Wearable electrochemical glove-based sensor for rapid and on-site detection of fentanyl

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\textbf{ABSTRACT}

Rapid, on-site detection of fentanyl is of critical importance, as it is an extremely potent synthetic opioid that is prone to abuse. Here we describe a wearable glove-based sensor that can detect fentanyl electrochemically on the fingertips towards decentralized testing for opioids. The glove-based sensor consists of flexible screen-printed carbon electrodes modified with a mixture of multiwalled carbon nanotubes and a room temperature ionic liquid, 4-(3-butyl-1-imidazolio)-1-butanesulfonate. The sensor shows direct oxidation of fentanyl in both liquid and powder forms with a detection limit of 10 μM using square-wave voltammetry. The “Lab-on-a-Glove” sensors, combined with a portable electrochemical analyzer, provide wireless transmission of the measured data to a smartphone or tablet for further analysis. The integrated sampling and sensing methodology on the thumb and index fingers, respectively, enables rapid screening of fentanyl in the presence of a mixture of cutting agents and offers considerable promise for timely point-of-need screening for first responders. Such a glove-based “swipe, scan, sense, and alert” strategy brings chemical analytics directly to the user’s fingertips and opens new possibilities for detecting substances of abuse in emergency situations.

1. Introduction

Wearable sensors are well positioned to revolutionize healthcare monitoring through seamless integration in our daily routines [1–3]. For example, wearable biosensing has been demonstrated with micro-needles [4–7], epidermal microfluidics [8–11], and mouthguard-based sensor platforms [12] for continuous monitoring of electrolytes and metabolites. Other examples of cutting-edge wearable devices include tattoo sensors for non-invasive analysis of biomarkers, such as alcohol [13], glucose [14,15], and lactate [16] in epidermal biofluids (ISF and sweat). Glove-based sensors have been shown for detection of environmental and security threats of organophosphates [17], explosives, and gunshot residues [18]. Incorporating chemical sensors into wearable platforms (e.g., gloves, microfluidics, tattoos, etc.) offer the supremacy of lab-centered chemical analysis in a convenient form at that can be measured anywhere [19].

While most early wearable sensors to date have focused on fitness, wellness, and healthcare, there are tremendous opportunities for security and forensic applications of these devices [20,21] and towards monitoring drugs of abuse [22,23]. Drug abuse and misuse have increased significantly in recent years and is a rapidly growing and major problem for our society with tremendous costs (> $400 billion annually) due to lost productivity, healthcare needs, and legal processes [24]. Specifically, the misuse of prescription pain medications has become a public health epidemic, including the diversion of prescribed medications, such as oxycodone, buprenorphine, morphine, and fentanyl [25].

Fentanyl is a sedative and a Schedule II drug under the controlled substance act. It is currently being used as an analgesic and anesthetic. It is 50 to 100× more potent than morphine and 30 to 50× more potent than heroin. Fentanyl has been progressively used as an adulterant in illegitimate substances, such as heroin [26]. In 2015, the Drug Enforcement Agency issued a nationwide alert calling fentanyl a “threat to health and public safety.” Moreover, recently the Centers for Disease Control and Prevention issued a new warning about the use of fentanyl in the illicit drug supply [27].

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Control and Prevention reported that fentanyl and associated analogues were connected with over half of the opioid overdoses in ten states during the second half of 2016 [27]. Opioids represent a grave threat to public health and safety, in particular to first responders trying to treat people who have overdosed. Even small amounts of opioids present can become aerosolized and toxic. Thus, identification of fentanyl in trace quantities is crucial to deal with the current widespread epidemic.

Currently, liquid chromatography (LC) with mass spectrometry (MS) and immunoassay techniques are the gold standard for fentanyl detection and quantification [28,29]. Recent advanced technologies include surface enhanced Raman spectroscopy (SERS), which offer sensitive detection of fentanyl in heroin samples with a limit of detection (LOD) of 100 ng/mL [30]. Electromembrane extraction coupled with voltammetric readout demonstrated detection of sufentanil (a fentanyl analogue) in urine and plasma samples, with an LOD of $2 \times 10^6$ mol/dm$^3$ [31]. Antibody-based biosensors, employing enzyme-linked immunosorbent assays (ELISAs) and automated homogeneous enzyme immunoassays have also shown potential applications for fentanyl detection with an LOD of 1 ng/mL [32]. However, these techniques all require laboratory-based instrumentation, as well as complex sample preparation and operation, precluding on-site detection by law enforcement officers or first responders. In this regard, electrochemical analysis provides a unique combination of high sensitivity, fast response, low cost, and ease of operation.

Herein, we report a wearable glove with flexible electrochemical sensors integrated on the fingertips for spot identification of illicit drugs, specifically fentanyl. This is the first demonstration of a wearable electrochemical opioid drug detection platform towards on-site fentanyl screening. Glove-based wearable sensors, capable of chemical sensing at the fingertips, have been developed recently for detecting gunshot residues and nerve-agent threats [17,18,33]. The new fentanyl glove detection device carries out the sampling and electrochemical sensing steps on different fingers, with the thumb finger used for collection of drug residues or powder samples along with the index finger (printed carbon electrodes coated with an ionic liquid/multiwalled carbon nanotube composite film) for fentanyl sensing (Fig. 1A and B). Electrochemical (voltammetric) measurements were performed after completing the “electrochemical cell” by joining the thumb (sampling) and index (sensing) fingers. These measurements relied on square-wave voltammetric analysis, where the anodic peak current response, corresponding to direct fentanyl oxidation, is sent to a nearby tablet (or smartphone) via a Bluetooth wrist-worn potentiostat. The resulting glove-based sensor system thus enables rapid (~1 min) screening of fentanyl in both powder and liquid forms, bringing the power of centralized lab testing to the point-of-need. This allows first responders to rapidly assess unknown suspicious powders in emergency situations and deliver appropriate care for the patient while taking necessary precautions themselves.

![Fig. 1. Overview of the proposed Lab-on-a-Glove concept (swipe, scan, and sense) for on-site detection of fentanyl; (A) Photograph of glove containing sensing finger modified with ionic liquid/MWCNT composition and sample collector. (B) Image showing the glove-based sensor with a portable electroanalyzer (back-view). The electrodes are connected via wires and a modified ring to a PalmSens potentiostat for on-site detection with wireless communication to a smartphone for rapid analysis and reporting of the SWV results. (C) Image showing suspicious sample collection in powder phase. (D) Joining of thumb (collector) and sensing (index) fingers after swiping a powder sample; inset shows the voltammograms of direct fentanyl detection in powder/liquid samples.](image-url)
2. Experimental

2.1. Chemicals and reagents

Fentanyl (1.0 mg/mL in methanol, certified reference material), acetaminophen, glucose, caffeine, theophylline, multi-walled carbon nanotubes (MWCNT, O.D. × L 6–13 nm × 2.5–20 μm), polyethyleneimine (PEI), the ionic liquid of 4-(3-Butyl-1-imidazolio)-1-butanesulfonate, xylene, dipotassium hydrogen phosphate (K2HPO4), potassium dihydrogen phosphate (KH2PO4), sodium acetate, potassium chloride, and agarose were all purchased from Sigma-Aldrich (MO, USA). Carbon and Ag/AgCl inks (Ercon, Inc., Wareham, MA) were used to print the sensors on green nitrile powder-free exam gloves purchased from Kimberly-Clark (Roswell, GA). Deionized water was used to prepare aqueous electrolyte solutions. All solvents and chemicals were of analytical grade and used without further purification.

2.2. Glove-based sensor fabrication

Fabrication of the disposable glove-based sensor was carried out utilizing a semiautomatic MPM-SPM screen printer (Speedline Technologies, Franklin, MA). A 125 μm thick stainless-steel stencil (Metal Etch Services, San Marcos, CA) was designed using AutoCAD and laser-cut. To facilitate a smooth and planar printing surface, a layer of carbon and an insulator layer to complete the sensing electrode. The screen-printed sensor was cured at 70 °C for 10 min after each layer was printed. The Ag/AgCl-based ink served as the reference electrode, whereas carbon ink was used for the working and counter electrodes. The transparent insulating layer was carefully printed on the sensor interconnects to provide a dielectric segregation of the three-electrode system and to subside sensor short-circuits. The same printing procedure was used to print a circular carbon pad (1 cm diameter) on the thumb finger.

2.3. Preparation of the ionic liquid modified glove-based sensors

Multiwalled carbon nanotubes (MWCNT) were added to an ethanol solution and sonicated for 1 h. Then, 2 mg IL, 4-(3-Butyl-1-imidazolio)-1-butanesulfonate, and PEI were added into the MWCNT organic phase solution and stirred for 2 h. The resulting solution was centrifuged to remove any unbound MWCNT, IL, and PEI in solution, resulting in the MWCNT-PEI-IL composite nanomaterial. Later, from the prepared nanomaterial (MWCNT-PEI composite (1:16 w/w ratio) and IL), a 10 mg sample was taken and dissolved in 50 μL of DI water. The mixture was sonicated for 1 min. The developed composition was further diluted 6 × in DI water and a final volume of 1.5 μL was drop-cast on the working electrode of the glove-based sensor. Each electrode was allowed to air dry for 10 min prior to use. For dry phase analysis, a semi-solid 1.0 wt % agarose gel was prepared in a mixture of 0.1 M PBS and dispersed on the sensor (index finger) to cover the surface and provide a conductive medium for electrochemical analysis. The thickness of the hydrogel was about 3 mm. Such hydrogel layer displayed high stability over 2 weeks when stored frozen for 10 min prior to use. For dry phase analysis, a semi-solid 1.0 wt % agarose gel was prepared in a mixture of 0.1 M PBS and dispersed on the sensor (index finger) to cover the surface and provide a conductive medium for electrochemical analysis. The thickness of the hydrogel was about 3 mm. Such hydrogel layer displayed high stability over 2 weeks when stored frozen for 10 min prior to use.

2.4. Electrochemical measurements

Electrochemical characterization was performed at room temperature using a PalmSens hand-held potentiostat (EmStat3 Blue with 10 × 6 × 3.4 cm³ dimensions, PalmSens, Houten, Netherlands) powered by a rechargeable Li–Po battery (Fig. 1B). The PalmSens electrochemical analyzer can perform square wave voltammetry. The obtained data were wirelessly transmitted to a tablet to perform the analysis. The electrochemical sensing studies were performed by completing the “electrochemical cell” by joining the thumb (sample collector) with the sensing (index finger) covered with 1.0 wt % agarose gel and recording fentanyl oxidation by square wave voltammetry (SWV) with optimized parameters of 10 Hz frequency, 20 mV amplitude, and 3 mV step potential.

2.5. Powder fentanyl swapping and scanning

Before contaminating the target surfaces (glass slides) with fentanyl, they were first rinsed with absolute ethanol, distilled water, and finally dried using air flow. Subsequently, the glass slides (10 × 10 mm²) were contaminated using a 5 μL aliquot of the fentanyl stock solution. The drop-casted fentanyl solution was dried to a residue through evaporation at room temperature for 3–4 h under a fume hood.

The glove-based sensors were worn for the sampling and detection steps and connected to an electrochemical analyzer. The fentanyl powder residues were collected by wiping contaminated glass slides using the thumb finger followed by subsequent fentanyl sensing performed by joining the collection and sensing (index finger) covered with 200 μL of 1 wt % agarose gel. After completing the electrochemical cell by joining the two fingers, the operator waited 2 min to perform the analysis. It should be noted that, to avoid any cross-contamination, each glove sensor was fabricated for a single use.

Safety note: According to United States Drug Enforcement Administration (DEA), fentanyl is potentially lethal, even at levels as low as 0.25 mg [34]. Therefore, all experiments including sample preparation and measurements that involved fentanyl handling were strictly conducted inside a fume hood along with wearing appropriate personal protective equipment (PPE) using a university approved handling protocol. To perform sample swapping (dried fentanyl solution on a glass slide), all preparations and experiments were performed in a sealed air bag-based glove box (4 gal. capacity). The complete setup was placed inside a fume hood with continuous air flow.

3. Results and discussion

3.1. Fentanyl sensing in liquid form

The glove-based wearable platform for fentanyl screening is shown in Fig. 1. The developed wearable sensor combines the screen-printed glove sensors with a portable electrochemical analyzer (Fig. 1C) to enable on-site fentanyl sampling and screening (Fig. 1D). The square wave voltammetry (SWV) technique exhibits a distinct electrochemical fingerprint for fentanyl based on its direct, irreversible oxidation on the carbon sensing electrode modified with the room temperature ionic liquid composite (MWCNT-PEI-IL)

The performance of the glove-based sensor was first assessed towards fentanyl screening in the liquid phase using buffer solution. Changes in the fentanyl concentrations were used to evaluate the dynamic behavior of the sensor. Fig. 2A shows square wave voltammograms of fentanyl in PBS (0.1 M, pH 7.4) on ionic liquid-modified sensors, upon increasing fentanyl concentrations over the range of 10–100 μM. These fentanyl levels result in a single and distinct oxidation peak at +0.65 V (vs. Ag/AgCl), which is attributed to the oxidation of the tertiary amine groups present in the fentanyl structure [31]. The resulting calibration curve (shown in Fig. 2B) demonstrates good linearity (R² = 0.988) with an estimated detection limit (LOD) of 10 μM (RSD = 3.2%, n = 6). It should be mentioned that the elevated background signal observed when increasing the fentanyl concentration on Fig. 2A could be due to the presence of methanol in the fentanyl samples. It is well known that ionic liquids improve electrochemical processes [35]; however, the 4-(3-Butyl-1-imidazolio)-1-butanesulfonate...
ionic liquid coating here is essential for the fentanyl detection. No distinguishable fentanyl signal was seen using bare carbon sensors without the ionic liquid modification.

3.2. Fentanyl sensing in powder form

After establishing the glove sensor response in liquid phase, the analysis was further extended to powder/dry phase to perform real-time and in-field analysis of fentanyl (see related Safety Note in Experimental section). As mentioned in the experimental section, a conducting agarose hydrogel (200μL of 1 wt %) that covers the sensing finger was used as an electrolyte for the electrochemical measurements. During the sampling of fentanyl residues (dried fentanyl residues on a glass slide), the collection finger with the printed circular carbon pad swiped the test surface. The presence of fentanyl was detected by joining the sensing and collection fingers where the conducting agarose gel completed the electrochemical cell. The resulting square wave voltammograms of fentanyl in powder samples are presented in Fig. 2C. It can be seen that a clear background signal (black curve) is recorded for the glove sensor before fentanyl swiping. After collecting a fentanyl sample, a sharp and distinct peak at +0.76 V (vs. Ag/AgCl) is observed, demonstrating the fentanyl electrochemical oxidation (Fig. 2C, red curve). Compared to the fentanyl oxidation peak of +0.65 V in liquid phase (Fig. 2A), lower electrolyte concentration in the powder form slows the oxidation kinetics and therefore a higher oxidation potential is expected. The analytical operation is based on the abrasive voltammetry technique (transfer of trace amounts of solid sample onto the surface of an electrode [36]) to rapidly identify the presence of the drug. According to Faraday’s law, ultra-small amounts of collected electroactive materials can give rise to significant currents (μA), sampling of trace amounts of a solid material onto the electrode surface is therefore sufficient for analysis.

As a control, Fig. 2D also illustrates the swiping and screening in the absence of fentanyl powders and shows no detectable signal. The elevated signal is likely due to the applied pressure while joining the figures, but there are no oxidation peaks present in the signal.

3.3. Electrochemical screening of fentanyl mixed with common cutting agents

Street drugs usually contain cutting agents or diluents. Thus, a sensitive and reliable sensor should be able to distinguish the target analyte in a heterogeneous sample. Common cutting agents such as acetaminophen, caffeine, glucose, and theophylline were selected to test the performance of the glove sensor. Fig. 3 demonstrates the high selectivity of the sensor toward fentanyl in both liquid and powder forms (Fig. 3D), along with the SWV response of different interferers. As shown in the measurements, only acetaminophen (Fig. 3B) and theophylline (Fig. 3F) undergo oxidation at around +0.12 V (vs. Ag/AgCl) and +0.80 V (vs. Ag/AgCl), respectively, which caused no interference for the fentanyl sensing.

Electrochemical sensing of fentanyl was performed in a mixture containing 100 μM of acetaminophen, caffeine and glucose in a liquid form (Fig. 4A). A well-defined fentanyl oxidation peak at +0.63 V (vs. Ag/AgCl) along with a gradual increase in signal upon raising the fentanyl concentration from 10–100 μM was observed. The results show
that these cutting agents do not introduce any drift or changes in the fentanyl peak, with results similar to the liquid-phase experiments presented in Fig. 2A. Similarly, in the mixed powder form (Fig. 4B), fentanyl oxidation showed a distinct peak around +0.72 V (vs. Ag/AgCl). These results demonstrate a highly selective glove-based sensor for fentanyl screening.

4. Conclusions

This work aimed at developing a glove-based wearable electrochemical sensor capable of detecting opioids. The concept was demonstrated by voltammetric detection of fentanyl in liquid and powder forms. The glove-based sensor was prepared by screen printing "sensing" and "sampling" fingers and coupling this with a wireless and portable electrochemical analyzer, suitable for on-site and one-touch sample-to-answer testing of suspicious samples. Our data demonstrate that the "Lab-on-a-Glove" sensor platform responds in a reproducible and selective manner and enables fentanyl detection in powder as well as in liquid samples in the range of 10–100 μM with a 10 μM LOD. Detection was also demonstrated in the presence of cutting agents with no observed interference. As such, the glove-based drug sensor addresses major challenges associated with reliable, rapid, on-site screening and field detection of opioids in connection with first responder safety, airport/border screening, and potential terror threats. Further improvement in the peak identification can be achieved using our recently developed cyclic-square-wave voltammetric operation that yields a distinct electrochemical fingerprint of fentanyl [37]. Continued advances in electronics, particularly integration with miniaturized on-ring electrochemical systems, will further enhance such important field screening applications.

Author contributions

A.B and R.K.M contributed equally to this work. R.S. prepared the MWCNT-PEI-IL composite nanomaterial. All authors contributed to the manuscript and have given approval to the final version of the manuscript.

Notes

The authors declare no competing financial interest.

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