RSC Advances



PAPER

View Article Online
View Journal | View Issue



Cite this: RSC Adv., 2018, 8, 29104

Exploring the nature of the clopidogrel—bromocresol green interaction *via* spectrophotometric measurements and quantum chemical calculations†

Sabrein H. Mohamed, *\overline{O} *a Alyaa I. Magdy* and Ashour A. Ahmed *\overline{O} c

Clopidogrel is an oral, thienopyridine class antiplatelet agent used to inhibit blood clots in coronary arteries, peripheral vascular and cerebrovascular diseases. A spectrophotometric method was developed for clopidogrel bisulfate (CLOP·H₂SO₄) determination using bromocresol green (BCG) as an ion-pairing agent. To explore the binding nature of CLOP·H₂SO₄ with BCG at a molecular level, quantum chemical calculations have been performed. DFT based full geometry optimization has been carried out for BCG and clopidogrel in basic (CLOP) and protonated (CLOP+) forms as well as for BCG ion-pairs with CLOP and CLOP·H₂SO₄. The DFT calculations referred to the stability of the BCG-CLOP⁺ ion-pair and its spontaneous formation reaction from BCG and CLOP·H₂SO₄ compared to the BCG-CLOP-ion-pair. Furthermore, the UV-visible spectra and their corresponding excited states and electronic transitions for BCG, BCG-CLOP⁺ ion-pair, and BCG-CLOP ion-pair have been investigated. These spectra provided a molecular level understanding of the nature of the different intra-molecular and intermolecular electronic transitions in the BCG ion-pairs with CLOP⁺. Moreover, the quantitative analysis based on extracting a yellow-formed ion-pair into chloroform from aqueous medium was carried out. The ionpair exhibits an absorption maximum at 413 nm. The optimum conditions of the reactions were studied experimentally and optimized. The calibration graph shows that CLOP·H₂SO₄ can be determined up to 100.0 μg mL⁻¹ with detection limit (LOD) of 0.57 μg mL⁻¹ and quantification limit (LOQ) of 1.86 μg mL⁻¹. The low relative standard deviation values, 0.16-1.16, indicate good precision. The results were compared to other published data and were treated statistically using F and t-tests.

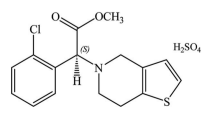
Received 17th June 2018 Accepted 8th August 2018

DOI: 10.1039/c8ra05187a

rsc.li/rsc-advances

1 Introduction

Clopidogrel bisulfate (CLOP· H_2SO_4), (2S)-2-(2-chlorophenyl)-2-(6,7-dihydro-4H-thieno[3,2-c]pyridine-5-yl)acetate, has a molecular weight of 419.9 g mol⁻¹, Scheme 1.¹ In medicinal chemistry, it is marketed globally with the primary goal of being able to reduce atherosclerotic events in patients with several comorbid conditions due to stroke, myocardial infarction and cardiovascular disease.² The recommended therapeutic dose of CLOP· H_2SO_4 is 75 mg per day and the initial pharmacodynamics activity can occur within 2 h after a single dose. CLOP· H_2SO_4 in combination with aspirin significantly reduced collagen-induced platelet aggregation.³



Scheme 1 Structural formula of clopidogrel bisulphate.

A review about the bioanalytical methods, pharmacokinetics and pharmacodynamics has been published regarding the update on the recent trends in drug interaction studies.⁴ The literature survey revealed that several analytical techniques have been reported for clopidogrel, where there is no previous study about its action with sulfonephthalein dyes. It was observed that there was a UV spectrophotometric method for its analysis in presence of its alkaline degredates⁵ and in its pharmaceutical formulation.⁶ Three visible spectrophotometric methods⁷ have been reported for its quantification in pure and pharmaceutical dosage forms based on the formation of ion-pair complexes of the drug with acidic dye Solochrome black T, oxidation of the

[&]quot;Chemistry Department, Faculty of Science, Cairo University, Giza, Egypt. E-mail: sabrein@sci.cu.edu.eg; sabrein_harbi@yahoo.com

^bChemistry Department, Faculty of Engineering, Madina High Institute for Engineering and Technology, Giza, Egypt

^{&#}x27;Institute of Physics, University of Rostock, D-18059 Rostock, Germany

[†] Electronic supplementary information (ESI) available. See DOI: 10.1039/c8ra05187a

Table 1 A comparison between the suggested and some other published spectrophotometric methods for the quantification of clopidogrel

Reagent	Linear range μg mL ⁻¹	$\begin{array}{c} LOD \ \mu g \\ mL^{-1} \end{array}$	Slope, $a \perp g^{-1} \text{ cm}^{-1}$	r^2	$\varepsilon \ \mathrm{L} \ \mathrm{mol}^{-1} \ \mathrm{cm}^{-1}$	RSD%	$\lambda_{ m max}$ nm	Ref.
BCG	Up to 100.0	0.57	1.00×10^{-2}	0.9946	5.67×10^{3}	0.31-0.96	415	C.S
Direct UV	40.0-70.0	0.40	1.00	1.0000	3.20×10^{2}	_	222	6
SBT	15.0-25.0	_	1.87×10^{-1}	0.9969	5.0781×10^4	0.69	510	7
(Fe(III)/PTL)	10.0-50.0	_	$7 imes 10^{-3}$	0.9980	3.873×10^{3}	0.66	515	7
$KMnO_4$	2.5-12.5	_	2.00×10^{-3}	0.9980	21.27×10^3	1.56	410	7

 $[^]a$ r^2 : correlation coefficient, ε : molar absorptivity, RSD%: relative standard deviation, C.S: current study, BCG: Bromocresol green, SBT: Solochrome black T, PTL: phenanthroline.

drug with phenanthroline and oxidation with potassium permanganate, Table 1.

Sulfonephthalein dyes are commonly used as indicators in acid–base titrations. They have been also used to build pH-sensors⁸ and for the spectrophotometric determination of different drugs.^{9,10} Correlation between the pK_a of these dyes with the nature of the substituent group was studied using the stability quotients from absorbance data.¹¹ The electron-donor group increases their pK_a values and the electron-attracting group decreases these values. Spectrophotometry has an inherent simplicity, a low cost and a wide availability in most quality control laboratories, so it considered as the most convenient analytical techniques.

Based on the wide use of sulfonephthalein dyes, the main goal of the present study is to perform density functional theory (DFT) based molecular modelling to investigate the binding nature of clopidogrel in basic (CLOP) and protonated (CLOP $^+$) forms with bromocresol green (BCG) at a molecular level. In addition, the use of BCG, Scheme 2, in the determination of the antiplatelet drug CLOP \cdot H $_2$ SO $_4$ as a way to get its concentration with a new simple, accurate and sensitive technique in its raw material and pharmaceutical formulations in comparison with the previously published literary.

2. Materials and methods

2.1. Apparatus

All the spectral measurements were carried out using a Jenway 6105 UV/Vis single beam spectrophotometer equipped with

Scheme 2 The structural formula of bromocresol green.

quartz cells of 1 cm optical path length. A Scientech SA 210 digital balance was used for weighing throughout the study.

2.2. Reagents and solutions

All chemicals and reagents used in this investigation were of the highest purity available, and bi-distilled water was used throughout the current study. Chloroform and absolute ethanol were obtained from Sigma-Aldrich (Germany). BCG was purchased from BDH Limited, Poole (England). CLOP·H₂SO₄ standard powder was obtained from Pfizer, Egypt, Plavix® (75.0 mg per tablet) was manufactured by Sanofi Aventis, France, and Borgavix® (75.0 mg per tablet) was obtained from Borg Pharmaceutical Industries. Standard stock solutions of 1.0 mmol L⁻¹ BCG was prepared by dissolving the exact weight in 10 mL of ethyl alcohol then completed to 50 mL with bidistilled water. CLOP·H₂SO₄ stock solution was prepared by dissolving 10.49 mg of the pure powder in 5.0 mL of ethyl alcohol and completed with bi-distilled water up to 25 mL to obtain a solution of 1.0 mmol L⁻¹.

2.3. Visible spectrophotometric method

The absorption spectra of 1.0 mmol L^{-1} BCG was measured in the range of 300–500 nm using 1.0 cm path length spectrophotometric cell. This has been done after extraction of BCG in 5.0 mL chloroform by shaking for 1.0 min into a 60.0 mL separating funnel. The same action was done for the formed ion-pair upon mixing 1.0 mL 1.0 mmol L^{-1} of CLOP· H_2SO_4 with 1.0 mL of 1.0 mmol L^{-1} of BCG. The absorption spectrum of the extract was measured also in the same range against a reagent blank. All measurements were carried out at room temperature (25 \pm 2 °C).

The stoichiometric ratios were determined utilizing Job's method of continuous variation. 12 Diverse volumes of 1.0 mmol L^{-1} CLOP· H_2SO_4 were treated with the corresponding complementary volume of 1.0 mmol L^{-1} BCG to give a total volume of 3.0 mL of both reagent and drug. Then the experiment was continued as mentioned above.

A more detailed examination for the ion-pair (1:1) and (1:2), drug to reagent, was continued by using Benesi–Hildebrand method.¹³ It was conducted by using variable concentrations of 0.2–1.2 of 0.5 mmol L⁻¹ BCG, to which 1.0 mL 5.0 mmol L⁻¹ CLOP·H₂SO₄ was added. Then the experiment was continued as mentioned above.

RSC Advances

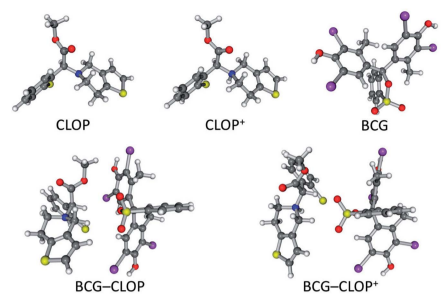


Fig. 1 Optimized geometries of CLOP, CLOP+, BCG, BCG-CLOP ion-pair, and BCG-CLOP+ ion-pair at the B3LYP/6-31+G(d,p) + Lanl2dz level of theory.

2.4. Procedure for tablets

10 Tablets (Plavix®, 75.0 mg per tablet or Borgavix®, 75.0 mg per tablet) were accurately weighed and ground using an agate mortar. The average weight equivalent to one tablet was calculated. An accurately weighed amount of the ground tablets powder equivalent to 75.0 mg CLOP·H2SO4 were dissolved in the least amount of ethyl alcohol. The resulting solution was filtered and the remained excipients were washed several times by using bi-distilled water to make sure that the entire active ingredient in the powder was completely dissolved, the volume was completed to 50 mL. Diverse volumes of the later solution were taken and treated by the same procedure for the pure one.

Quantum chemical calculations 2.5.

The interaction between BCG and clopidogrel was simulated through 1:1 ion-pair formation between them, i.e. 1 BCG + 1 clopidogrel → 1 BCG-clopidogrel ion-pair. Here, to study effect of the presence of bisulphate and its protonation to the clopidogrel N atom, clopidogrel was modelled in two different ways. Clopidogrel was introduced one time as a basic molecule (CLOP) and in another time as a protonated molecule (CLOP⁺, Fig. 1). This means that we have modelled two reactions, 1 BCG + 1 CLOP \rightarrow 1 BCG-CLOP ion-pair and 1 BCG + 1 CLOP⁺ \rightarrow 1 BCG-CLOP⁺ ion-pair. The initial geometries for each ion-pair were constructed by selecting the expected preferential binding situations between BCG and CLOP as well as between BCG and CLOP⁺. Full geometry optimization was performed for the initial ion-pair geometries as well as for the individual species (BCG, CLOP, and CLOP⁺) using the gradient minimization technique. Harmonic vibrational frequencies were calculated for all optimized geometries. These geometries are characterized by having zero gradient norms and positive harmonic vibrational frequencies. All calculations have been performed using the Gaussian09 program package.14 The

calculations were performed using density functional theory (DFT) with the Becke, three-parameter, Lee-Yang-Parr hybrid functional (B3LYP). 15,16 Here, the 6-31+G(d,p) basis set17 was used for H, C, N, O, and S atoms while the Lanl2dz basis set18,19 was used for Cl and Br atoms. In addition, the Grimme's D3 dispersion correction²⁰ has been applied.

For the ion-pair formation reaction, 1 BCG + 1 CLOP(CLOP⁺) \rightarrow 1 BCG-CLOP(CLOP⁺) ion-pair, the binding energy $(E_{\rm B})$ between BCG and CLOP or between BCG and CLOP+ is calculated as follows:

$$E_{\rm B} = E_{\rm BCG-CLOP(CLOP^+)ion\text{-}pair} - (E_{\rm BCG} + E_{\rm CLOP(CLOP^+)})$$
 (1)

where, $E_{\text{BCG-CLOP(CLOP}^+)\text{ion-pair}}$, E_{BCG} , and $E_{\text{CLOP(CLOP}^+)}$, are the electronic energies of the BCG-CLOP ion-pair or BCG-CLOP ion-pair, BCG, and CLOP or CLOP⁺, respectively. Here, the effect of the basis set superposition error (BSSE) for calculating the binding energy has been corrected using the counterpoise scheme.21 Similarly, the corresponding binding free energies $(\Delta G_{\rm B})$ were calculated by including the zero point energy and thermal correction to the Gibbs free energy. Moreover, UVvisible spectra (twelve excited states) for BCG and its ion-pairs with CLOP and CLOP+ have been calculated using linear response time-dependent DFT for the corresponding optimized geometries. In addition, the solvent effect on these UV-visible spectra has been involved by introducing chloroform as a solvent through an implicit treatment. This has been done using the conductor-like polarizable continuum model (CPCM).22

Results and discussion 3.

Quantum chemical calculations

The optimized geometries for CLOP, CLOP+, BCG, BCG-CLOP ion-pair, and BCG-CLOP+ ion-pair are gathered in Fig. 1. Based Paper

on the spatial configuration for the optimized structures, the root mean square deviation (RMSD) of the xyz coordinates between the BCG-CLOP- and BCG-CLOP+ ion-pairs is larger than that between CLOP and CLOP⁺. This means that the basic and protonated clopidogrel molecular configurations are closer to each other compared to the molecular configurations of the basic and protonated clopidogrel ion-pairs with BCG. Here, the main kind of the molecular interaction in both BCG-CLOP and BCG-CLOP⁺ ion-pairs is resulting from the dispersion interaction. Both ion-pairs have not shown any covalent bond or Hbond. Further, the BCG-CLOP ion-pair has exhibited electrostatic interaction between the positively charged CLOP⁺ N atom and the BCG electron density. This yields a stronger interaction between BCG and CLOP+ than between BCG and CLOP. In addition, the calculated binding energy for the BCG-CLOP⁺ ionpair, -19.5 kcal mol⁻¹, is almost twice of that for the BCG-CLOP ion-pair $(-9.9 \text{ kcal mol}^{-1}, \text{ Table 2})$. This indicates to the substantial role of presence the sulfuric acid in the ion-pair formation process between BCG and CLOP·H2SO4. Here, the sulfuric acid strengthens the interaction between BCG and CLOP+ in the BCG-CLOP+ ion-pair two times compared to the case of absence of the sulfuric acid. Moreover, the calculated binding free energies between BCG and clopidogrel for the BCG-CLOP and BCG-CLOP ion-pairs are +1.9 and $-8.9 \text{ kcal mol}^{-1}$, respectively. These values indicate again that the BCG-CLOP⁺ ion-pair is stronger and more stable than the BCG-CLOP ion-pair. Furthermore, one can explore from the binding free energy values that the BCG-CLOP⁺ ion-pair is thermally stable and can be formed spontaneously. In contrast,

Table 2 Calculated binding energies, enthalpy, and free energies between BCG and CLOP as well as CLOP $^+$ in the BCG $^-$ CLOP, and BCG $^-$ CLOP $^+$ ion-pairs at the B3LYP/6-31+G(d,p) + Lanl2dz level of theory

the BCG-CLOP ion-pair is not thermally stable ion-pair and

	$E_{ m B}$			
	(kcal mol ⁻¹)	$E_{ m B}^{ m ZPE+thermal}$	$\Delta H_{ m B}$	$\Delta G_{ m B}$
BCG-CLOP BCG-CLOP ⁺	-9.90 -19.49	-8.69 -17.74	-9.28 -18.33	+1.88 -8.90

needs activation to be formed. This eventually indicates that the formation of the ion-pair between BCG and CLOP⁺ will be carried out in two steps. First, the protonated clopidogrel (CLOP⁺) will be formed by proton transfer process from the sulfuric acid to clopidogrel. Consequently, a spontaneous interaction process will take place between BCG and the protonated clopidogrel (CLOP⁺).

Next, the UV-visible spectra for BCG and its ion-pairs with CLOP and CLOP⁺ were calculated theoretically (Fig. 2) for further understanding for the nature of the intra-molecular and inter-molecular interaction in BCG, BCG–CLOP ion-pair, and BCG–CLOP⁺ ion-pair. Regardless the shift in $\lambda_{\rm max}$ between the experimental and theoretical UV-visible spectra, Fig. 2 provides two evidence points for the stronger interaction between BCG and CLOP⁺ compared to that between BCG and CLOP. First, the intensity or absorbance at $\lambda_{\rm max}$ for the BCG–CLOP ion-pair is almost 63 times of that for the BCG–CLOP ion-pair. Second, the shift in $\lambda_{\rm max}$ between the BCG and BCG–CLOP spectra is larger than that between the BCG and BCG–CLOP⁺ spectra. For more details, the possible excited states and the corresponding electronic transitions for BCG and its ion-pairs with CLOP and CLOP⁺ will be discussed briefly below.

For BCG, the calculated UV-visible spectrum showed excited states at wavelength range from 256.0 to 287.2 nm. The intensity of these states has been increased in the order 277.6 < 256.0 <269.1 < 269.8 < 257.9 < 287.2 < 261.4 < 262.3 < 285.1 < 270.7 < 280.0 < 265.5 nm. The most intense excited state at 265.5 nm consists of several electronic transitions involving HOMO $-3 \rightarrow$ LUMO+2 (16.3%), HOMO-1 \rightarrow LUMO+1 (14.4%), HOMO-3 \rightarrow LUMO+1 (14.2%), HOMO-1 \rightarrow LUMO+2 (13.7%), HOMO-2 \rightarrow LUMO+2 (6.8%), HOMO-3 \rightarrow LUMO (6.5%), and HOMO-1 \rightarrow LUMO+5 (6.2%). Focusing on the frontier molecular orbitals in Fig. 3, this explains that the most intense excited state involved the $\pi \to \pi^*$ and $\pi \to \sigma^*$ electronic transitions. These electronic transitions take place from the BCG benzene rings "A" and "B" to benzene ring "C" as well as to the Br atoms involved in the benzene rings "A" and "B". The excited state at 280.0 nm occurred due to combination of HOMO \rightarrow LUMO+1 (48.5%), HOMO \rightarrow LUMO+5 (12.1%), HOMO \rightarrow LUMO+2 (8.4%), and $HOMO \rightarrow LUMO+4$ (7.6%) electronic transitions. Like the previous excited state, this state involved $\pi \to \pi^*$ and $\pi \to \sigma^*$

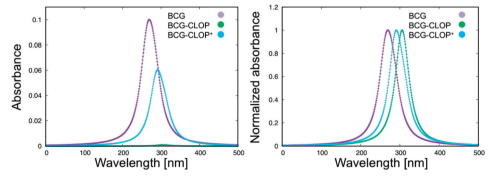


Fig. 2 Calculated UV-visible spectra for BCG and its ion-pairs with CLOP and CLOP $^+$ at the B3LYP/6-31+G(d,p) + Lanl2dz level of theory. In the right panel, the absorbance is normalized for every separate spectrum.

RSC Advances Paper

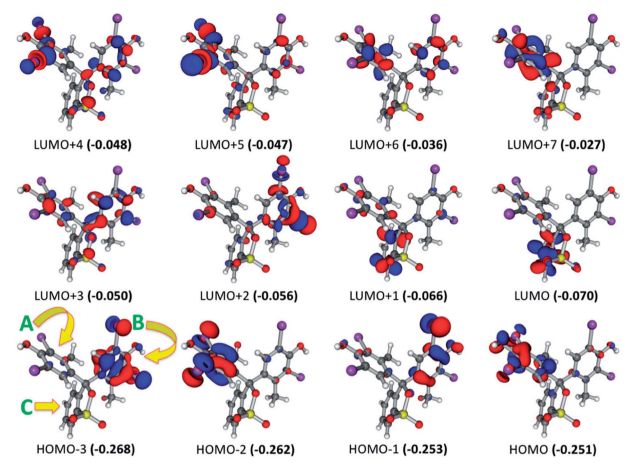


Fig. 3 Frontier molecular orbitals for the optimized geometry of BCG at the B3LYP/6-31+G(d,p) + Lanl2dz level of theory. Values in parentheses are the electronic energies in a.u. of the corresponding frontier molecular orbitals. For simplicity, we have referred the symbols "A", "B", and "C" to the three benzene rings in BCG (see HOMO-3). Band gap = 0.181 a.u.

electronic transitions as well. Here, these electronic transitions happen from the benzene ring "A" to the benzene rings "B" and "C" and also to the C-Br bonds in the benzene rings "A" and "B". At 270.7 nm, combination of the electronic transitions $HOMO \rightarrow LUMO+4 (15.8\%), HOMO \rightarrow LUMO+1 (15.2\%),$ $HOMO-2 \rightarrow LUMO (14.1\%), HOMO \rightarrow LUMO+3 (13.1\%),$ $HOMO \rightarrow LUMO+2$ (6.0%), and $HOMO \rightarrow LUMO$ (5.0%) has been observed. This means that this excitation state involved π $\rightarrow \pi^*$ and $\pi \rightarrow \sigma^*$ electronic transitions from the benzene ring "A" to the benzene rings "A", "B" and "C" and to the C-Br bonds. Furthermore, the electronic transitions HOMO → LUMO (69.3%), HOMO-1 \rightarrow LUMO (8.5%), and HOMO-1 \rightarrow LUMO+1 (5.3%) have been observed at 285.1 nm that are related to $\pi \to \pi^*$ transitions from the benzene rings "A" and "B" to the benzene ring "C". While $\pi \to \pi^*$ electronic transitions have been observed at 262.3 nm from the benzene ring "A" to the benzene ring "C" through the HOMO $-2 \rightarrow LUMO$ (47.6%) and $HOMO-2 \rightarrow LUMO+1$ (39.9%) transitions. For the rest of the excited states, only the most contributed electronic transition for every excited state will be mentioned briefly. Here, the electronic transitions HOMO → LUMO+2 (256.0 nm), $HOMO-1 \rightarrow LUMO+3$ (261.4 nm), $HOMO-1 \rightarrow LUMO+1$ (269.8 and 287.2 nm), HOMO-1 \rightarrow LUMO (277.6 nm),

HOMO–2 \rightarrow LUMO+5 (257.9 nm), and HOMO–2 \rightarrow LUMO (269.1 nm) have been observed at the corresponding wavelength (given in parentheses). These are related to electronic transitions from π of the benzene rings "A" and "B" to π^* of the benzene rings "A", "B", and "C" and σ^* of C–Br bonds. The last exited sates can summarize all the possible electronic transitions that take place in BCG.

For the BCG-CLOP⁺ ion-pair, the intensity of the most intense excited states has the order 291.7 > 288.0 > 309.7 > 284.8 > 305.8 > 285.5 nm. For the excited state at 291.7 nm, the main electronic transitions are HOMO → LUMO+3 (72.8%) and $HOMO-2 \rightarrow LUMO+1$ (18.6%) transitions. At 288.0 nm, the $HOMO-2 \rightarrow LUMO (44.2\%), HOMO-2 \rightarrow LUMO+1 (33.6\%)$ electronic excitations have been observed while at 309.7 nm the $HOMO \rightarrow LUMO+1$ (68%) and $HOMO \rightarrow LUMO$ (30%) transitions were the most abundant electronic transitions. For the excited state at 284.8 nm, the HOMO $-1 \rightarrow LUMO+3$ electronic transition was the main transition. Eventually, the electronic transitions HOMO → LUMO, HOMO → LUMO+1, HOMO-1 \rightarrow LUMO, HOMO-1 \rightarrow LUMO+3, HOMO-2 \rightarrow LUMO, and HOMO-2 → LUMO+1 have been observed at 305.8 and 285.5 nm. All these intense transitions can be simply classified into intra-molecular as well as intermolecular $\pi \to \pi^*$

Paper

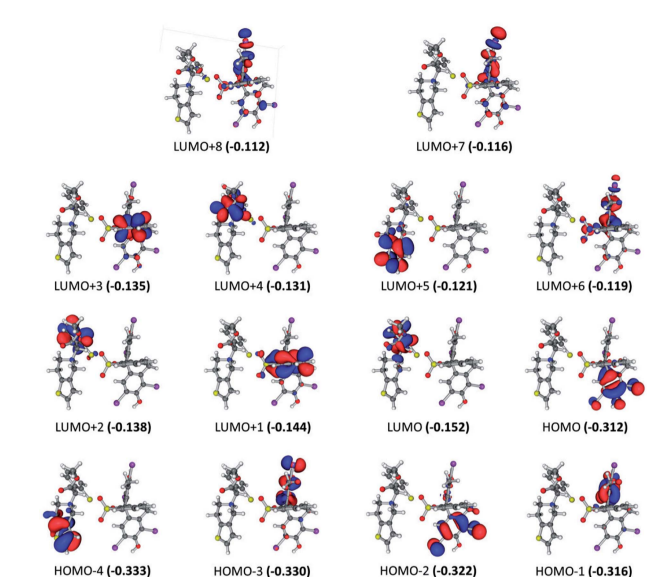


Fig. 4 Frontier molecular orbitals for the optimized geometry of $BCG-CLOP^+$ ion-pair at the B3LYP/6-31+G(d,p) + Lanl2dz level of theory. Values in parentheses are the electronic energies in a.u. of the corresponding frontier molecular orbitals. Band gap = 0.160 a.u.

electronic transitions (Fig. 4). Specifically, the intra-molecular transitions take place from the BCG benzene rings "A" and "B" to the BCG benzene ring "C". On the other hand, the intermolecular electronic transitions take place from the BCG benzene rings "A" and "B" to the clopidogrel benzene ring. The latter transitions can strengthen the interaction between BCG and clopidogrel in the BCG-CLOP⁺ ion-pair. Moreover, low intense excited states are observed at 303.6, 297.4, 279.6, 279.3, and 277.7 nm due to the electronic transitions HOMO \rightarrow LUMO, HOMO \rightarrow LUMO+2, HOMO-1 \rightarrow LUMO, HOMO-1 \rightarrow LUMO+1, HOMO-1 \rightarrow LUMO+2, HOMO-3 \rightarrow LUMO, and $HOMO-4 \rightarrow LUMO$ (Fig. 4). These excited states involved mainly intermolecular $\pi \to \pi^*$ electronic transitions from the BCG benzene rings "A" and "B" to the clopidogrel benzene ring in addition to one intramolecular $\pi \to \pi^*$ electronic transition from the BCG benzene ring "A" to the BCG benzene ring "C". Furthermore, the HOMO-4 \rightarrow LUMO transition is related to

intramolecular $\pi \to \pi^*$ electronic transition in clopidogrel from the thiophene ring to benzene one.

For the BCG–CLOP ion-pair, very low intense excited states are observed at 360.8, 355.1, 347.4, 341.3, 322.5, and 304.0 nm. These excited states involved the electronic transitions from HOMO to LUMO, LUMO+1, LUMO+2, and LUMO+3; from HOMO–1 to LUMO, LUMO+1, and LUMO+2; and from HOMO–2 to LUMO (Fig. 5). Excited states of intermediate intensity have been observed at 298.7, 313.3, 313.8, and 304.5 nm. Mainly these excited states have been arisen due to transitions from HOMO to LUMO+2, LUMO+3, LUMO+4, and LUMO+5; and from HOMO–1 to LUMO+2, LUMO+3, and LUMO+4. The most intense excited states, at 303.6 and 306.22 nm, have been explored via the electronic transitions HOMO \rightarrow LUMO+4, HOMO \rightarrow LUMO+7; HOMO–1 \rightarrow LUMO+4, and HOMO–1 \rightarrow LUMO+7. This means that the most abundant excited states included intramolecular and

This article is licensed under a Creative Commons Attribution-NonCommercial 3.0 Unported Licence.

Open Access Article. Published on 15 August 2018. Downloaded on 12/6/2025 5:52:48 AM

RSC Advances Paper

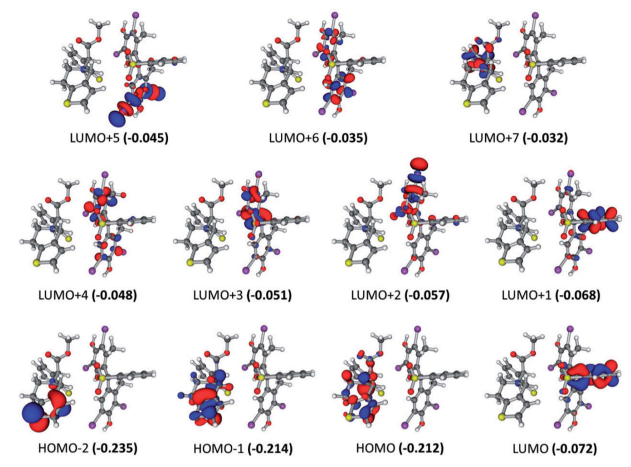


Fig. 5 Frontier molecular orbitals for the optimized geometry of BCG-CLOP ion-pair at the B3LYP/6-31+G(d,p) + Lanl2dz level of theory. Values in parentheses are the electronic energies in a.u. of the corresponding frontier molecular orbitals. Band gap = 0.140 a.u.

intermolecular $\pi\to\pi^*$ electronic transition from the BCG benzene rings "A" and "B" to the BCG benzene ring "A" and the clopidogrel benzene ring. In addition, a $\pi\to\sigma^*$ intra-molecular electronic transition has been observed to the C–Br bonds. Eventually, the UV-visible electronic transitions show that this BCG–CLOP ion-pair is weaker than the BCG–CLOP $^+$ ion-pair although it shows intermolecular electronic transitions from BCG to clopidogrel. This is due to the intensity of the electronic transitions in the BCG–CLOP ion-pair is very small compared to that in the BCG–CLOP $^+$ ion-pair. This conclusion comes in a good agreement with the calculated binding (free) energies for both ion-pairs.

Finally, the effect of chloroform as a solvent on the calculated UV-visible spectra for BCG and its ion-pairs with CLOP and CLOP $^+$ (see Fig. 2) has been introduced into Fig. 6. Here, the differences and similarities between the UV-visible spectra in vacuum and in chloroform will be discussed briefly. All the calculated UV-visible spectra in chloroform shifted to shorter wavelength compared to the spectra in the vacuum case. This is due to lower band gaps in vacuum (band gap: 0.181 a.u. for BCG, 0.160 for BCG–CLOP $^+$ ion-pair, 0.140 for BCG–CLOP ion-pair) than in chloroform (band gap: 0.186 for BCG, 0.181 for BCG–CLOP $^+$ ion-pair, 0.159 for BCG–CLOP ion-pair). Moreover, the shift in $\lambda_{\rm max}$ between BCG and BCG–CLOP $^+$ ion-pair is

smaller in chloroform than in vacuum. In addition, higher intensities have been observed for all spectra in chloroform compared to the vacuum cases. Regardless these differences, the order of intensities of $\lambda_{\rm max}$ and the order of the shift in $\lambda_{\rm max}$ between the spectra calculated in chloroform are the same as in

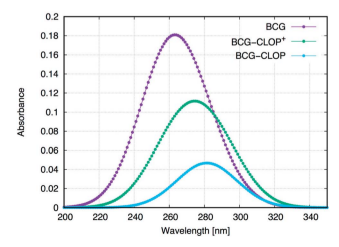


Fig. 6 Calculated UV-visible spectra for BCG and its ion-pairs with CLOP and $CLOP^+$ at the B3LYP/6-31+G(d,p) + Lanl2dz level of theory and using CPCM.

the vacuum cases. Mainly, the shift in λ_{max} between BCG–CLOP and BCG–CLOP⁺ ion-pairs is similar for the chloroform and vacuum cases. Moreover, the possible excited states and the corresponding electronic transitions are almost similar for both vacuum and chloroform cases (for more details see Fig. S1–S3 and Tables S1–S3 in the ESI†).

3.2. Experimental part

Having a molecular level investigation for the interaction between the clopidogrel drug in its basic (CLOP) and acidic (CLOP⁺) form with BCG in vacuum as well as in chloroform, this motivates us to introduce a valid and precise detection method to quantify clopidogrel. Here, several extractive spectrophotometric methods have been published for the basic nitrogenous compounds, in which the acidic dyes form ion-pairs that can be extracted from aqueous solution to organic phase.²³ Hence BCG as a selective reagent can be used for the determination of CLOP·H₂SO₄. After selection of maximum wavelength 413 nm, Fig. 7, some variables in the reaction conditions were studied and the influence of these variables on the reaction was tested as reagent concentration, effect of time, *etc.*

3.2.1. Optimization of the analytical procedures. The influence of BCG concentration on the intensity of the developed colour was studied. Different volumes (1.0-5.0) mL 1.0 mmol L^{-1} of BCG were added to 1.0 mL 1.0 mmol L^{-1} CLOP· H_2SO_4 . The data showed that 3 mL 1.0 mmol L^{-1} of BCG was sufficient to produce a maximum and reproducible colour.

The effect of time on the formation of the reaction product at $(25\pm2~^\circ\text{C})$ was investigated by allowing the reaction to proceed at different times. The results revealed that the reaction was completed within 2 min. The absorbance remains stable for 24 h. This allowed the processing of large number of samples. Moreover, this increases the reliability of the methods as well as made it applicable for large number of samples.

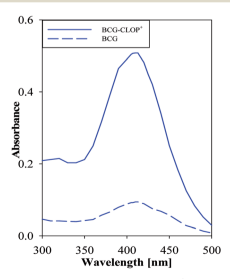


Fig. 7 Absorption spectrum of 1.0 mmol L^{-1} CLOP·H₂SO₄ with 1.0 mmol L^{-1} BCG against a blank and 1.0 mmol L^{-1} BCG against chloroform as a blank.

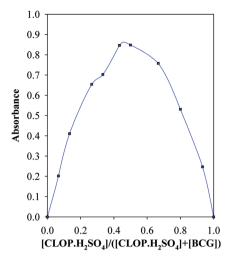


Fig. 8 Job's method of continuous variation for BCG-CLOP⁺ ion pair.

Under the optimum conditions, the stoichiometry of the reaction between $CLOP \cdot H_2SO_4$ and BCG was investigated by Job's method. The bell shape of Job's plot indicated that ion-pair formation between $CLOP \cdot H_2SO_4$ and BCG followed 1:1 and 1:2 reaction stoichiometries, Fig. 8.

3.2.2. The association and stability constants of the BCG–CLOP⁺ ion pair. The stability constant of the ion-pair (K_f) was determined by substituting the data of continuous variation in the following equation:²⁴

$$K_{\rm f} = \left(\frac{A}{A_{\rm m}}\right) / \left(\frac{1 - A}{A_{\rm m}}\right)^{n+1} C^n n^n \tag{2}$$

where, A and $A_{\rm m}$: are the observed absorbance and the absorbance maximum value when all the present drug is associated, respectively. C: is the molar concentration of the drug at the

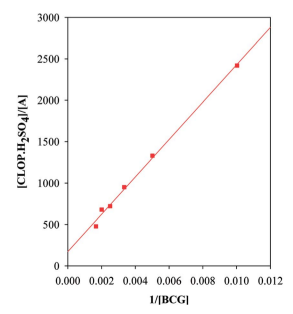


Fig. 9 Benesi-Hildebrand plots for BCG-CLOP+ ion-pair.

RSC Advances

Table 3 Analytical parameters for the determination of $CLOP \cdot H_2SO_4$ using BCG^a

Parameter	Value
λ_{\max} (nm)	413
Beer's law (μg mL ⁻¹)	Up to 100.0
Molar absorptivity (L mol ⁻¹ cm ⁻¹)	5.67×10^{3}
Sandell sensitivity (µg cm ⁻²)	8.06×10^{-2}
Ringbom range ($\mu g \text{ mL}^{-1}$)	17.37-81.28
Limit of detection (LOD) ($\mu g \text{ mL}^{-1}$)	0.57
Limit of quantification (LOQ) (µg mL ⁻¹)	1.86
Slope $(b)^*$ (mL μg^{-1} cm ⁻¹)	0.01
Intercept (a)*	0.06
Correlation coefficient (r^2)	0.9946
Relative standard deviation (RSD) % $(n = 3)$	0.31-0.96
K_C^{AD} (L mol ⁻¹)	5.34×10^{3}
$\varepsilon_{\lambda}^{AD}$ (L mol ⁻¹ cm ⁻¹)	2.43×10^{3}
$K_{\rm f}$ $(n=1)$	1.65×10^{5}
$K_{\rm f}(n=2)$	7.56×10^{18}

 a $K_C^{\rm AD}$ is the association constant calculated applying Benesi–Hildebrand method. $K_{\rm f}$ is the formation constants calculated from the continuous variation data.

maximum absorbance and n is the stoichiometry in which the reagent associates with the drug.

The association constant of 1:1 and 1:2 ion-pairs was determined using Benesi-Hildebrand method applying the following equation.¹⁰

$$\frac{[A_o]}{[A_\lambda^{\rm AD}]} = \frac{1}{\varepsilon^{\rm AD}} + \left(\frac{1}{K_C^{\rm AD}}\varepsilon_\lambda^{\rm AD}\right) \frac{1}{[D_o]} \tag{3}$$

where, $A_{\rm o}$ and $D_{\rm o}$ are the total concentration of reagent and drug, respectively, $A_{\lambda}^{\rm AD}$ and $\varepsilon^{\rm AD}$ are the absorbance and the molar absorptivity of the ion-pair at 413 nm, and $K_C^{\rm AD}$ is the association constant of the ion-pair. By plotting the values of $[A_{\rm o}]/[A_{\lambda}^{\rm AD}]$ versus $1/[D_{\rm o}]$, a straight line was obtained with a slope equals $(1/K_C^{\rm AD}\varepsilon_{\lambda}^{\rm AD})$, and intercept of $(1/\varepsilon^{\rm AD})$, Fig. 9. However, it

should be noted that ε^{AD} , which is the molar absorptivity of the ion-pair itself should not be confused with any stoichiometric values calculated regarding the amount of any analyte being determined. The latter is best described, as Beer's value while the former is Benesi–Hildebrand value. The association constants and the molar absorptivity are listed in Table 3.

3.2.3. Method validation, linearity. Under the above described experimental conditions, standard calibration curves for $CLOP \cdot H_2SO_4$ with BCG was constructed by plotting absorbance *versus* concentration, Fig. 10.

Conformity with calibration curve²⁵ and Ringbom are evident in the concentration ranges up to 100.0 and 17.37–81.28 $\mu g \ mL^{-1}$, respectively. The high molar absorptivity (5.67 \times 10³ L $mol^{-1} \ cm^{-1}$) and the low Sandell sensitivity values (8.06 \times 10⁻² $\mu g \ cm^{-2}$) reflect the good and high sensitivity of this method. The linearity of the calibration curve was approved by the high correlation coefficient (r^2), 0.9946, and the small value of y-intercept (0.06) of the regression equation, Table 3.

3.2.4. Sensitivity. The limit of detection (LOD) and quantitation (LOQ) for the suggested method were calculated using the following equations:

$$LOD = 3s/k, LOQ = 10s/k \tag{4}$$

where, s is the standard deviation of the blank and k is the sensitivity (the slope of the calibration graph). The method LOD and LOQ are 0.57 and 1.86 μ g mL⁻¹, respectively, Table 3.

3.2.5. Precision and accuracy, robustness and ruggedness. The reaction of the nitrogenous drug $CLOP \cdot H_2SO_4$ and the acid dye BCG is selective and sensitive. The precision of this method was studied by four replicate measurements of three different concentrations from $CLOP \cdot H_2SO_4$ solution within the calibration curve limits. The later was carried out by the study of the relative standard deviation (% RSD). The percentage error was also studied to show the method accuracy. Results in Table 4 show that this new method is repeatable.

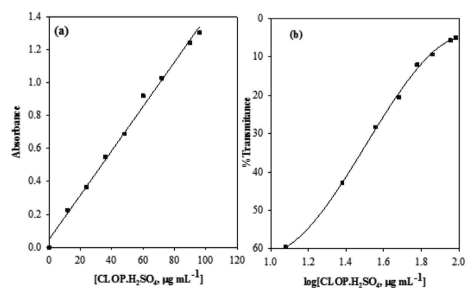


Fig. 10 Calibration curve (a) and Ringbom plot (b) for the determination of CLOP·H₂SO₄ with BCG.

Table 4 Evaluation of precision of the proposed methods on clopidogrel bisulfate pure

Taken (μg mL ⁻¹)	^a Recovery% ± SD	RSD%	t-Test	F-test
23.99	100.23 ± 0.31	0.31	0.94	2.31
35.99	101.00 ± 0.97	0.96	4.89	1.58
59.99	99.95 ± 0.91	0.91	0.31	1.27

Table 5 Analyses of clopidogrel bisulfate in its pharmaceutical formulations

Taken ($\mu g \; mL^{-1}$)	a Recovery% \pm SD	RSD%	t-Test	F-test			
Plavix® tablets (75.0 mg CLOP·H ₂ SO ₄ /tablet)							
23.99	98.00 ± 0.47	0.48	4.90	2.53			
35.99	100.47 ± 0.51	0.51	0.47	2.36			
59.99	99.89 ± 0.56	0.56	0.39	2.13			
Borgavix® tablets (75.0 mg CLOP·H ₂ SO ₄ /tablet)							
23.99	98.57 ± 0.76	0.77	2.65	1.55			
35.99	100.13 ± 0.57	0.57	0.49	1.16			
59.99	100.42 ± 0.69	0.69	0.08	1.42			

^a Average of four replicates.

The robustness of the method was evaluated by making small changes in some parameters as, the volume of dye, volume of extracting solvent and wavelength range. The effects of these changes were studied on the absorbance of the ion-pair. The changes had negligible influence on the results as revealed by small intermediate precision values expressed as % RSD (0.29–1.41%). Method ruggedness was demonstrated by carrying the analysis by two analysts, and also by a single analyst performing analysis using three different cuvettes. Intermediate precision values in both instances were in the range 0.58–1.63 indicating acceptable ruggedness.

3.2.6. Pharmaceutical analyses. It is evident from the above-mentioned results that the proposed methods gave satisfactory results with $CLOP \cdot H_2SO_4$ in pure solution. Thus, its pharmaceutical dosage forms (Plavix® and Borgavix® tablets) were subjected to the analysis of its contents by proposed method which is based on direct measurement of the absorbance of the formed ion-pair. The recovery percentages are within the range 98.0 to 101.0%, Table 5. The proposed methods for the determination of all pharmaceutical formulations were carried out on the same batch of samples. The results were statistically compared with those of the reference one²⁶ by applying the students *t*-test for accuracy and *F*-test for precision, Table 5.

4. Conclusions

Spectrophotometric method has advantageous over many of the reported methods due to their sensitivity, high percentage of recovery, wide application range, low relative standard deviation, and they do not need expensive sophisticated apparatus. Furthermore, all the analytical reagents are inexpensive, have excellent shelf life, and are available in many analytical laboratories. Therefore, the methods are practical and valuable for routine application in quality control laboratories for analysis of clopidogrel bisulfate. Moreover, investigation of the binding nature between it and BCG has been explored at molecular level. It was found that, BCG binds to the protonated form of clopidogrel (CLOP⁺) stronger and more spontaneously than to the corresponding neutral form (CLOP). Further, the UV-visible spectra for BCG, BCG–CLOP⁺ ion-pair, and BCG–CLOP ion-pair have been calculated. These spectra showed different intramolecular and intermolecular $\pi \to \pi^*$ and $\pi \to \sigma^*$ electronic transitions in the BCG ion-pairs with clopidogrel bisulfate.

Conflicts of interest

There are no conflicts to declare.

References

- 1 A. C. Moffat, M. D. Osselton and B. Widdop, *Clarke's Analysis of Drugs and Poisons*, Pharmaceutical Press, London, UK, 4th edn, 2011.
- 2 D. Taubert, A. Kastrati, S. Harlfinger, O. Gorchakoca, A. Lazar, N. Von Beckerath, A. Schomig and E. Schomig, *Thromb. Haemostasis*, 2004, 92, 311–316.
- 3 G. L. Plosker and K. A. Lyseng-Williamson, *Drugs*, 2007, **67**(4), 613–646.
- 4 R. Mullangi and N. R. Srinivas, *Biomed. Chromatogr.*, 2009, 23, 26–41.
- H. E. Zaazaa, S. S. Abbas, M. Abdelkawy and M. M. Abdelrahman, *Talanta*, 2009, 78, 874–884.
- 6 P. B. Cholke, R. Ahmed, S. Z. Chemate and K. R. Jadhav, Arch. Appl. Sci. Res., 2012, 4(1), 59–64.
- 7 B. Anupama, V. Jagathi, A. Aparna, M. Madhubabu and V. L. Annapurna, *Int. J. Pharma Bio Sci.*, 2011, 2, 105–108.
- 8 K. Vytras and J. Vytrasova, Chem. Zvesti, 1974, 28(6), 779-788.
- 9 K. Basavaiah, S. A. M. Abdulrahman and K. B. Vinay, *J. Food Drug Anal.*, 2009, 17(6), 434–442.
- 10 N. Rahman, N. A. Khan and S. N. H. Azmi, *Il Farmaco*, 2004, **59**(1), 47–54.
- 11 R. Casula, G. Crisponi, F. Cristiani and V. Nurchi, *Talanta*, 1993, **40**(12), 1781–1788.
- 12 P. Job, Ann. Chem., 1936, 16, 97.
- 13 H. A. Benesi and J. H. Hildebrand, *J. Am. Chem. Soc.*, 1949, 71(8), 2703–2707.
- 14 M. J. Frisch, G. W. Trucks, H. B. Schlegel, G. E. Scuseria, M. A. Robb, J. R. Cheeseman, G. Scalmani; V. Barone; B. Mennucci; G. A. Petersson; H. Nakatsuji; M. Caricato; X. Li; H. P. Hratchian, A. F. Izmaylov, J. Bloino, G. Zheng, J. L. Sonnenberg, M. Hada, M. Ehara, K. Toyota, R. Fukuda, J. Hasegawa, M. Ishida, T. Nakajima, Y. Honda, O. Kitao, H. Nakai, T. Vreven, J. A. Montgomery Jr; J. E. Peralta; F. Ogliaro; M. Bearpark; J. J. Heyd; E. Brothers; K. N. Kudin; V. N. Staroverov, R. Kobayashi,

- J. Normand, K. Raghavachari, A. Rendell, J. C. Burant, S. S. Iyengar, J. Tomasi, M. Cossi, N. Rega, J. M. Millam, M. Klene, J. E. Knox, J. B. Cross, V. Bakken, C. Adamo, J. Jaramillo, R. Gomperts, R. E. Stratmann, O. Yazyev, A. J. Austin, R. Cammi, C. Pomelli, J. W. Ochterski, R. L. Martin, K. Morokuma, V. G. Zakrzewski, G. A. Voth, P. Salvador, J. J. Dannenberg, S. Dapprich, A. D. Daniels, O. Farkas, J. B. Foresman, J. V. Ortiz, J. Cioslowski and D. J. Fox, *Gaussian 09, Revision D.01*, Gaussian, Inc., Wallingford CT, 2013.
- 15 A. D. Becke, Phys. Rev. A: At., Mol., Opt. Phys., 1988, 38, 3098-3100.
- 16 C. Lee, W. Yang and R. G. Parr, Phys. Rev. B: Condens. Matter Mater. Phys., 1988, 37, 785–789.
- 17 W. J. Hehre, R. Ditchfield and J. A. Pople, *J. Chem. Phys.*, 1972, **56**, 2257–2261.

- 18 T. H. Dunning Jr and P. J. Hay, in *Modern Theoretical Chemistry*, ed. H. F.Schaefer III, Plenum, New York, 1977, vol. 3, pp. 1–28.
- 19 P. J. Hay and W. R. Wadt, J. Chem. Phys., 1985, 82, 270-283.
- 20 S. Grimme, S. Ehrlich and L. Goerigk, J. Comput. Chem., 2011, 32, 1456–1465.
- 21 H. B. Jansen and P. Ros, Chem. Phys. Lett., 1969, 3, 140-143.
- 22 M. Cossi, N. Rega, G. Scalmani and V. Barone, *J. Comput. Chem.*, 2003, 24, 669–681.
- 23 I. Suslu and A. Tamer, *J. Pharm. Biomed. Anal.*, 2002, **29**, 545–554.
- 24 J. Inczédy, *Analytical Application of Ion-pair Equilibria*, Budapest, John Wiley & Sons Inc, 1976.
- 25 J. C. Miller and J. N. Miller, *Statistics for Analytical Chemistry*, Horwood, Chichester: UK, 3rd edn, 1993.
- 26 S. Gurav, R. Tembare, V. Salunkhe, Devprakash and G. P. Senthilkumar, *Am. J. PharmTech Res.*, 2011, 1(4), 258–263.