Unraveling the complex electrochemistry of serotonin using engineered graphitic sensors†

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Fast-scan cyclic voltammetry (FSCV) with micron-sized carbon sensors is a promising approach for monitoring the fast dynamics of serotonin (5-HT) neuromodulatory signals in the brain. However, sensor performance using FSCV can be compromised by complex chemical reactions associated with the reduction and oxidation of 5-HT, posing considerable challenges to detection of 5-HT in vivo. Herein we describe the use of engineered graphitic sensors to characterize the complex electrochemistry of 5-HT under a wide range of measurement conditions, with the aim of optimizing the FSCV conditions for in vivo quantitative 5-HT detection. These measurements reveal that water plays a significant role in driving side reactions during low-voltage FSCV measurements, leading to the observation of a well-defined secondary redox couple associated with the redox reaction of tryptamine 4,5-dione. Remarkably, these side reactions can persist subsequent to the primary redox events associated with 5-HT. Furthermore, the results reveal a critical deviation from this ideal redox behavior if the FSCV anodic limit exceeds +0.8 V, which can be attributed to the generation of radical species from water oxidation. These new insights could lead to new FSCV protocols for more reliable 5-HT detection.

Serotonin (5-HT) is a widespread neurotransmitter in the brain with implications for psychiatric and neurological disorders.1–5 Hence, there is significant research interest in developing technologies for monitoring the dynamics of 5-HT release and uptake in the brain with high spatial (microscale) and temporal (milliseconds) resolution, including genetically-encoded fluorescent protein sensors6–11 and fast-scan cyclic voltammetry (FSCV) with micron-size carbon electrodes.12–23 FSCV is a particularly attractive choice for human translation owing to the absence of inherent ethical and safety barriers. Carbon sensors foul during FSCV detection of 5-HT, however.24 While this fouling is often attributed to byproducts from side reactions of 5-HT,13,14,25,26 a knowledge gap pertaining to the identity and mechanism of 5-HT side reactions under FSCV conditions remains.

A pioneering work by Wrona et al., provides the most comprehensive account for the possible electrochemical reactions in linear sweep voltammetric measurements of 5-HT at physiological pH.27 Their study described a series of electrochemical and chemical reaction schemes that can occur during the redox of 5-HT at slow sweep rates (i.e., sweep rate $\nu = 200$ mV s$^{-1}$). An important finding of this study was the identification of a dominant electroactive side product formed in the presence of water. This side product displayed a distinct characteristic with increasing $\nu$ over a fixed potential sweep range from $-0.35$ V to $+0.6$ V. Critically, those experiments revealed a considerable suppression of this side reaction with increasing $\nu$, such that only the primary redox peaks of 5-HT were observable at $\nu = 50$ V s$^{-1}$. This finding suggests that employing fast sweep rates could mitigate sensor fouling caused by detrimental side reactions of 5-HT.

Sensor fouling remains a severe problem in FSCV detection of 5-HT regardless of the high sweep rates (>100 V s$^{-1}$).13,25 Although this problem is broadly attributed to side products of 5-HT redox reactions,23,26 the identity and mechanism of those reactions in FSCV measurements remain unknown. Indeed, the typical cyclic voltammograms (CVs) of 5-HT in FSCV measurements with the classic waveform ($\nu = 400$ V s$^{-1}$, potential sweep from $-0.4$ V to $+1.3$ V) display a shoulder next to the primary oxidation peak of 5-HT, which suggests the presence of following side reactions.25,28 Jackson et al., hypothesized that this shoulder peak is due to the addition of water to the phenylene ring, which they aimed to suppress in their study by modifying the parameters of the FSCV waveform (i.e., potential limits and sweep rate).25 The observation of this
A closer look at these previous reports suggests that the potential sweep range could play a role in producing unknown side reactions in the FSCV measurements of 5-HT. However, studying the effect of the potential sweep range under the FSCV conditions is not possible using existing carbon sensors, as they require a wide potential sweep range (at least from −0.6 V to +1.0 V) to reveal redox peaks of 5-HT and its side products under the typical FSCV conditions (i.e., \( \nu \geq 100 \text{ V s}^{-1} \)). Another significant challenge when studying the side reactions using existing sensors is the broad and generally ill-defined shapes of their corresponding redox peaks in the CVs of 5-HT. This poses a considerable challenge to accurately determine \( E^0 \), preventing a reliable identification of electroactive compounds from the CVs. Due to these shortcomings of existing sensors, a comprehensive study of 5-HT electrochemistry under FSCV conditions has remained elusive.

Herein we describe a comprehensive study of 5-HT electrochemistry to address three major questions. First, how does the potential sweep range under the FSCV measurement conditions (\( \nu \geq 100 \text{ V s}^{-1} \)) influence side reactions of 5-HT oxidation? Second, how do those side reactions evolve by varying \( \nu \) from a relatively slow rate (10 V s\(^{-1}\)) to a fast rate (200 V s\(^{-1}\))? Third, what is the underlying mechanism of those side reactions? The key to this study here is the use of a new generation of FSCV sensors made from engineered graphitic carbon materials, which were recently developed by our group.\(^{29}\) These sensors offer high sensitivity. They also provide sharp and well-defined redox peaks, allowing FSCV operation at low potentials inaccessible to the existing carbon sensors.\(^{30}\) Our experiments provide new insights into the identity and mechanism of 5-HT side reactions, which reconcile the inconsistent past observations. Lastly, we demonstrate the application of these insights for optimizing the FSCV protocols for selective and sensitive detection of 5-HT.

## Results and discussion

### Engineered graphitic sensors

Engineered graphitic sensors with a typical size of 20 × 20 µm\(^2\) were used for the experiments described herein. These sensors were fabricated on oxide-coated silicon substrates following protocols previously described by our laboratory\(^{29}\) (see Methods). Fig. 1a–c illustrate the schematic illustration, typical optical and SEM images of a graphitic sensor. The strong D peak in the Raman spectrum of Fig. 1d is consistent with a high density of defects, which we demonstrated previously to be crucial for the high FSCV sensitivity of the resulting sensors.\(^{29,31}\) Furthermore, our materials engineering approach yields fully graphitic carbon materials, evidenced by the sharp G and strong 2D Raman peaks (see Fig. 1d). This structural property of our materials results in electrochemical sensors with a near-ideal capacitive behavior, which was confirmed by the electrochemical impedance measurements (see Fig. 1e). The frequency response plots indicate a slight change of capacitance (<10%) and a phase response close to −90° at frequencies below 3 kHz. These results demonstrate that our sensors can maintain their near-ideal capacitive behavior in the frequency range of FSCV measurements (gray shading in Fig. 1e). This characteristic features a crucial distinction of these engineered graphitic sensors from conventional carbon sensors, which have a mixture of resistive-capacitive behavior.\(^{22,34}\)

The engineered structure of these graphitic carbon materials results in electrochemical sensors that produce sharp and well-defined peaks in FSCV measurements of electroactive analytes.\(^{29,30,34}\) As discussed below, this unique feature allowed us to examine the effects of the FSCV potential sweep range, revealing the unexpected role of the anodic potential limit (\( U_0 \)) in driving 5-HT side reactions.

### Effect of the FSCV waveform parameters

A series of FSCV measurements were performed to examine how the potential sweep range influences the 5-HT electrochemistry. All CVs in this section were recorded while flowing the 5-HT solution over the sensor (see Methods for details). In Fig. 2a–d, we plotted the background-subtracted CVs of 1 µM 5-HT. These CVs were obtained from the same sensor at \( \nu = 200 \text{ V s}^{-1} \) and a repetition scan frequency of 10 Hz. The potential sweep range in these measurements was selected such that the CVs can capture the full shape of all observable oxidation and reduction peaks.

The measurements started by obtaining the CV using \( U_0 = +1.15 \text{ V} \), as shown in Fig. 2a. The general shape of the CV is consistent with the previous observation of the 5-HT measurements using the classic FSCV waveform.\(^{28}\) Sharp oxidation and reduction peaks at +380 mV and +270 mV are observable, corresponding to a redox couple with \( E^0 = +325 \text{ mV} \). This \( E^0 \) is the signature of 5-HT molecules (\( E^i_{5HT} = +320 \text{ mV} \)). The product of 5-HT oxidation is expected to be \( p \)-quinone imine (compound II in Scheme 1[1]). Unlike the measurements at low sweep rates,\(^{27}\) the CV revealed more features that indicated the presence of unknown chemical reactions during the FSCV of 5-HT. An asymmetric and broad oxidation peak (marked with an arrow in Fig. 2a) is observable at the shoulder of the 5-HT oxidation peak. The CV also shows a second reduction peak at a negative potential of −400 mV and a significant level of residual current between the two observable reduction peaks. The peculiarities of these features suggest that the 5-HT side reactions could be more complex than previously thought.\(^{25}\)

The FSCV measurements were repeated with an identical waveform except for \( U_0 \), and their background-subtracted CVs were then plotted, as shown in Fig. 2b–d. These results correspond to three measurements with \( U_0 \) of +1.0, +0.8, and +0.6 V, respectively. The side-by-side comparison of these results...
Fig. 1  Engineered graphitic sensors. (a) Schematic illustration of the engineered graphitic sensor and the measurement test setup. The sensors were fabricated on SiO$_2$-coated silicon substrates. (b) The optical image of an example sensor. The scale bar is 20 µm. (c) The representative SEM image of a graphitic sensor, showing the relatively smooth surface morphology of these sensors. The scale bar is 1 µm. (d) The typical Raman signature of an engineered graphitic sensor. (e) The measured frequency responses of capacitance and phase indicate the near-ideal capacitive behavior of the engineered graphitic sensors arising from the graphitic structure of the sensors. The gray panel marks the approximate bandwidth of the FSCV measurements.

Fig. 2  Effect of anodic potential. Typical background-subtracted CVs of 1 µM 5-HT measured at $\nu = 200$ V s$^{-1}$ using several waveforms from −0.6 V to different anodic potentials of (a) +1.15, (b) +1.0, (c) +0.8, and (d) +0.6 V. The arrow in panel (a) indicates the oxidation peak due to side reactions. The letters “pr” and “sc” denote the primary and secondary redox peaks of 5-HT reactions. (e) Comparison of the transient response of the two oxidative peaks ($I_{ox,pr}$ and $I_{ox,sc}$) of the 5-HT CV in panel (d) reveals the slower response of the secondary redox reaction. (f) The background-subtracted CVs at three different times, corresponding to data in panel (e). (g) The background-subtracted CV of 5-HT measured at $\nu = 10$ V s$^{-1}$, revealing the presence of two redox couples with identical $E^\circ$ values to those in panel (d). In panel (g) the cathodic potential is −0.4 V and the anodic potential is 0.5 V.
revealed crucial insights. We observed that the change of $U_a$ did not alter the peak positions of the primary redox couple (see ESI† for detailed analysis). This suggests that the apparent electron transfer kinetics of this redox couple remained unchanged in these measurements. In contrast, the features associated with the side reactions undergo stark transformations, suggesting the critical effect of the anodic potential limit on 5-HT side reactions. Interestingly, we also observed that other waveform parameters, such as cathodic potential and sweep rate, have barely any effect on 5-HT side reactions (see ESI†).

A close look at these results revealed that the amplitude of the oxidation shoulder begins to grow by decreasing $U_a$. Curiously, this feature appeared to complete its transformation into a nearly symmetric peak in FSCV with $U_a = +0.8$ V. It then maintained its shape characteristics at $U_a = +0.6$ V. Moreover, the amplitude of the reduction peak of the side reactions increased by lowering $U_a$. It also completed its shape transformation at $U_a = +0.8$ V. Remarkably, an observable residual current between the two reduction peaks also began to diminish at $U_a < +0.8$ V. This result indicates that the observed residual current in CVs with $U_a > +0.8$ V is an actual response of the sensor to 5-HT-related reactions.

Next, we probed the origin of the secondary redox peaks in the CVs of 5-HT in Fig. 2 by initially focusing on the CV with $U_a = +0.6$ V (Fig. 2d). Due to their sharp and well-defined shapes, we hypothesized that the secondary peaks in this CV originate from the redox reactions of a single electroactive compound. These properties of the secondary peaks allowed us to assign an $E^0$ of $-210$ mV to this compound. Moreover, analyzing the corresponding transient response of the sensor revealed the slower rise time of the secondary oxidation peak relative to the primary peak (Fig. 2e). In Fig. 2f, the CVs taken from three different times in this measurement illustrate the appearance of the secondary redox peaks following the primary reaction. By combining this observation with the assigned $E^0$ of $-210$ mV, we hypothesized that the secondary peaks are associated with the redox reactions of tryptamine 4,5-dione (i.e., compound IV in Scheme 1(2)).

This hypothesis is based on the proposed reaction mechanism by Wrona et al., suggesting that this compound arises from the nucleophilic attack by water on the quinone imine (compound II). Scheme 1(2) outlines the simplified version of this reaction mechanism. However, this assignment of the secondary redox couple in this analysis contradicts the observation of Wrona et al., which indicated the absence of this redox couple in CVs with a fast sweep rate.

Therefore, we tested the validity of this hypothesis next by studying the CV of 5-HT measured at a slow sweep rate of 10 V s$^{-1}$, as shown in Fig. 2g. This sweep rate was chosen because the redox couple in Scheme 1(2) was still observable at 10 V s$^{-1}$ in that previous study. The data in Fig. 2g indicated an $E^0$ of $-210$ mV for the secondary redox couple, which agrees with the $E^0$ in our FSCV measurements and that reported by Wrona et al. This analysis provided strong evidence that the secondary redox couple in Scheme 1(2) can persist in measurements employing fast sweep rates.

With the aid of Scheme 1(2), let us examine the observed evolution of the secondary redox peaks in Fig. 2a–d. A close comparison of the CVs indicates that $U_a = +0.8$ V corresponds to the onset of the transformation in shape and amplitude of the peaks associated with side reactions. Remarkably, this $U_a$ coincides with the onset of the water oxidation reaction at pH 7.4, which is known to generate chemically active species (e.g., hydroxyl radicals). It is reasonable to expect that these radical species react with 5-HT intermediate compounds, generating additional side reactions that are distinct from those of the secondary redox couple shown in Scheme 1(2). Accordingly, these new reactions will exhaust the intermediate compounds underlying the transformation of the well-defined secondary oxidation peak into a broad and asymmetric shoulder peak (e.g., see Fig. 2b and c). This picture is consistent with the sudden decrease in the amplitude of the secondary redox peaks in experiments with $U_a > +0.8$ V. The observed evolution of the secondary redox peak is a fundamental phenomenon and can be generalized to other electrodes, such as commercially available carbon fiber microelectrodes (see ESI†).

These other side reactions—arising from radical species due to water oxidation—have not been accounted for in the past FSCV studies of 5-HT$^{1,25,26}$ and could be an additional cause for the sensor fouling. Therefore, as we demonstrate later, a desirable strategy for robust detection of 5-HT is to design new FSCV waveforms with a potential sweep range that avoids the water oxidation reaction.

**Experimental understanding of the secondary redox couple**

Our results in the previous section suggest that Scheme 1(2) is the mechanism behind the observed secondary redox couple. To corroborate this finding, we designed two experiments described next. These experiments aimed to visualize the two major features of Scheme 1(2): (i) the secondary couple is a follow-up product of the primary reaction, and (ii) the secondary reaction is reversible. All experiments in this section were performed without the flow of 5-HT solution (i.e., static 5-HT solution in the chamber; see Methods).

According to Scheme 1(2), the secondary redox couple occurs subsequent to the electrochemical oxidation of 5-HT. Therefore, the first experiment examined whether the ox-
vation of 5-HT is essential for the formation of the secondary redox couple. To do so, four FSCV experiments were performed using the same sensor. The parameters of the FSCV waveforms were identical among these experiments except for their $U_a$. The waveforms had $\nu = 200$ V s$^{-1}$, a 100 ms time interval between the scans, and a rest potential of $-0.6$ V. In Fig. 3a–d, the color plots display the amplitude of the background-subtracted current of 5-HT measurements corresponding to each waveform.

Fig. 3a displays the color plot of the measurement performed with $U_a = +0.2$ V. This anodic potential is not high enough (i.e., $U_a < E^\circ_{5-HT}$) to oxidize 5-HT, allowing us to study the potential range where only the secondary reaction should happen. No visible peaks are observable in this plot (see ESI† for the representative CVs). This result reveals that the secondary reaction does not occur in the absence of 5-HT oxidation.

The subsequent experiments used $U_a > E^\circ_{5-HT}$. Fig. 3b shows the color plot for $U_a = +0.35$ V. Although this anodic potential is not high enough to obtain a full oxidation peak of 5-HT in the CV plot (see ESI†), it is sufficient to initiate the oxidation of 5-HT. Remarkably, redox peaks associated with the secondary couple became discernable in this plot (marked with arrows). This experiment provides unambiguous evidence that the secondary redox couple forms after the electrochemical oxidation of 5-HT, hence supporting the mechanism in Scheme 1(2).

As larger $U_a$ (+0.45 and +0.6 V) was used in the two subsequent experiments, the steady-state amplitude of the secondary redox peaks became more prominent, as shown in Fig. 3c and d (see ESI† for representative CVs). This observation is consistent with the mechanism in Scheme 1(2).

Fig. 3 Dependence of the secondary couple on 5-HT oxidation. Color plots of 500 nM 5-HT redox reactions measured using waveforms from $-0.6$ V to different anodic potentials of (a) +0.2 V, (b) +0.35, (c) +0.45, and (d) +0.6 V. The data indicate that the secondary redox peaks appear only when the anodic potential is sufficiently high to initiate the oxidation of 5-HT. The red arrows show the location of the secondary redox peaks. The other two visible bands in the color plots (not marked with arrows) are associated with the redox peaks of the primary couple. All the measurements were performed in the absence of flow.

According to Scheme 1(2), the secondary redox couple involves reversible redox reactions. A reaction should sustain itself if it is reversible. On this basis, we designed another experiment to examine this property of the secondary couple by posing a question. How will the secondary redox reactions evolve if, after their stabilization, the primary redox reactions are interrupted?

The experiment for answering this question consisted of two sets of FSCV measurements. All measurements were made without the flow of the 5-HT solution. The first set of measurements used an FSCV waveform with a potential sweep range of $-0.6$ to $+0.6$ V, capturing the full redox peaks for both the primary and secondary redox couples. The scans were repeated for a duration of $\sim 30$ s at 100 ms time intervals and at $\nu = 200$ V s$^{-1}$. Fig. 4a displays the color plot of the background-subtracted current amplitudes, indicating the presence of the primary and secondary redox peaks. The plot in Fig. 4c shows three representative CVs of 5-HT at three different times (corresponding to the dashed lines in Fig. 4a). The data indicate that the measurement duration is long enough to permit all redox peaks to stabilize.

The second set of measurements interrupted the primary redox reaction by decreasing $U_a$ from +0.6 to +0.2 V. The other FSCV waveform parameters were kept unchanged. The CVs were then recorded for an additional $30$ s. Fig. 4b–d show the color plot of the current and three representative CVs. The
data revealed that the amplitude of the secondary redox peaks declined initially and then stabilized as the reaction appeared to reach an equilibrium. This experiment provides evidence for the reversible nature of the secondary redox reactions.

These are remarkable observations: we demonstrated two experiments employing an identical FSCV waveform (see Fig. 3a and 4b) that yielded two drastically different outcomes. These experiments established that although the formation of the secondary redox couple requires the oxidation of 5-HT (Fig. 3a), it can persist in the absence of the primary redox couple (Fig. 4b).

Application to engineering FSCV protocols

This section provides an example of how the insights of this study can be applied to engineer FSCV waveforms for sensitive, selective, and robust detection of 5-HT using engineered graphitic sensors.

According to the comprehensive study by Wrona et al., the intermediate products of the side redox reactions can serve as precursors that yield dimers and more complex molecules. These complex products can gradually cover the sensor surface through adsorption. This will consequently degrade the sensitivity by interrupting the electron transfer between the sensor and the target analyte. Therefore, avoiding the 5-HT side reactions is crucial for improving the reliability of sensors in repeated measurements of 5-HT.

Our findings guide us to employ an anodic potential limit below +0.8 V to avoid side reactions due to radical species generated by the water-splitting reaction. The earlier experiments at \( v = 200 \text{ V s}^{-1} \) indicated that \( U_a = +0.6 \text{ V} \) is high enough to capture the full shape of the primary oxidation peak of 5-HT.

Our experiments further indicated that under this condition, the byproduct of the secondary redox reaction has \( E^0 \) of −210 mV. Our experiments revealed that implementing a waveform with a hold potential \( (U_h) > -100 \text{ mV} \) can effectively mitigate the oxidation of this compound.

The choice of \( U_h \) is also crucial for obtaining an engineered waveform for specific detection of 5-HT. Therefore, we set out to experimentally engineer this parameter of the waveform to achieve highly sensitive detection of 5-HT, while at the same time the sensors remained insensitive to dopamine (DA)—another major neurotransmitter in the brain. To do so, we performed multiple FSCV measurements of DA and 5-HT with concentrations of 1 µM and 500 nM, respectively. In all experiments, we used an N-shape waveform with cathodic and anodic potentials of −0.2 and 0.6 V and only varied \( U_h \).

Fig. 5a shows the schematic illustration of this waveform. Fig. 5b–c show the corresponding CVs of DA and 5-HT measured at different \( U_h \). Fig. 5d shows the summary plot, illustrating the evolution of the normalized \( I_{\text{ox,peak}} \) for both DA and 5-HT plotted as a function of \( U_h \).

These experiments revealed that the normalized \( I_{\text{ox,peak}} \) of DA and 5-HT remained nearly unchanged below a critical hold potential \( (U_{h,c}) \) and began to decrease monotonically beyond it. From the data, we identified \( U_{h,c} \) for DA and 5-HT to be +50 and +210 mV. Moreover, the data also indicate that our graphitic sensors become insensitive to DA at \( U_h \) of about +120 mV. Therefore, for our engineered waveform, we chose \( U_h = +200 \text{ mV} \), for which our graphitic sensors simultaneously maintain their high sensitivity to 5-HT. Overall, this \( U_h \) ensures a waveform sensitive to 5-HT, insensitive to DA and it is high enough to avoid the oxidation of the secondary product.
Finally, the FSCV waveform should also permit the completion of the reduction process, which is crucial for balancing the electrochemical reactions by minimizing the accumulation of the reductive products. Furthermore, a well-defined reduction peak can aid in identifying the detected analytes. For these reasons, we set the lower potential limit of the FSCV waveform at +50 mV.

An N-shaped FSCV waveform allows the incorporation of these potential limits. For the proof-of-concept experiments, we created an N-shaped waveform with a repetition scan frequency of 10 Hz and \( \nu = 200 \text{ V s}^{-1} \). Fig. 6a illustrates this waveform, which we refer to as the "SER" waveform. We employed the SER waveform for the in vitro detection of 5-HT and DA using the engineered graphitic sensors. Fig. 6b shows the superimposed CVs of 5-HT and DA measured using the SER waveform, indicating the high sensitivity of 37 ± 0.3 nA \( \mu \text{M}^{-1} \). The nearly unchanged CVs of 5-HT after 15 trials (see Fig. 6d) indicate the prospects of the SER waveform in improving the robustness of the sensor performance for 5-HT sensing experiments.

Conclusions

The study reported here revealed three critical new insights into the complex electrochemistry of 5-HT under FSCV conditions. First, it established that the secondary redox couple (Scheme 1(2)) persists at fast sweep rates. Second, the strength of the secondary reactions depends on the number of water molecules in the solution. Our experiments revealed a sudden decline in the amplitude of the secondary redox peaks once the anodic potential limit exceeded +0.8 V, which coincides about 5 nM—which is at the lower end of the physiological levels in the brain.

Lastly, we examined how the sensor performance evolves in the repeated experiments of 5-HT detection employing the SER waveform. Past FSCV studies have shown that the 5-HT-related sensor fouling typically manifests itself within the first few trials, where each FSCV experiment is one trial. With this in mind, the sensor performance was examined by performing 15 consecutive trials. The nearly unchanged CVs of 5-HT after 15 trials (see Fig. 6d) indicate the prospects of the SER waveform in improving the robustness of the sensor performance for 5-HT sensing experiments.

Fig. 5 Waveform engineering for 5-HT-specific detection over DA. (a) Schematic of the N-shape waveform used for determining \( U_h \). In all measurements the cathodic potential is −200 mV and the anodic potential is +600 mV. \( U_h \) varies for different measurements. (b) Superimposed CVs of 1 \( \mu \text{M} \) DA measured with \( U_h \) of −50, 0, 50, 70, 90, 100, and 120 mV. (c) Superimposed CVs of 500 nM 5-HT measured with \( U_h \) of 0, 50, 100, 200, 220, 250, 270, and 300 mV. The dashed lines in panels b and c mark \( U_{h,c} \). (d) Normalized \( I_{\text{ox,peak}} \) plotted against \( U_h \). The dotted line shows the +200 mV position.

Fig. 6 Application to waveform engineering for 5-HT detection. (a) Schematic of the proposed SER waveform, which was engineered by applying the insights of our study. (b) Superimposed CVs of 50 nM 5-HT and 500 nM DA measured using the SER waveform, revealing the high selectivity for detecting 5-HT over DA. (c) 5-HT calibration curve displaying the high sensitivity of the engineered graphitic sensor and its high degree of linearity. The slope of the linear fit corresponds to a sensitivity of 37 ± 0.3 nA \( \mu \text{M}^{-1} \). (d) CVs of 200 nM 5-HT indicate minimal degradation after 15 repeated trials.
with the onset of the water-oxidation reaction at pH 7.4. Lastly, and most critically, the side reactions begin to deviate from the ideal redox behavior in Scheme 1(2) at anodic potentials exceeding +0.8 V. Additional unknown side reactions appear to occur due to the generation of radical species by the water-oxidation reaction. Although this study unraveled the mechanism of these new side reactions, their identity remains unknown. Future studies should focus on identifying these unknown side products in the mixture by combining the electrochemical measurements with analytical techniques (e.g., liquid chromatography). Finally, we established, through a design example, how these insights could contribute to new FSCV protocols with the goal of improving the quantitative detection of 5-HT in vivo.

Methods
Fabrication of engineered graphitic sensors
The synthesis of engineered graphitic carbon followed the same steps described in our previous studies. Briefly, we first patterned micron-size SU8 islands directly on SiO2-coated silicon substrates using electron-beam lithography (EBL). The SU8 islands were then annealed at 450 °C in a non-oxidizing ambient (80/20 mixture of Ar/H2 gas) and converted into a fully sp2 amorphous carbon. Next, we deposited an ultra-thin layer of nickel (<1 nm) in an ultra-high vacuum e-beam evaporator, followed by annealing at 1100 °C in a dual heating zone MTI furnace. We then converted the graphitic carbon islands into sensors employing standard nanofabrication techniques. High-quality Cr/Au metal leads and contact pads were formed through a sequence of EBL, e-beam metal evaporation, and metal lift-off. The formation of the SU-8 protection layers on the metal leads through EBL completed the sensor fabrication process.

Electrochemical measurements
We used 1X phosphate-buffered saline (PBS) with pH 7.4 as the buffer solution. The serotonin (5-HT) and dopamine (DA) solutions with known concentrations were prepared by dissolving their corresponding chemical powders in PBS. All solutions were delivered into a custom-made Y-shaped microfluidic chamber using two program-controlled syringe pumps (World Precision Instruments). A chlorinated silver wire (Ag/AgCl) was used as the reference electrode. The current output of the sensors during FSCV was recorded using a custom-made detection circuitry and digitized using a data acquisition instrument (NI USB-6363 series, National Instruments). The data acquisition and analysis were performed using a custom-made user interface program developed in MATLAB. For experiments in Fig. 2 and 5, the FSCV data were recorded while the PBS and 5-HT solutions flowed into the chamber. For experiments in Fig. 3 and 4, the FSCV data were obtained after stopping the flow of these solutions.

Chemicals
Serotonin hydrochloride powder was first dissolved in 1X PBS to form a stock concentration of 2 mM. 5-HT solutions with the desired concentrations were then prepared by diluting the stock solution in 1X PBS. The same procedure was used to prepare DA solutions, starting from dopamine hydrochloride. The 1X PBS (pH of 7.4) was prepared by dissolving 16 g of sodium chloride, 2.88 g of sodium phosphate dibasic, 480 mg of potassium phosphate monobasic, and 400 mg potassium chloride in 2 liters of deionized water. All chemicals were purchased from Sigma Aldrich.

Conflicts of interest
The authors declare no competing interests.

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