RSC Advances



PAPER

View Article Online
View Journal | View Issue



Cite this: RSC Adv., 2021, 11, 13366

Widening the applications of the Just-Dip-It approach: a solid contact screen-printed ion-selective electrode for the real-time assessment of pharmaceutical dissolution testing in comparison to off-line HPLC analysis†

Khadiga M. Kelani, ab Osama M. Badran, Mamdouh R. Rezk, Mohamed R. Elghobashy ac and Sherif M. Eid 6 *c*

Over past years, the field of pharmaceutical dissolution testing has significantly expanded to cover not only the quality control of dosage forms, but also to play an important role in the bioavailability testing paradigm and screening of most formulations. These tests usually need a very long time sampling and monitoring, so that the automation of sampling is laborsaving. Problems often occur with these automatic devices due to sampling lines that may disconnect, crimp, carry over, become mixed up, or are inadequately cleaned. Potentiometric sensors, such as liquid contact (LC-ISE) or solid contact ion-selective electrodes (SC-SP-ISE), can provide timely data to be used for the real-time tracking of the amount of active pharmaceutical ingredients (APIs) released in the dissolution medium without these problems. In this work, we adopted the Just-Dip-It approach as a process analytical technology solution with the ultimate goal of advancing the ion selective sensors to their most effective use in pharmaceutical analysis. Two sensors were fabricated, the traditional LC-ISE and SC-SP-ISE. The sensing poly-vinyl chloride membranes of two electrodes were prepared using 2-nitrophenyl octyl ether as a plasticizer to soften the membrane, and the reduction in resistance to pioglitazone ions (PIO) permeability was achieved through the incorporation of sodium tetraphenylborate and calix[8]arene as a cationic exchanger salt and inclusion complexing ligand, respectively. Finally, prepared membranes were turned into the flexible perm-selective slices of hydrophobic plastic, which work as a barrier to other compounds, except for the PIO cation in the concentration range of 1 \times 10⁻⁶ to 1 \times 10⁻² M and 1 \times 10⁻⁵ to 1 \times 10⁻² M for SC-SP-ISE and LC-ISE, respectively. The challenges and opportunities of both sensors in comparison to a developed HPLC method were discussed for the dissolution testing of the combination dosage forms of pioglitazone. Potentiometric methods were validated according to IUPAC guidelines, while HPLC was validated according to ICH guidelines to ensure accuracy and precision.

Received 3rd January 2021 Accepted 22nd March 2021

DOI: 10.1039/d1ra00040c

rsc.li/rsc-advances

1. Introduction

Electroanalytical techniques have a great potential to become the technique of choice for the real-time tracking of active processes, such as drug release from nanoparticles, 1,2 the kinetics and thermodynamics of formation and the decay of nanoparticles, 3 cell apoptosis, 4 dopamine release from brain tissue, 5 and adsorption

processes.^{6,7} Since 1970s, with the development of more sensitive electronics, the applications of ISEs have expanded tremendously. The solid contact screen printed ion-selective electrodes (SC-SP-ISE) have emerged as an essential technique of the potentiometric electrochemical analysis with hundreds of publications.^{8,9} This may be due to the fact that SC-SP-ISE has many principle advantages concerning the small electrode area, small thickness, high-temperature tolerance, elimination of the liquid filling solution that is found in the traditional liquid contact ion selective electrode (LC-ISE), the fact that many substances (such as catalysts) can be incorporated in the screen-printing paste, and it is disposable and cheap so that the statistical validation of experimental results can be performed easily due to the presence of several electrode replicates.¹⁰

The Just-Dip-It approach provides the highest level of realtime monitoring using ion selective electrodes (ISE), in which

^aAnalytical Chemistry Department, Faculty of Pharmacy, Cairo University, Kasr El-Aini Street, ET-11562 Cairo, Egypt

 $[^]b$ Analytical Chemistry Department, Faculty of Pharmacy, Modern University for Technology and Information, Cairo, Egypt

^cAnalytical Chemistry Department, Faculty of Pharmacy, October 6 University, 6 October City, Giza, Egypt. E-mail: Sheriefmohammed@o6u.edu.eg; sherief055@icloud.com

[†] Electronic supplementary information (ESI) available. See DOI: 10.1039/d1ra00040c

Paper

the potentiometric method is fully integrated with the medium unit under investigation. This integration is a vital tool for tracking bio/chemical changes with time without losing information due to sampling intervals.³ The sensing membrane is just dipped into the medium in question to obtain instant results. Abd El-Rahman *et al.*¹¹ described a detailed study considering the Just-Dip-It approach for the inline quantitative determination of ipratropium bromide using ISEs in comparison to HPLC techniques in light of the 12 principles of green analytical chemistry. This study showed equivalent analytical results like HPLC, while significantly reducing the amount of solvent, energy consumption, and waste generation.

Another study¹² described the use of a dip-it sampler as an extraction device in mass spectrometry. The ISEs capability of providing continuously updated information during the real-time tracking of bio/chemical changes^{3,13} opened the way for its utilization for inline quantitative determinations and the dissolution testing of pharmaceutical preparations.

Dissolution testing can provide key information about the release of active pharmaceutical ingredients (API) from the market dosage forms, and this information is associated with the API bioavailability.14 The dissolution parameters and procedures are described in the European and Japanese pharmacopeias. In these procedures, when multiple sampling times are required, frequent samples have to be withdrawn for analysis. The withdrawn samples have to be replaced by a fresh dissolution medium at the specified temperatures.¹⁵ These tedious protocols may lead to frequent sampling for long periods, frequent solution replacement, re-adjustment of dissolution conditions after each measurement, the consumption of a large number of solvents, the generation of a large amount of waste, high energy consumption due to long operation periods, and great effort from the operator. Most of the traditional methods used for dissolution monitoring such as HPLC chromatographic separations or UV/VIS spectroscopy suffer from these obstacles that may lead to discontinuous dissolution profiles of drugs. Moreover, HPLC devices consume large amounts of samples, solvents, energy, and time during operation and trials.16 Great efforts have been spent to overcome these obstacles, such as using dissolution devices attached to UV/VIS detectors with automatic samplers or potentiometric ion-selective electrodes.17-19

ISE electrodes offer many advantages over the traditional techniques for quantitative real-time tracking of the dissolution of pharmaceutical preparations due to their simplicity, portability, economical operation, and being unaffected by excipients and colors. Moreover, these electrodes are insensitive towards possible air bubbles or undissolved substances in the dissolution solution in contrast to UV/VIS measurements, where these may lead to serious light scattering. Although the traditional LC-ISE has an inner filling solution that may be subjected to evaporation in the case of immersion in analyte solutions at high temperatures, it showed good performance for tracking the API release from their dosage forms. SC-SP-ISE can provide better performance for the real-time tracking of dissolution testing.

In the current work, pioglitazone hydrochloride (IUPAC name: 5-(11methyl)-1,3-thiazolidine-2,4-dione) was selected as

a model drug for this study (Fig. 1S†). It is mostly administered in combination dosage forms as an antidiabetic agent that acts primarily by decreasing insulin resistance. A single graphitic carbon disc SC-SP-ISE has been selected, as it creates a solid contact between the electron-conducting substrate and the sensing membrane to replace the liquid contact solution in the LC-ISE, working as an ion-to-electron transducer.²² This approach eliminates the solutions inside the electrode, permitting the usage of SC-SP-ISE for quantification at different conditions (such as high temperatures) with a better response in comparison to the traditional LC-ISE.

This work aimed to fabricate a sensitive sensor, such as an SC-SP-ISE and LC-ISE. Then, the performance of each electrode was tested towards the real-time tracking of the dissolution profile of pioglitazone in its combined dosage form to highlight the opportunities and obstacles of each sensor. The obtained results were compared with the results of a newly developed HPLC method to validate the results and discuss the performance of the proposed methods, and this was compared to the traditional technique. This work could greatly enhance the predictions of the *in vivo* dissolution performance of oral dosage forms, and be considered as a step forward toward the biowaiver concept.

2. Experiment section

2.1. Chemicals and reagents

Pioglitazone (PIO), alogliptin (ALG), calix[8]arene, and sodium tetrakis[3,5-bis(trifluoromethyl)phenyl]borate (Na-TFPB), 2nitrophenyl octyl ether (2-NPOE) were purchased from Sigma-Aldrich. Polyvinyl chloride (PVC), sodium phosphomolybdate hydrate (PM), sodium phosphotungstate tribasic (PT), and dibutyl sebathate (DBS) were obtained from Fluka (St. Louis, MO, USA). Tetrahydrofuran (THF) was obtained from (Poole, England). Methanol (HPLC grade) was obtained from Fischer Scientific (UK). Potassium chloride (KCl) and potassium dihydrogen phosphate (analytical grade) were obtained from Oxford Laboratory (Mumbai, India). Orthophosphoric acid was obtained from El-Nasr Pharmaceutical Company (Egypt). All other reagents were of analytical grade, and bi-distilled water was used throughout the work. The pharmaceutical dosage form; Oseni® tablets (alogliptin, 25 mg/pioglitazone, 45 mg per tablet), a product of Takeda Pharmaceuticals (USA) was purchased from the local market.

2.2. Instrumentation

2.2.1. The potentiometric method device. The potentiometric measurements were conducted using the JENWAY digital device Model 3505 (Essex, UK). The device was attached to the Calomel double junction $Hg/HgCl_2$ reference electrode (Beckman, USA). The screen-printed electrodes (type: single disk graphitic carbon SE101) with dimensions of 50×13 mm ($h \times w$) were purchased from (CH Instruments, USA). Glass pH electrodes (JENWAY, UK) were used for pH measurements. A magnetic stirrer with digitally controllable heater (IKEA, UK) and a sonicator Rx510S (Budapest, Hungarian) were also used.

2.2.2. The HPLC method device. The HPLC device was a Shimadzu LC-20AT series equipped with a degasser, quaternary pump, an automatic sampler, and a dual-wavelength detector (SPD-20AD Model). The Inertsil® C_{18} (150 \times 4.6 mm, 5 μ m) column was purchased from GL Science® (Tokyo, Japan). The device is controlled by LC solution software V.1.2 (Shimadzu, Japan) for data acquisition and manipulation.

2.2.3. The dissolution test device. The dissolution test was carried out using a Dissolution System Distek® model 2500 equipped with an Evolution Sampler 4300 (Distek, USA). The device is equipped with a USB-II paddle and bucket.

2.3. Standard solutions

2.3.1. For potentiometric method. About 0.098 g of PIO powder was accurately weighed and dissolved in the least amount of 0.01 M HCL into a 25 mL volumetric flask, and then the volume was made to mark using phosphate buffer solution (pH 4) to get a stock solution of 1×10^{-2} M. The remaining serial dilutions from 1×10^{-7} to 1×10^{-3} M were prepared by repeated transfer of 2.5 mL from the higher concentration to get the lower concentration, and then the volumes were made up to the mark using phosphate buffer. The prepared solutions were stored in the dark at -4 °C. There was no need to control the ionic strength of the buffer, so we did not add any salts.

2.3.2. For the HPLC method. Amounts corresponding to 100 mg of PIO and ALG powder were accurately weighed and placed into 100 mL volumetric flasks. The weights were dissolved in the least amount of mobile phase [20 mm phosphate buffer solution of pH 3: methanol (30:70, v/v)], and the volumes were made to the mark using the same solvent.

2.4. The process of fabrication of the sensors and their optimization

2.4.1. The preparation of the membranes and conditioning. The SC-SP-ISE membrane was prepared by transferring 0.01 g of Na-TFPB as an ion exchanger into a glass Petri-dish. Then, 0.015 g of calix[8]arene was added to the Petri dish as an ionophore, and mixed with 0.4 mL of 2-NPOE as a plasticizer. About 0.19 g of PVC was transferred into the Petri dish by mixing with 8 mL THF to get a homogeneous membrane cocktail. Exactly 10 microliters of the prepared cocktail was dropped and cast over the disk of SP-ISE using a micropipette to fabricate the SC-SP-ISE. The membrane was left for five hours at room temperature until the complete evaporation of THF, leaving a dry solid membrane on the SP-IE. The drop-casting²³ steps must be done carefully with a calibrated micropipette over a horizontal carbon disk to ensure homogeneous distribution of the cocktail over the carbon disk.

The LC-ISE membrane was fabricated using the same components in the same ratio by mixing the weights of Na-TFPB, 2-NPOE, and PVC in a 5 cm diameter clean glass Petri dish, followed by mixing with THF using a glass rod. Calix[8] arene was added to the mixture, and covered by paper to allow for slow solvent evaporation. The Petri dish components were left to dry for 12 hours at 25 °C. The obtained membrane with a thickness of about 0.1 mm in diameter is considered as

a master membrane, from which a small disk with an appropriate size (about 8 mm in diameter) has been cut and glued by THF onto the flat end of the PVC tube. Then, the PVC tube was attached to a glass micropipette from the other end to form an electrode body (LC-ISE), which was filled with equal concentrations (1 \times 10 $^{-2}$ M) of PIO and KCl aqueous solution as an inner filling solution.

Fig. 1 describes the design of the two sensors. The conditioning of the two sensors was performed through overnight soaking of the membranes with a 1 \times 10⁻² M pioglitazone solution.

2.4.2. Influence of pH on the electrode response. The response of the two sensors towards the change of pH was recorded in the pH range of 1–12 using two PIO concentrations 1×10^{-3} and 1×10^{-4} M. The pH values were adjusted using the appropriate amounts of 0.01 M NaOH and 0.01 M HCl.

2.4.3. Evaluation of the time required to get a steady response. The time required by the two sensors to get an acceptable stable potential with time was studied. This was done by the gradual increase of the concentration of the studied drugs by 10-fold within their linearity range by observing the change in the potential. Once a stable response was obtained, the time needed to obtain this constant response was determined using a stopwatch. A plot of the time in minutes against the electromotive force (emf) change was constructed for the accurate determination of the dynamic response time.

2.4.4. Construction of the calibration graphs. The conditioned sensors (LC-ISE and SC-SP-ISE) were immersed in combination with the calomel reference electrode into a series of 50 mL beakers containing different concentrations of PIO (1 \times 10 $^{-7}$ to 1 \times 10 $^{-2}$ M), prepared in a phosphate buffer solution of pH 4. It was then allowed to equilibrate, and the emf was recorded within ± 1 mV with continuous stirring. Pure water was used as a washing solution between measurements. The emf was then plotted against the logarithm of PIO concentration. The regression equations were then calculated from the linear range of the curve, and used for the subsequent calculation of the unknown concentration of PIO.

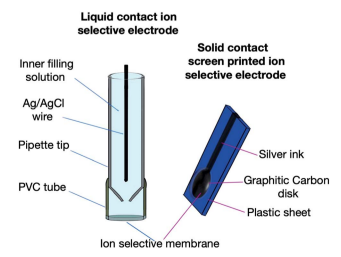


Fig. 1 Design and components of LC-ISE and SC-SP-ISE.

Paper RSC Advances

2.4.5. Evaluation of the membrane selectivity. The selectivity of the two sensors towards PIO was evaluated in the presence of the expected interfering substances, such as alogliptin, and other pharmaceutical excipients, such as mannitol, cellulose, magnesium stearate, lactose, talc, and ferric oxide. The separate solution method (SSM)²⁴ was followed, where the potential of 1×10^{-3} M of PIO and 1×10^{-3} M of the expected interferents were separately measured. The selectivity coefficients values ($k_{\rm ion\ interferent}^{\rm pot}$) were computed from the following equation:

$$\log \left(k_{\text{ion interferent}}^{\text{pot}}\right) = \frac{E_2 - E_1}{S} + \left(1 - \frac{A}{B}\right) \log aA$$

where the potentiometric selectivity coefficient is symbolized ($k_{\rm lon}^{\rm pot}$ interferent). "S" is the slope of the calibration plots for each sensor. " E_1 and E_2 " are potentials measured in 1×10^{-3} M of under-investigation ions and interferent ions, respectively. "A and B" represent the charges of the under-investigation and the interferent ions, respectively. "a" is the activity of the underinvestigation ions.

2.5. HPLC separation conditions and calibration

The ideal performance was obtained using a filtered and degassed mobile phase consisting of 20 mM phosphate buffer solution (pH 3): methanol (30:70, v/v), with isocratic elution at a flow rate of 1.35 mL min $^{-1}$, oven temperature at 30 $^{\circ}$ C, and the eluent was monitored at 265 nm. The separation was done using an Inertsil C_{18} (150 \times 4.6 mm, 5 μ m) column.

The linearity range was determined by transferring different volumes from the ALG and PIO standard solutions (100 $\mu g~mL^{-1}$) into a series of 10 mL volumetric flasks. Then, the volumetric flasks were completed to the mark using the mobile phase to get a concentration range of (10–80 $\mu g~mL^{-1}$) and (5–40 $\mu g~mL^{-1}$) for PIO and ALG, respectively. About 10 μL from each volumetric flask was injected and separated under the chromatographic condition. A linear relationship was obtained by graphing the peak areas against the corresponding concentrations of PIO and ALG, and then the actual concentrations were computed from the corresponding regression equations.

2.6. Real-time tracking of pioglitazone HCl release from the pharmaceutical dosage form (dissolution testing)

Drug absorption from the oral tablet dosage form depends on the drug release and dissolution under different physiological conditions and permeability through the gastrointestinal tract. Because of the crucial nature of these conditions, *in vitro* dissolution testing can be deemed relevant to try to expect the *in vivo* performance. Based on that, the dissolution test for PIO was carried out. The dissolution analysis was carried out in 0.01 M HCl as a recommended medium in FDA guidelines. The dissolution test was conducted by dropping one of the Oseni® tablets in one of the vessels (n=6) containing 900 mL of 0.01 M HCl preheated to 37 °C \pm 0.5 °C using the paddle (USP apparatus II), at a speed of stirring equal to 75 rpm. Then, the dissolution testing was constructed using the following steps, as outlined in the next sub-sections.

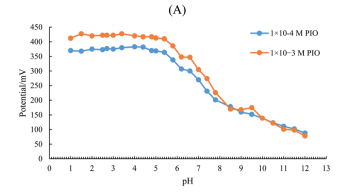
2.6.1. Dissolution test using the potentiometric method. The fabricated electrodes (LC-ISE and SC-SP-ISE) were immersed in the dissolution medium in combination with the reference electrode. The change in the potential with time was recorded directly (continuous sampling).

2.6.2. Dissolution test using the HPLC method. Manual sampling was performed by frequent withdrawing of solution samples (each sample was 1 mL) from the dissolution tube every two minutes (for the first ten minutes), and then every five minutes for the remaining time. The collected samples were filtered separately using a nylon filter (pore size equal to 0.45 μm), and the filtrate was analyzed by the optimized separation conditions. The withdrawn samples were replaced by the same amounts from a fresh dissolution medium, which was preheated to 37 °C \pm 0.5 °C, and these dilutions were taken into consideration in the calculations.

3. Results and discussion

3.1. The performance characteristics of the fabricated sensors

- **3.1.1. Membrane cocktail optimization.** The sensing membranes were fabricated carefully to show high selectivity towards the PIO ions. According to many previous studies, the sensitivity and selectivity of the proposed ion-selective membranes are significantly affected by the type and proportion of constituents involved in their fabrication, such as PVC, ionophore, ion exchanger, and plasticizer.²⁵ Accordingly, the mixture of these components was carefully chosen to be of the highest degree of selectivity required for PIO. This was performed by fixation of the membrane components, except for the targeted component:
- Initially, polyvinyl chloride (PVC) was selected because of its high lipophilicity ($\log P = 0.593$) and high molecular weight.²⁶ Moreover, its ability to increase the dielectric constant was beneficial.
- Different types of commonly used plasticizers, such as DBS, 2-NPOE, and DOP, were examined to obtain the most appropriate one. 2-NPOE was selected as the best plasticizer to obtain the optimum Nernstian slope of the monovalent cations. This may be due to the higher dielectric constant of 2-NPOE compared to DOP or DBS, which reaches 24, and its ability to increase the polarity of the membrane compared to the others. Moreover, 2-NPOE can solubilize the ionophore, soften the PVC membrane, and it is inert to ion exchangers and insoluble in water.²⁷
- Since the studied drug (PIO) acts as a cation, this fact suggests the selection of a suitable cationic exchanger. Several cationic exchangers have been tested, such as PT, Na-TFPB, and PM. However, non-Nernstian slopes were obtained using PM and PT, in addition to the non-reproducible response, poorer selectivity, lower sensitivity, and short life-time. On the other hand, the sensory membrane prepared using Na-TFPB provided a rapid response, reproducible Nernstian slope, and lower LOD. Therefore, it was elected as a suitable ion exchanger.
- Selectivity enhancement was performed through the incorporation of an ionophore. Two ionophores were examined,



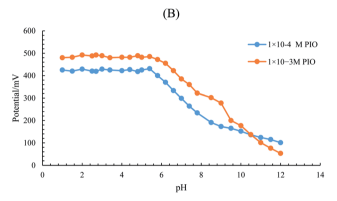


Fig. 2 EMF-pH profiles of 1 \times 10⁻³ and 1 \times 10⁻⁴ M concentrations of PIO using (A) LC-ISE and (B) SC-SP-ISE.

calix[8] arene and β -cyclodextrin (β -CD). Both ionophores have a large number of OH groups, which are considered as high donation sites that provide the ability for the formation of complementary hydrogen or ionic bonds and inclusion complexes with the positive PIO ions. The two ionophores were incorporated in the fabrication of two different membranes, and their performance was tested. The incidence of appropriate chelation does not occur with β -CD, while with

calix[8]arene, a stabilized host–guest complex occurs by the electrostatic attraction between the anionic calix[8]arene and the cationic PIO. Moreover, the internal cavity size of calix[8] arene (9.5 Å) is larger than that of β -CD (6 Å). As a result, calix[8] arene was suggested as a suitable ionophore for LC-ISE and SC-SP-ISE.

After the election of 2-NPOE and calix[8] arene as the suitable plasticizer and ionophore, respectively, their appropriate molar ratio of calix[8] arene to 2-NPOE was prepared to be 1:2 mole percent (mol%) (*i.e.*, the molar ratio of the ion exchanger is 50% relative to the used ionophore). This molar ratio follows many studies, 3,29 which proved the fitting of the analyte (PIO) in the inclusion sites of calix[8] arene.

The membrane polymeric matrix (PVC) is an impermeable layer due to its hydrophobic characteristics ($\log P = 0.593$), which necessitates the incorporation of hydrophobic 2-NPOE ($\log P = 4.858$) as a plasticizer to soften the membrane. Further reduction in the resistance to ion permeability was achieved through the incorporation of Na-TFPB ($\log P = 12.971$) and calix [8]arene ($\log P = 10.729$) as the cationic exchanger salt and inclusion complexing ligand, respectively. Finally, the prepared membrane was turned into a flexible perm-selective slice of hydrophobic plastic, which works as a barrier to other compounds except for the PIO cation ($\log P = 3.328$). The membranes for LC-ISE and SC-SP-ISE were fabricated using the same membrane components.

3.1.2. Effect of pH on the membrane response. The effect of the fluctuation of pH values on the electrode response was studied using 10^{-4} and 10^{-3} M solutions of PIO, where the two proposed LC-ISE and SC-SP-ISE revealed a stable response over the pH range (1.0–5.5), as shown in Fig. 2. Therefore, this pH range is considered as the working pH range of the two fabricated sensors. An irregular decrease was observed in the EMF response outside this range.

3.1.3. The dynamic response time of the fabricated sensor. The main factor that is usually used to evaluate the performance of the membrane sensors is the time it takes to obtain

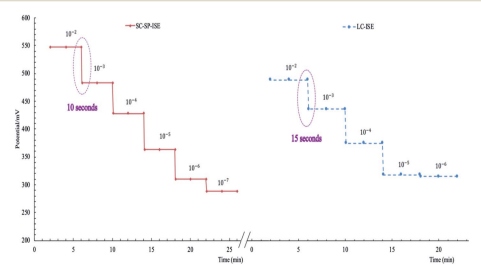


Fig. 3 Dynamic response time for SC-SP-ISE and LC-ISE showing the time required to obtain a stable potential.

consistent emf values (± 1 mV), which is known as the dynamic response time. Interestingly, the speed of the response is a good indicator that permits accurate monitoring of the drug release. Studying the practical response time was performed for the two studied sensors over their entire linearity range, and the appropriate time needed to get a steady response ($\pm 1 \text{ mV}$) was found to be 10 and 15 seconds for sensors I and II, respectively. The time traces of the two sensors (I and II) showed the superior performance of the SC-SP-ISE, as shown in Fig. 3, which indicates its ability to collect all required real-time information instantaneously and fits the idea of the real-time tracking of drug release.

3.1.4. The method calibration and validation data. Fig. 4 shows the linear range of the PIO concentrations between 1 \times 10^{-6} to 1×10^{-2} M and 1×10^{-5} to 1×10^{-2} M, with a good Nernstian slope equal to 59.4 and 57.2 mV per decade for SC-SP-ISE and LC-ISE, respectively. The limit of detection for the two proposed SC-SP-ISE and LC-ISE reached 2.4 imes 10 $^{-7}$ and 1.7 imes10⁻⁶ M, respectively.

The potentiometric performance was validated following the guidelines of IUPAC.30 The accuracy of the potentiometric readings of both sensors was confirmed by measuring blind PIO concentrations of $(1 \times 10^{-5}, 1 \times 10^{-4}, \text{ and } 1 \times 10^{-3} \text{ M})$. Repeatable and precise measurements were obtained, while applying the method on the same day to the same concentrations using the same conditions. Moreover, the intermediate precision was ensured by measuring the same concentrations on different days, and the calibration regression equations were used to calculate the unknown concentrations. Table 1

summarizes the calculated results, indicating high accuracy and precision from the calculated values of RSD ≥ 2 and the mean recovery percent within the range of 97-101%. The results indicate good performance of the potentiometric analysis for the detection of unknown PIO concentrations with high accuracy and precision.

3.1.5. The sensor selectivity towards pioglitazone. The selectivity of the sensors towards PIO was examined in the presence of different substances, including the co-formulated drug ALG and the expected tablets excipients (mannitol, cellulose, magnesium stearate, lactose, talc, and ferric oxide), in addition to some organic cations (like Ca2+ and Mg2+) that may be present in water. The separate solution method (SSM) has been applied, and the results in Table 2 indicate the high applicability of the method with high selectivity towards PIO.

3.2. HPLC method calibration and validation

Chromatographic methods are widely used for the quantitative analysis of active pharmaceutical ingredient release profiles. Therefore, the establishment and validation of an efficient, robust, and reliable RP-HPLC method for tracking the rate of drug release from the Oseni® tablet was one of the goals of our study. Different chromatographic conditions have been tested, including different pH values, mobile phase components, flow rate, buffer components, and stationary phase aiming for the separation of ALG from PIO with high resolution. The selected pH values at which the good resolution and sharp peaks for the two studied drugs were obtained was 3. The optimum separation of ALG and PIO was ideally performed utilizing the

Table 1 The characteristics of the response of the two sensors towards pioglitazone and the validation parameters^a

Parameters		SC-SP-ISE	LC-ISE	HPLC		
Analytes		PIO	PIO	PIO	ALG	
Slope		59.40	57.20	864.64	54.64 1464.70	
Intercept 663.80		663.80	604.80	-33.57	188.20	
The correlation coefficient (R^2)		0.9990	0.9990	0.9999	0.9998	
The linear range		1×10^{-6} to 1×10^{-2}	1×10^{-5} to 1×10^{-2}	$2.55 \times 10^{-5} \text{ to } 20.36 \times 10^{-5}$	1.27×10^{-5} to 10.18 \times 10^{-5}	
(molar)						
Working pH range		1-5.5	1-5.5	_	_	
Response time (s)		10	15	_	_	
Retention time (min)		_	_	2.824	1.179	
Validation param	neters					
Accuracy (mean recovery \pm SD)		99.27 ± 0.38	99.67 \pm 0.78	100.07 ± 0.48	100.13 ± 0.95	
Repeatability (RSD%)		0.66	0.73	0.75	0.55	
Intermediate precision (RSD%)		0.61	0.52	0.98	0.47	
LOD	,	$2.4 imes 10^{-7} ext{ M}$	$1.7 imes 10^{-6} ext{ M}$			
Robustness*	a	_	_	100.05 ± 0.80	99.94 ± 0.32	
	b			99.15 ± 0.58	100.24 ± 0.91	
	c			100.87 ± 0.63	99.84 ± 0.72	
	d			100.61 ± 0.78	99.08 ± 0.23	

^a * is the average recoveries of three determinations of both PIO and ALG using: (a) mobile phase at pH = 2.8, instead of 3; (b) mobile phase contains 67% methanol, instead of 70%; (c) flow rate at 1.15, instead of 1.35; and (d) column temperature at 28 instead of 30 °C.

Table 2 The selectivity of the two proposed sensors towards pioglitazone a

	Selectivity coefficient (mean \pm S.D.)					
Interferent ^b	LC-ISE		SC-SP-ISE			
Mannitol	1.41×10^{-3}	± 0.65	8.32×10^{-4}	± 0.11		
Cellulose	5.91×10^{-3}	± 0.78	6.98×10^{-4}	± 1.09		
Magnesium stearate	7.91×10^{-3}	± 1.23	11.07×10^{-3}	± 0.56		
Talc	6.81×10^{-3}	± 0.34	1.82×10^{-3}	± 0.88		
Lactose	5.14×10^{-3}	± 0.44	3.83×10^{-4}	± 1.23		
Ferric oxide	2.94×10^{-3}	± 0.89	2.81×10^{-3}	± 0.76		
Starch	10.37×10^{-3}	± 1.20	9.93×10^{-3}	± 0.58		
Ca ²⁺	11.07×10^{-3}	± 1.32	8.18×10^{-3}	± 0.90		
Mg^{2+}	9.24×10^{-4}	± 0.77	5.19×10^{-4}	± 1.11		
Alogliptin benzoate	8.86×10^{-3}	± 0.56	7.67×10^{-4}	± 0.67		

 $[^]a$ Each value is the average of three determinations. b All interferents are in the form of 1 \times 10 $^{-3}$ mol L $^{-1}$ solution.

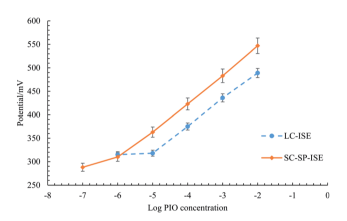


Fig. 4 Profile of the potential in mV *versus* log concentrations of PIO in mol L^{-1} obtained with (—) SC-SP-ISE and (- - -) LC-ISE.

degassed mobile phase of 20 mM phosphate buffer solution (pH 3): methanol (30:70, v/v), isocratic elution, flow rate at 1.35 mL min $^{-1}$, column temperature of 30 °C, and the eluent was monitored at a wavelength of 265 nm (the wavelength at which ALG and PIO have appropriate absorbance). The separation was done using an Inertsil C_{18} (150 \times 4.6 mm, 5 μm) column. The ALG and PIO retention times were 1.1 and 2.8 minutes, respectively, Fig. 5.

A linear relationship was obtained in the concentration range of (10–80 $\mu g\ mL^{-1}$), as shown in Fig. 2S.† The obtained linearity range was from 20–160% of the expected released PIO in a dissolution medium. The actual amount released from PIO in the dissolution medium at the first and last interval is completely covered by the range.

The HPLC method was validated according to the ICH guideline.31 Adequate accuracy for the proposed chromatographic method was established through the calculation of the mean recovery and RSD for different pure PIO concentrations. Assessment of precision was also employed, and the findings from inter-day and intra-day precision indicate that the chromatographed method was highly precise. The method's robustness has been assessed by evaluating the effect of small changes in the experimental conditions, such as flow rate, mobile phase composition, oven temperature, and pH. In all of the deliberate varied chromatographic conditions, good recoveries were obtained, as shown in Table 1. Also, the resolution between the two drugs was slightly affected, indicating a highly robust method. Table 1S† shows the system suitability parameters, and the results indicate that the method is suitable. Thus, it can be used for quantification of the drugs using representative chromatograms.

3.3. Tracking of pioglitazone ion activity during the dissolution test

Pioglitazone and alogliptin were formulated in a combination dosage form called Oseni® tablets. Oseni® tablets are film-

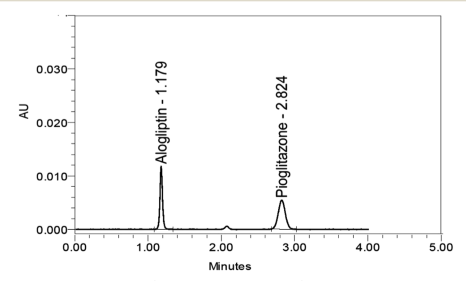
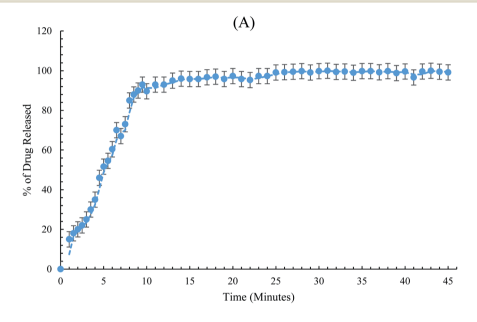


Fig. 5 HPLC chromatogram for alogliptin (20 μ g mL⁻¹) and pioglitazone (40 μ g mL⁻¹) using a C₁₈ column and 20 mM phosphate buffer (pH = 3): methanol (30 : 70, v/v) as the mobile phase and UV detection at 265 nm.

coated tablets that have been listed as an oral dosage form that should not be crushed before swallowing, as their content must be slowly released in the stomach. The tracking of its active component slow-release using traditional methods that require frequent sampling, such as HPLC, is a tedious process. Thus, the use of inline monitoring by ion-selective electrodes is advantageous. In particular, pioglitazone is more lipophilic (log P=2.12) than alogliptin (log P=-1.88), which increases the selectivity of the prepared lipophilic membrane towards pioglitazone. Also, the different lipophilicities facilitated the chromatographic separation.

3.3.1. In-line potentiometric method. Tracking of the PIO release from its tablet dosage form was performed utilizing the

selected SC-SP-ISE, where its superiority over the LC-ISE in potentiometric measurement has been approved. Some points have to be taken into consideration concerning the conventional LC-ISE, such as the fact that it has to be manually fabricated and optimized for each analyte as it must be incorporated not only in the membrane, but also in the inner filling solution. Moreover, the inner filling solvent may be susceptible to be evaporated during long periods of usage, leading to a change of the concentration of its component and occasional precipitation of KCl salt. Also, its performance may be affected by the unfavorable transmembrane fluxes from the concentrated internal components to the external membrane surface, as reported in the literature, ³² causing deterioration of the analytical



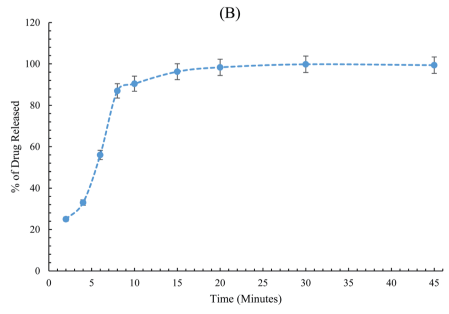


Fig. 6 Dissolution profiles for PIO in the Oseni® tablet obtained by (A) SC-SP-ISE and (B) HPLC method.

sensitivity and selectivity with time. Thus, we recommend switching to ISEs without an inner filling solution, such as the solid contact SP-ISE, which eliminates the inner filling solution with all of its expected problems as the release monitoring requires long durations of dipping the electrodes into the acidic solution at 37 °C for a long duration of time.³³ All solid contact SP-ISEs are internal solution-free. They employ a cheap highly conductive layer made from polymer or nanostructures, such as nanoparticles, as a solid contact beneath the ISE membrane layer. This allows for the availability of these SC-SP-ISEs commercially at very cheap prices, allowing for miniaturization of the ISEs for different *in vitro* and *in vivo* applications.³⁴

The conditioned SC-SP-ISE was dipped in conjunction with the reference electrode into the dissolution medium, and the emf was recorded in a continuous manner. Then, the percentage of release was determined using the related regression equation. The construction of the dissolution curve was traditionally performed *via* plotting the percentage of dissolution *versus* release time in minutes, Fig. 6.

3.3.2. Off-line HPLC method. To compare the results of the in-line potentiometric measurement, the tracking of the drug release was performed *via* the proposed HPLC chromatographic separation. The withdrawn samples were filtered and degassed, and then chromatographed using the above-mentioned conditions. This off-line quantitative determination was found to be time and chemical consuming as a result of the multiple sampling and different preparation steps, which led to a profile that contained gaps of time as shown in Fig. 6.

The dissolution testing showed a complete release of PIO, where more than 90% of PIO was released from the dosage form after 15 minutes. The results are consistent with the acceptance dissolution criteria, as stated by USP.³⁵ From Fig. 6, it can be easily concluded that SC-SP-ISE has equivalent performance to the traditional HPLC for tracking the PIO release from the pharmaceutical preparations.

Conclusion

It can be concluded from the results of the proposed method that the solid contact single carbon screen-printed electrodes (SC-SP-ISE) have shown higher performance over the liquid contact ion-selective electrodes (LC-ISE) for the real-time tracking of the release of PIO from its commercial tablets. This may be due to the elimination of the inner filling solution that is present in the LC-ISE, which is susceptible to evaporation during analysis or the undesirable fluxes through the membrane from the internal filling concentrated components to the outer surface of the membrane, causing deterioration of the analytical sensitivity and selectivity with time. The results obtained using the SC-SP-ISE showed a full profile of PIO release with high selectivity and accuracy in the presence of the expected interfering substances, such as the excipients and ALG.

In comparison to the developed HPLC method, the proposed method showed reliable performance with a very good result, contentious profiles of release, lower solvent consumption, lower generation of solvents, and environmentally friendly technique. Finally, using SP-ISEs could greatly open the door for simple performance of the forthcoming dissolution studies, delineating them as an outstanding alternative to other offline techniques commonly utilized for monitoring the dissolution profile.

Conflicts of interest

The authors declare no conflict of interest.

Acknowledgements

We would like to express our deep thanks to Prof. Dr Mohamed K. Abd El-Rahman, a Visiting Scholar at the Whitesides group, Harvard University, and a Professor of Analytical Chemistry, Faculty of Pharmacy, Cairo University, for his guidance and valuable discussions during replying to review comments on this work. Also, Dr Sherif M Eid would like to thank his supervisor, Prof. Dr M. K. Abd El-Rahman, for the patient guidance, discussions and advices he has provided throughout the time as his electrochemistry student.

References

- L. Mora, K. Chumbimuni-Torres, C. Clawson, L. Hernandez,
 L. Zhang and J. Wang, J. Controlled Release, 2009, 140, 69–73.
- 2 A. Ensafi, E. Heydari-Bafrooei and B. Rezaei, *Electroanalysis*, 2014, **26**, 776–785.
- 3 S. M. Eid, Talanta, 2020, 218, 121135.
- 4 S. Takano, S. Shiomoto, K. Inoue, K. Ino, H. Shiku and T. Matsue, *Anal. Chem.*, 2014, **86**, 4723–4728.
- 5 M. Sun, S. Kaplan, R. Gehringer, R. Limbocker and M. Johnson, *Anal. Chem.*, 2014, **86**, 4151–4156.
- 6 M. Antilen, F. Amiama, M. Otaiza, F. Armijo, M. Escudey, C. Pizarro and N. Arancibia-Miranda, *J. Nanopart. Res.*, 2015, 17, 212.
- 7 D. Araujo, A. Faria, R. Takeuchi and A. Santos, *Electroanalysis*, 2016, **28**, 596–600.
- 8 M. Li, Y.-T. Li, D.-W. Li and Y.-T. Long, *Anal. Chim. Acta*, 2012, 734, 31–44.
- 9 M. K. Abd El-Rahman, S. M. Eid, M. R. Elghobashy and K. M. Kelani, Sens. Actuators, B, 2019, 285, 216–223.
- 10 C. Zuliani and D. Diamond, Electrochim. Acta, 2012, 84, 29-34.
- 11 M. El-Rahman, H. Zaazaa, N. ElDin and A. Moustafa, ACS Sustainable Chem. Eng., 2016, 4, 3122–3132.
- 12 X. Wang, X. Li, Y. Bai and H. Liu, *Chem. Commun.*, 2015, **51**, 4615–4618.
- 13 K. Y. Chumbimuni-Torres, E. Bakker and J. Wang, *Electrochem. Commun.*, 2009, **11**, 1964–1967.
- 14 V. Shah, in *Pharmaceutical Dissolution Testing*, 2005, pp. 81–96.
- 15 W. Brown, in *Pharmaceutical Dissolution Testing*, 2005, pp. 69–80.
- 16 S. Eid, K. Kelani, O. Badran, M. Rezk and M. Elghobashy, *Anal. Chim. Acta*, 2020, **1117**, 60–73.
- 17 M. Petruševska, M. Horvat, L. Peternel and K. Kristan, *Drug Dev. Ind. Pharm.*, 2015, **42**, 1149–1157.

18 Z. Legnerova, Talanta, 2002, 58, 1151-1155.

Paper

- 19 K. Peeters, R. De Maesschalck, H. Bohets, K. Vanhoutte and L. Nagels, *Eur. J. Pharm. Sci.*, 2008, 34, 243–249.
- 20 M. K. A. El-Rahman, R. A. Sayed, M. S. El-Masry, W. S. Hassan and A. Shalaby, *J. Electrochem. Soc.*, 2018, 165, B143-B149.
- 21 E. S. Elzanfaly and A. S. Saad, *ACS Sustainable Chem. Eng.*, 2017, 5, 4381–4387.
- 22 M. R. Rezk, A. S. Fayed, H. M. Marzouk and S. S. Abbas, *J. Electrochem. Soc.*, 2017, **164**, H628–H634.
- 23 A. Kumar, Y. Zhang, D. Li and R. Compton, *Electrochem. Commun.*, 2020, **121**, 106867.
- 24 Y. Umezawa, K. Umezawa and H. Sato, *Pure Appl. Chem.*, 1995, **67**, 507–518.
- 25 W. E. Morf, *The Principles of Ion-Selective Electrodes and of Membrane Transport*, Elsevier, 2012.
- 26 R. D. Armstrong and M. Todd, J. Electroanal. Chem. Interfacial Electrochem., 1987, 237, 181–185.

- 27 D. Wang, J. Hu, Y. Li, M. Fu, D. Liu and Q. Chen, J. Membr. Sci., 2016, 501, 228–235.
- 28 I. A. Pechenkina and K. N. Mikhelson, *Russ. J. Electrochem.*, 2015, **51**, 93–102.
- 29 Z. Yan, Y. Lu and X. Li, Sens. Actuators, B, 2007, 122, 174-181.
- 30 Y. Umezawa, P. Bühlmann, K. Umezawa, K. Tohda and S. Amemiya, *Pure Appl. Chem.*, 2000, 72, 1851–2082.
- 31 I. H. T. Guideline, Q2 (R1), 2005, vol. 1, pp. 1-15.
- 32 M. Sohail and R. De Marco, in *Reference Module in Chemistry, Molecular Sciences and Chemical Engineering*, 2013.
- 33 J. Hu, A. Stein and P. Bühlmann, *TrAC, Trends Anal. Chem.*, 2016, 76, 102–114.
- 34 H. M. Mohamed, TrAC, Trends Anal. Chem., 2016, 82, 1-11.
- 35 J. L. Cohen, B. B. Hubert, L. J. Leeson, C. T. Rhodes, J. R. Robinson, T. J. Roseman and E. Shefter, *Pharm. Res.*, 1990, **07**, 983–987.