doi: 10.1039/C7AN01465A.
- [22] E. W. Schluter, A. R. Mitz, J. F. Cheer, and B. B. Averbeck, "Real-time dopamine measurement in awake monkeys.," *PLoS One*, vol. 9, no. 6, 2014, doi: 10.1371/journal.pone.0098692.
- [23] J. A. Johnson, N. T. Rodeberg, and R. M. Wightman, "Failure of Standard Training Sets in the Analysis of Fast-Scan Cyclic Voltammetry Data.," *ACS Chem. Neurosci.*, vol. 7, no. 3, pp. 349–359, Mar. 2016, doi: 10.1021/acscchemneuro.5b00302.
- [24] P. Puthongkham and B. J. Venton, "Recent advances in fast-scan cyclic voltammetry.," *Analyst*, vol. 145, no. 4, pp. 1087–1102, 2020, doi: 10.1039/c9an01925a.
- [25] P. Takmakov, C. J. McKinney, R. M. Carelli, and R. M. Wightman, "Instrumentation for fast-scan cyclic voltammetry combined with electrophysiology for behavioral experiments in freely moving animals.," *Rev. Sci. Instrum.*, vol. 82, no. 7, pp. 1–6, 2011, doi: 10.1063/1.3610651.
- [26] J. J. Clark, A. L. Collins, C. A. Sanford, and P. E. M. Phillips, "Dopamine encoding of Pavlovian incentive stimuli diminishes with extended training.," *J. Neurosci.*, vol. 33, no. 8, pp. 3526–32, 2013, doi: 10.1523/JNEUROSCI.5119-12.2013.
- [27] C. M. Cameron, R. M. Wightman, and R. M. Carelli, "Dynamics of rapid dopamine release in the nucleus accumbens during goal-directed behaviors for cocaine versus natural rewards.," *Neuropharmacology*, vol. 86, no. August, pp. 319–328, 2014, doi: 10.1016/j.neuropharm.2014.08.006.
- [28] N. T. Rodeberg, J. A. Johnson, C. M. Cameron, M. P. Saddoris, R. M. Carelli, and R. M. Wightman, "Construction of Training Sets for Valid Calibration of in Vivo Cyclic Voltammetric Data by Principal Component Analysis.," *Anal. Chem.*, vol. 87, no. 22, pp. 11484–11491, 2015, doi: 10.1021/acs.analchem.5b03222.
- [29] C. J. Kimble *et al.*, "Wireless instantaneous neurotransmitter concentration sensing system (WINCS) for intraoperative neurochemical monitoring.," in *2009 Annual International*

Conference of the IEEE Engineering in Medicine and Biology Society, Sep. 2009, no. September 2009, pp. 4856–4859, doi: 10.1109/IEMBS.2009.5332773.

- [30] Y.-T. Li, J. R. Wickens, Y.-L. Huang, W. H. T. Pan, F.-Y. B. Chen, and J.-J. J. Chen, “Integrated wireless fast-scan cyclic voltammetry recording and electrical stimulation for reward-predictive learning in awake, freely moving rats.,” *J. Neural Eng.*, vol. 10, no. 4, p. 046007, 2013, doi: 10.1088/1741-2560/10/4/046007.
- [31] S. D. Adams, E. H. Doeven, S. J. Tye, K. E. Bennet, M. Berk, and A. Z. Kouzani, “TinyFSCV: FSCV for the Masses,” *IEEE Trans. Neural Syst. Rehabil. Eng.*, vol. 28, no. 1, pp. 133–142, 2020, doi: 10.1109/TNSRE.2019.2956479.
- [32] T. Scardochio, I. Trujillo-Pisanty, K. Conover, P. Shizgal, and P. B. S. Clarke, “The Effects of Electrical and Optical Stimulation of Midbrain Dopaminergic Neurons on Rat 50-kHz Ultrasonic Vocalizations,” *Front. Behav. Neurosci.*, vol. 9, no. December, pp. 1–15, 2015, doi: 10.3389/fnbeh.2015.00331.
- [33] A. L. Collins, V. Y. Greenfield, J. K. Bye, K. E. Linker, A. S. Wang, and K. M. Wassum, “Dynamic mesolimbic dopamine signaling during action sequence learning and expectation violation,” *Sci. Rep.*, vol. 6, no. February, pp. 1–15, 2016, doi: 10.1038/srep20231.
- [34] T. Claycomb *et al.*, *Analog Engineer’s Circuit Cookbook: Amplifiers*, 2nd ed. Texas Instruments, 2019.
- [35] D. J. Michael, J. D. Joseph, M. R. Kilpatrick, E. R. Travis, and R. M. Wightman, “Improving data acquisition for fast-scan cyclic voltammetry,” *Anal. Chem.*, vol. 71, no. 18, pp. 3941–3947, 1999, [Online]. Available: http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Citation&list_uids=10500480.
- [36] J. J. Clark *et al.*, “Chronic microsensors for longitudinal, subsecond dopamine detection in behaving animals,” *Nat. Methods*, vol. 7, no. 2, pp. 126–129, 2010, doi: 10.1038/nmeth.1412.
- [37] G. Paxinos and C. Watson, *The rat brain in stereotaxic coordinates*, 5th ed. Academic Press, 2004.
- [38] L. M. O. Oliveira and D. Dimitrov, “Surgical Techniques for Chronic Implantation of Microwire Arrays in Rodents and Primates,” in *Methods for Neural Ensemble Recordings*, 2nd ed., M. A. L. Nicolelis, Ed. Boca Raton: CRC Press, 2008.
- [39] F. Moussy and D. J. Harrison, “Prevention of the Rapid Degradation of Subcutaneously Implanted Ag/AgCl Reference Electrodes Using Polymer Coatings,” *Anal. Chem.*, vol. 66, no. 5, pp. 674–679, 1994, doi: 10.1021/ac00077a015.
- [40] P. Hashemi *et al.*, “Chronically implanted, nafion-coated Ag/AgCl reference electrodes for neurochemical applications,” *ACS Chem. Neurosci.*, vol. 2, no. 11, pp. 658–666, 2011, doi: 10.1021/cn2000684.
- [41] S. N. Berger and P. Hashemi, “Brain Chemistry: Neurotransmitters,” in *Reference Module in Chemistry, Molecular Sciences and Chemical Engineering*, 3rd ed., Elsevier, 2018, pp. 316–331.
- [42] E. S. Bucher and R. M. Wightman, “Electrochemical Analysis of Neurotransmitters,” *Annu. Rev. Anal. Chem.*, vol. 8, no. April, pp. 239–261, 2015, doi: 10.1146/annurev-anchem-071114-040426.