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# Chemical reactivity of 2-methylchromone with some nitrogen-based nucleophiles: reactions, DFT analysis, anticancer activity and molecular docking studies

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The current study investigates the reactivity of 2-methylchromone toward nitrogen-containing heterocyclic amines, with the aim of assessing how substitution pattern and nucleophile type influence chromone ring opening and subsequent heterocycle formation. Reaction of 2-methylchromone (**1**) with selected heterocyclic amines afforded a series of heterocycle-linked enaminones **5**, **8** and **9**, which were found to preferentially adopt the *Z*-configuration based on spectroscopic analysis. In addition, reactions of substrate **1** with 1,3-binucleophilic reagents enabled the construction of novel pyrimidine-fused heterocycles *via* ring opening followed by intramolecular cyclization. DFT calculations (B3LYP/6-311+G(d,p)) were performed to study the electronic and nonlinear optical properties of the present compounds. Compound **5** was found to be the most stable than compound **3** and compound **4**, consistent with experimental results. Theoretical IR and NMR spectra agreed well with experimental data. The high hyperpolarizability values of the prepared compounds suggest potential nonlinear optical applications. The synthesized compounds were evaluated for anticancer activity against HepG-2 cells, revealing compound **9** as the most active derivative, with potency exceeding that of cisplatin. Molecular docking studies against CDK1 supported the observed biological activity, while drug-likeness analysis indicated favorable pharmacokinetic profiles. Overall, this work highlights the utility of 2-methylchromone with diverse nucleophiles in generating structurally varied heterocycles, validated by combined experimental and theoretical studies.

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## 1. Introduction

Chromones constitute an important class of oxygen-containing heterocycles that function not only as bioactive scaffolds but also as highly versatile synthetic intermediates in heterocyclic chemistry.<sup>1–3</sup> Beyond their intrinsic biological and photo-physical relevance,<sup>4–8</sup> chromones are distinguished by their electron-deficient conjugated systems, which impart pronounced electrophilic character and render them particularly susceptible to nucleophilic attack. This reactivity facilitates a wide range of subsequent structural transformations, underscoring the value of chromones as privileged building blocks in organic synthesis.<sup>9–12</sup>

Among these derivatives, 2-methylchromones represent an important class of oxygen-containing heterocyclic compounds that have attracted considerable interest due to their diverse biological activities and synthetic versatility.<sup>13,14</sup> Typically, 2-methylchromones are prepared through the Claisen condensation of 2-hydroxyacetophenones with ethyl acetate, followed by intramolecular acid-catalyzed cyclocondensation, providing efficient access to the chromone framework with structural control.<sup>15–17</sup> The reactivity and electronic structure of 2-methylchromones have been systematically examined, especially in the earlier work by Ibrahim *et al.*,<sup>18</sup> who demonstrated that the chromone ring possess two principal electrophilic centers located at C-2 and C-4. These positions play a pivotal role in governing the compound's behavior toward nucleophiles and thus determine the synthetic pathways available for functionalizing the core.<sup>18</sup> Owing to the electron-deficient character of the chromone scaffold and the strong conjugation with carbonyl group, the C-2 position represents the most reactive electrophilic center toward nucleophiles. As a result, nucleophilic attack predominantly occurs at this site, often generating subsequent opening of the pyran ring. This ring-opening pathway is a well-

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established mechanistic feature in chromone chemistry and has been extensively documented in the literature.<sup>19</sup>

Over the past decade, theoretical studies have gained increasing importance, with computational and experimental chemistry being recognized as complementary approaches in modern research. Among computational methods, density functional theory (DFT) has emerged as a powerful tool, offering profound insights into both the kinetic stability and chemical reactivity of molecular systems.<sup>20,21</sup> Herein, DFT calculations were employed not to predict unknown outcomes, but to quantitatively rationalize experimental observations. These include charge distribution, electrophilicity at the C-2 position, preferences for ring opening, and the relative stability of the resulting enaminone and heterocyclic structures. Such calculations provide a molecular-level insight that complements the experimental data and enables meaningful comparison with previously reported chromone systems.<sup>8–10</sup>

The aim and novelty of the present work differ fundamentally from our earlier studies. Whereas previous investigations focused on 3-substituted chromones bearing electron-withdrawing groups that markedly enhance the electrophilicity at C-2,<sup>8–12</sup> the current study concisely on 2-methylchromone, in which the methyl substituent introduces both electron-donating and steric effects. This distinct substitution pattern creates a substantially different electronic environment, enabling a systematic evaluation of how reduced electrophilicity and altered charge distribution affect reaction pathways. Moreover, while the previous work on 2-methylchromones primarily involved simple nitrogen nucleophiles, the use of amino-functionalized nitrogen heterocycles as nucleophiles is explored here for the first time. This choice was motivated by the objective of generating enaminone systems incorporating heterocyclic fragments, which are synthetically valuable motifs with recognized relevance in medicinal chemistry and molecular recognition.

In this study, we report the synthesis of a series of novel heterocyclic derivatives *via* the reaction of 2-methylchromone (**1**) with selected heterocyclic amines and 1,3-binucleophilic reagents, followed by evaluation of their anticancer activity against HepG-2 cells. To gain deeper insight into their structural and electronic properties, DFT calculations were carried out at the B3LYP/6-311++G(d,p) level, where global reactivity descriptors were derived from HOMO–LUMO energies, and experimental NMR and FT-IR data were validated through GIAO-based theoretical simulations. Additionally, nonlinear optical (NLO) properties and molecular electrostatic potential (MEP) surfaces were examined. Molecular docking studies were performed against the CDK1 protein (PDB ID: 4Y72) to explore possible binding interactions, and drug-likeness assessment was conducted according to Lipinski, Egan, Veber, and Ghose rules.

## 2. Experimental

Melting points were determined on a digital Stuart SMP3 apparatus. Elemental microanalyses were performed on a PerkinElmer CHN-2400 analyzer. Infrared spectra were measured on FTIR Nicolet IS10 spectrophotometer (cm<sup>-1</sup>), using KBr

disks. <sup>1</sup>H NMR spectra were measured on mercury-400BB (400 MHz), using DMSO-*d*<sub>6</sub> as a solvent and TMS ( $\delta$ ) as the internal standard. <sup>13</sup>C NMR spectra were measured on mercury-400BB (100 MHz), using DMSO-*d*<sub>6</sub> as a solvent and TMS ( $\delta$ ) as the internal standard. Mass spectra were obtained using GC-2010 Shimadzu Gas chromatography instrument mass spectrometer (70 eV). The purity of the synthesized compounds was tested using TLC. 1-Amino-4,6-dimethyl-2-oxo-1,2-dihydropyridine-3-carbonitrile (**2**),<sup>22</sup> 4-amino-5-methyl-2,4-dihydro-3*H*-1,2,4-triazole-3-thione (**6**)<sup>23</sup> and 4-amino-6-methyl-3-thioxo-3,4-dihydro-1,2,4-triazin-5(2*H*)-one (**7**)<sup>24</sup> were synthesized according to literature.

### 2.1. Synthesis and characterization of compounds

**2.1.1. 1-[(2*Z*)-4-(2-Hydroxyphenyl)-4-oxobut-2-en-2-yl]amino]-4,6-dimethyl-2-oxo-1,2-dihydropyridine-3-carbonitrile (**5**).** A mixture of 2-methylchromone (**1**) (0.32 g, 2 mmol) and 1-amino-4,6-dimethyl-2-oxo-1,2-dihydropyridine-3-carbonitrile (**2**) (0.32 g, 2 mmol) in absolute ethanol (10 mL) containing DBU (0.2 mL) was heated under reflux for 2 h. The pale yellow crystals obtained during heating were filtered and crystallized from EtOH, mp 205–206 °C, yield (0.47 g, 73%). IR (KBr, cm<sup>-1</sup>): 3406 (OH), 3279 (NH), 2217 (C≡N), 1676 (C=O<sub>pyridone</sub>), 1642 (C=O<sub>enaminone</sub>), 1588 (C=C). <sup>1</sup>H NMR (DMSO-*d*<sub>6</sub>,  $\delta$ ): 2.10 (s, 3H, CH<sub>3</sub>), 2.31 (s, 3H, CH<sub>3</sub>), 2.39 (s, 3H, CH<sub>3</sub>), 6.20 (s, 1H, H- $\alpha$ <sub>olefinic</sub>), 6.34 (s, 1H, H-5<sub>pyridine</sub>), 7.38 (t, 1H, *J* = 7.2 Hz, Ar-H), 7.61 (d, 1H, *J* = 7.2 Hz, Ar-H), 7.81 (t, 1H, *J* = 7.5 Hz, Ar-H), 8.12 (d, 1H, *J* = 7.5 Hz, Ar-H), 9.80 (bs, 1H, NH exchangeable with D<sub>2</sub>O), 12.22 (bs, 1H, OH exchangeable with D<sub>2</sub>O). <sup>13</sup>C NMR (DMSO-*d*<sub>6</sub>,  $\delta$ ): 15.9 (CH<sub>3</sub>), 17.5 (CH<sub>3</sub>), 19.4 (CH<sub>3</sub>), 94.2 (C- $\alpha$ <sub>olefinic</sub>), 106.2 (C-3<sub>pyridone</sub>), 108.3 (C-5<sub>pyridone</sub>), 116.4 (C≡N), 122.2, 124.9, 126.8, 128.4, 129.5 (5Ar-C), 132.8 (C-6<sub>pyridone</sub>), 142.3 (C-4<sub>pyridone</sub>), 148.2 (C- $\beta$ <sub>olefinic</sub>), 152.1, (C-OH), 163.4 (C=O<sub>pyridone</sub>), 182.6 (C=O<sub>ketone</sub>). Mass spectrum, *m/z* (*I*<sub>r</sub>%): 323 (M<sup>+</sup>, 11), 308 (17), 215 (16), 176 (100), 161 (34), 146 (16), 121 (22), 93 (62), 77 (42), 65 (13). Anal. calcd for C<sub>18</sub>H<sub>17</sub>N<sub>3</sub>O<sub>3</sub> (323.34); C, 66.86; H, 5.30; N, 13.00%. Found: C, 66.58; H, 5.12; N, 12.84%.

**2.1.2. (2*Z*)-1-(2-Hydroxyphenyl)-3-[(3-methyl-5-thioxo-1,5-dihydro-4*H*-1,2,4-triazol-4-yl)amino]but-2-en-1-one (**8**).** A mixture of 2-methylchromone (**1**) (0.32 g, 2 mmol) and 4-amino-5-methyl-2,4-dihydro-3*H*-1,2,4-triazole-3-thione (**6**) (0.26 g, 2 mmol) in absolute ethanol (10 mL) containing DBU (0.2 mL) was heated under reflux for 2 h. The yellow crystals obtained during heating were filtered and crystallized from isopropanol, mp 236–237 °C, yield (0.46 g, 77%). IR (KBr, cm<sup>-1</sup>): 3409 (OH), 3248, 3211 (2NH), 1644 (C=O<sub>enaminone</sub>), 1613 (C≡N), 1587 (C=C), 1223 (C=S). <sup>1</sup>H NMR (DMSO-*d*<sub>6</sub>,  $\delta$ ): 2.27 (s, 3H, CH<sub>3</sub>), 2.37 (s, 3H, CH<sub>3</sub>), 6.32 (s, 1H, H- $\alpha$ <sub>olefinic</sub>), 7.54 (t, 1H, *J* = 7.5 Hz, Ar-H), 7.72 (d, 1H, *J* = 7.8 Hz, Ar-H), 7.82 (t, 1H, *J* = 7.5 Hz, Ar-H), 8.14 (d, 1H, *J* = 7.5 Hz, Ar-H), 9.94 (bs, 1H, NH exchangeable with D<sub>2</sub>O), 10.74 (bs, 1H, NH exchangeable with D<sub>2</sub>O), 11.81 (bs, 1H, OH exchangeable with D<sub>2</sub>O). <sup>13</sup>C NMR (DMSO-*d*<sub>6</sub>,  $\delta$ ): 15.6 (CH<sub>3</sub>), 16.4 (CH<sub>3</sub>), 96.1 (C- $\alpha$ <sub>olefinic</sub>), 122.6, 123.3, 126.4, 125.1, 128.6 (5Ar-C), 144.3 (C-3<sub>triazole</sub>), 148.1 (C- $\beta$ <sub>olefinic</sub>), 150.8, (C-OH), 182.3 (C=O<sub>ketone</sub>), 194.5 (C=S). Mass spectrum, *m/z* (*I*<sub>r</sub>%): 290 (M<sup>+</sup>, 21), 245 (13), 219 (18), 176 (100), 163 (31), 148 (19), 122 (27), 105



(54), 94 (76), 77 (49), 65 (22). Anal. calcd for  $C_{13}H_{14}N_4O_2S$  (290.34); C, 53.78; H, 4.86; N, 19.30, S, 11.04%. Found: C, 53.61; H, 4.79; N, 19.13, S, 10.86%.

**2.1.3. 4-[[[(2Z)-4-(2-Hydroxyphenyl)-4-oxobut-2-en-2-yl]amino]-6-methyl-3-thioxo-3,4-dihydro-1,2,4-triazin-5(2H)-one (9).** A mixture of 2-methylchromone (1) (0.32 g, 2 mmol) and 4-amino-6-methyl-3-thioxo-3,4-dihydro-1,2,4-triazin-5(2H)-one (7) (0.48 g, 2 mmol) in absolute ethanol (10 mL) containing DBU (0.2 mL) was heated under reflux for 2 h. The pale yellow crystals obtained during heating were filtered and crystallized from EtOH, mp 251–252 °C, yield (0.48 g, 75%). IR (KBr,  $cm^{-1}$ ): 3402 (OH), 3237, 3198 (2NH), 1692 (C=O<sub>triazine</sub>), 1646 (C=O<sub>enaminone</sub>), 1570 (C=C), 1226 (C=S). <sup>1</sup>H NMR (DMSO-*d*<sub>6</sub>,  $\delta$ ): 2.18 (s, 3H, CH<sub>3</sub>), 2.40 (s, 3H, CH<sub>3</sub>), 6.21 (s, 1H, H- $\alpha$ <sub>olefinic</sub>), 7.40 (t, 1H, *J* = 7.2 Hz, Ar-H), 7.55 (d, 1H, *J* = 7.2 Hz, Ar-H), 7.84 (t, 1H, *J* = 7.5 Hz, Ar-H), 8.06 (d, 1H, *J* = 7.5 Hz, Ar-H), 9.81 (bs, 1H, NH exchangeable with D<sub>2</sub>O), 11.65 (bs, 1H, NH exchangeable with D<sub>2</sub>O), 12.36 (bs, 1H, OH exchangeable with D<sub>2</sub>O). <sup>13</sup>C NMR (DMSO-*d*<sub>6</sub>,  $\delta$ ): 15.7 (CH<sub>3</sub>), 17.5 (CH<sub>3</sub>), 95.9 (C- $\alpha$ <sub>olefinic</sub>), 122.5, 124.7, 126.3, 128.6, 130.5 (5Ar-C), 144.8 (C-6<sub>triazine</sub>), 148.6 (C- $\beta$ <sub>olefinic</sub>), 151.7 (C-OH), 166.5 (C=O<sub>triazine</sub>), 182.7 (C=O<sub>ketone</sub>), 195.8 (C=S). Mass spectrum, *m/z* (*I*<sub>r</sub>%): 318 (M<sup>+</sup>, 15), 303 (27), 275 (23), 231 (24), 176 (100), 142 (39), 120 (36), 93 (45), 77 (34), 65 (23). Anal. calcd for  $C_{14}H_{14}N_4O_3S$  (318.35); C, 52.82; H, 4.43; N, 17.60, S, 10.07%. Found: C, 52.67; H, 4.35; N, 17.41, S, 9.83%.

**2.1.4. 6-(2-Hydroxyphenyl)-4-methyl-1,2-dihydro-3H-pyrazolo[3,4-*b*]pyridin-3-one (10).** A mixture of 2-methylchromone (1) (0.32 g, 2 mmol) and 5-amino-2,4-dihydro-3H-pyrazol-3-one (0.25 g, 2 mmol), in sodium ethoxide (prepared by adding 0.2 g sodium in 20 mL absolute ethanol) was heated under reflux for 2 h. After cooling, the reaction mixture was neutralized with conc. HCl. The solid obtained were filtered and crystallized from DMF/H<sub>2</sub>O as yellow crystals, mp 296–297 °C, yield (0.37 g, 77%). IR (KBr,  $cm^{-1}$ ): 3405 (OH), 3291 (2NH), 1659 (C=O<sub>pyrazole</sub>), 1609 (C=N), 1577 (C=C). <sup>1</sup>H NMR (DMSO-*d*<sub>6</sub>,  $\delta$ ): 2.09 (s, 3H, CH<sub>3</sub>), 7.27 (t, 1H, *J* = 7.2 Hz, Ar-H), 7.42 (d, 1H, *J* = 7.5 Hz, Ar-H), 7.76 (t, 1H, *J* = 7.5 Hz, Ar-H), 8.00 (d, 1H, *J* = 7.2 Hz, Ar-H), 8.37 (s, 1H, H-3<sub>pyridine</sub>), 9.84 (bs, 1H, NH exchangeable with D<sub>2</sub>O), 10.72 (bs, 1H, NH exchangeable with D<sub>2</sub>O), 11.90 (bs, 1H, OH exchangeable with D<sub>2</sub>O). <sup>13</sup>C NMR (DMSO-*d*<sub>6</sub>,  $\delta$ ): 15.6 (CH<sub>3</sub>), 112.2 (C-3a), 122.3 (C-3), 123.7, 125.4, 126.4, 128.7, 129.9 (5Ar-C), 142.2 (C-4), 144.5 (C-6), 146.3 (C-7a), 151.6 (C-OH), 165.4 (C-3 as C=O). Mass spectrum, *m/z* (*I*<sub>r</sub>%): 241 (M<sup>+</sup>, 66), 226 (22), 214 (15), 198 (35), 184 (27), 148 (36), 120 (100), 93 (63), 77 (55), 64 (38). Anal. calcd for  $C_{13}H_{11}N_3O_2$  (241.24); C, 64.72; H, 4.60; N, 17.42%. Found: C, 64.68; H, 4.45; N, 17.38%.

**2.1.5. 6-(2-Hydroxyphenyl)-3,8-dimethyl-4H-pyrimido[2,1-*c*][1,2,4]triazin-4-one (12).** A mixture of 2-methylchromone (1) (0.32 g, 2 mmol) and 3-amino-6-methyl-1,2,4-triazin-5(4H)-one (11) (0.25 g, 2 mmol), in sodium ethoxide (prepared by adding 0.2 g sodium in 20 mL absolute ethanol) was heated under reflux for 2 h. After cooling, the reaction mixture was neutralized with conc. HCl. The solid obtained were filtered and crystallized from AcOH as pale-yellow crystals, mp > 300 °C, yield (0.35 g, 65%). IR (KBr,  $cm^{-1}$ ): 3405 (OH), 1655 (C=O<sub>triazine</sub>), 1608 (C=N), 1566 (C=C). <sup>1</sup>H NMR (DMSO-*d*<sub>6</sub>,  $\delta$ ): 2.15 (s, 3H,

CH<sub>3</sub>), 2.29 (s, 3H, CH<sub>3</sub>), 7.38 (t, 1H, *J* = 7.5 Hz, Ar-H), 7.58 (d, 1H, *J* = 7.5 Hz, Ar-H), 7.82 (t, 1H, *J* = 7.2 Hz, Ar-H), 8.01 (d, 1H, *J* = 7.5 Hz, Ar-H), 8.24 (s, 1H, H-5<sub>pyrimidine</sub>), 11.04 (bs, 1H, OH exchangeable with D<sub>2</sub>O). <sup>13</sup>C NMR (DMSO-*d*<sub>6</sub>,  $\delta$ ): 16.7 (CH<sub>3</sub>), 18.5 (CH<sub>3</sub>), 108.7 (C-7), 122.7, 124.3, 126.4, 127.2, 128.6 (5Ar-C), 142.3 (C-6), 144.3 (C-3), 145.8 (C-8), 147.4 (C-9a), 151.6 (C-OH), 170.5 (C-3 as C=O). Mass spectrum, *m/z* (*I*<sub>r</sub>%): 268 (100), 240 (62), 225 (20), 199 (29), 175 (43), 120 (56), 91 (38), 77 (26), 65 (13). Anal. calcd for  $C_{14}H_{12}N_4O_2$  (268.27); C, 62.68; H, 4.51; N, 20.88%. Found: C, 62.57; H, 4.50; N, 20.73%.

**2.1.6. 5-(2-Hydroxyphenyl)-7-methyl[1,2,4]triazolo[4,3-*a*]pyrimidine (13).** A mixture of 2-methylchromone (1) (0.32 g, 2 mmol) and 3-amino-1,2,4-triazole (0.17 g, 2 mmol) in sodium ethoxide (prepared by adding 0.2 g sodium in 15 mL absolute ethanol) was heated under reflux for 2 h. After cooling, the reaction mixture was poured onto crushed ice (50 g) and neutralized with conc. HCl. The solid obtained filtered and crystallized from DMF/H<sub>2</sub>O, mp 268–269 °C, yield (0.32 g, 71%). IR (KBr,  $cm^{-1}$ ): 3416 (OH), 1608 (C=N), 1587 (C=C). <sup>1</sup>H NMR (DMSO-*d*<sub>6</sub>,  $\delta$ ): 2.25 (s, 3H, CH<sub>3</sub>), 7.35 (t, 1H, *J* = 7.2 Hz, Ar-H), 7.53 (d, 1H, *J* = 7.2 Hz, Ar-H), 7.69 (t, 1H, *J* = 7.2 Hz, Ar-H), 7.99 (d, 1H, *J* = 7.2 Hz, Ar-H), 8.38 (s, 1H, H-5<sub>pyrimidine</sub>), 9.12 (s, 1H, H-3<sub>triazole</sub>), 11.75 (bs, 1H, OH exchangeable with D<sub>2</sub>O). <sup>13</sup>C NMR (DMSO-*d*<sub>6</sub>,  $\delta$ ): 18.4 (CH<sub>3</sub>), 121.7, 123.5, 125.9, 127.3, 128.7 (5Ar-C), 132.3 (C-5<sub>pyrimidine</sub>), 140.4 (C-4<sub>pyrimidine</sub>), 140.9 (C-6<sub>pyrimidine</sub>), 141.2 (C-3), 142.5 (C-8a), 150.3, (C-OH). Mass spectrum, *m/z* (*I*<sub>r</sub>%): 226 (M<sup>+</sup>, 44), 214 (11), 195 (27), 148 (16), 133 (40), 107 (21), 94 (100), 77 (56), 64 (39). Anal. calcd for  $C_{12}H_{10}N_4O$  (226.23); C, 63.71; H, 4.46; N, 24.76%. Found: C, 63.58; H, 4.28; N, 24.54%.

**2.1.7. 4-(2-Hydroxyphenyl)-2-methylpyrimido[1,2-*a*]benzimidazole (14).** A mixture of 2-methylchromone (1) (0.32 g, 2 mmol) and 2-aminobenzimidazole (0.27 g, 2 mmol) in sodium ethoxide (prepared by adding 0.2 g sodium in 15 mL absolute ethanol) was heated under reflux for 2 h. After cooling, the reaction mixture was poured onto crushed ice (50 g) and neutralized with conc. HCl. The solid obtained filtered and crystallized from AcOH, mp > 300 °C, yield (0.40 g, 74%). <sup>13</sup>C NMR (DMSO-*d*<sub>6</sub>,  $\delta$ ): 17.5 (CH<sub>3</sub>), 122.7, 123.2, 124.1, 125.3, 126.2, 126.7, 127.2, 128.5, 129.1, 129.7, 130.4, (Ar-C), 131.5 (C-5<sub>pyrimidine</sub>), 140.1 (C-6<sub>pyrimidine</sub>), 140.9 (C-4<sub>pyrimidine</sub>), 142.2 (C-2<sub>pyrimidine</sub>), 150.6 (C-OH). IR (KBr,  $cm^{-1}$ ): 3410 (OH), 1615 (C=N), 1583 (C=C). <sup>1</sup>H NMR (DMSO-*d*<sub>6</sub>,  $\delta$ ): 2.26 (s, 3H, CH<sub>3</sub>), 7.29–7.40 (m, 4H, Ar-H), 7.54 (t, 1H, *J* = 7.8 Hz, Ar-H), 7.74 (d, 1H, *J* = 7.8 Hz, Ar-H), 7.83 (t, 1H, *J* = 7.5 Hz, Ar-H), 8.08 (d, 1H, *J* = 7.8 Hz, Ar-H), 8.37 (s, 1H, H-5<sub>pyrimidine</sub>), 11.39 (bs, 1H, OH exchangeable with D<sub>2</sub>O). Mass spectrum, *m/z* (*I*<sub>r</sub>%): 275 (M<sup>+</sup>, 100), 260 (25), 244 (16), 183 (46), 169 (32), 117 (26), 93 (70), 77 (45), 65 (23). Anal. calcd for  $C_{17}H_{13}N_3O$  (275.30); C, 74.17; H, 4.76; N, 15.26%. Found: C, 74.02; H, 4.73; N, 15.15%.

## 2.2. Computational methods

Computational chemistry analyses for the synthesized compounds were carried out using DFT at the B3LYP level with the standard 6-311++G(d,p) basis set, employing the GAUSSIAN 09W software. This method is among the most reliable for assessing compound stability and reactivity.<sup>25,26</sup> Molecular



visualizations were generated using GaussView 5.0.<sup>27–29</sup> No symmetry constraints were applied during geometry optimization. Quantum chemical calculations were performed to evaluate the molecular electrostatic potential (MEP), optimized geometries, and molecular orbital (MO) energy levels. Additionally, chemical shift predictions for <sup>1</sup>H and <sup>13</sup>C NMR were performed using the GIAO method using the same basis set at the B3LYP level,<sup>30</sup> and the results were compared with the experimental data. Further, vibrational frequencies were computed for all new compounds. Polarizability, first hyperpolarizability and electronic dipole moment were also determined at the same DFT level of theory.<sup>31</sup>

### 2.3. Biological evaluation

**2.3.1. Antitumor activity.** The antitumor activity of the synthesized compounds was tested against HepG2 cells using the published procedure of evaluating the effect of the test samples on cell morphology and viability.<sup>32–34</sup> Mammalian cell lines: HepG-2 cells (human Hepatocellular carcinoma) were obtained from the American Type Culture Collection (ATCC, Rockville, MD).

**2.3.2. ADME analysis.** A computational ADME analysis was conducted using SwissADME to evaluate the physicochemical and drug-likeness properties of the studied compounds.<sup>35,36</sup>

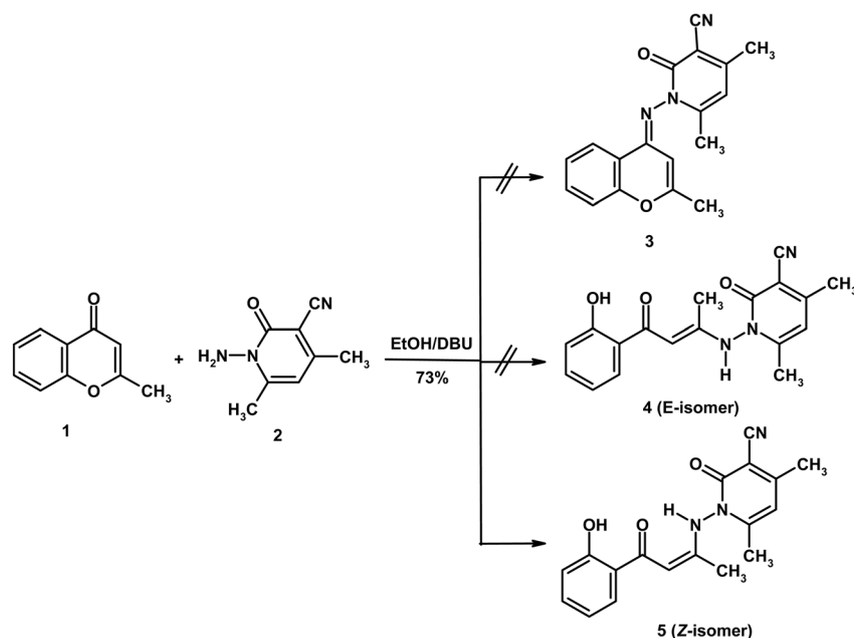
**2.3.3. Molecular docking study.** Molecular docking studies were performed using AutoDock Vina<sup>37</sup> to explore the interaction profiles of the synthesized compounds with cyclin-dependent kinase 1 (CDK1; PDB ID: 4Y72).<sup>38</sup> The crystal structure of the target protein was obtained from the Protein Data Bank. Prior to docking, the receptor was prepared by removing all crystallographic water molecules, co-crystallized ligands, and heteroatoms, followed by the addition of polar hydrogens and Kollman charges to ensure proper protonation and electrostatic configuration.<sup>39,40</sup>

Ligand structures were sketched in ChemDraw, subjected to energy minimization, and subsequently converted into PDBQT format. Docking was carried out using a grid box centered at coordinates ( $X = 29.509$ ,  $Y = -70.550$ ,  $Z = 185.362$ ) with dimensions ( $X = 13.738$  Å,  $Y = 12.295$  Å,  $Z = 11.474$  Å), ensuring coverage of the active binding region. Binding energies (kcal mol<sup>-1</sup>) were used as the primary criterion for evaluating ligand–protein interactions and predicting binding affinity.<sup>41,42</sup>

## 3. Results and discussion

### 3.1. Characterization of the synthesized compounds

The present work focuses on exploring the reactivity of 2-methylchromone (**1**) toward a variety of heterocyclic amines. Initially, its behavior was examined with 1-amino-4,6-dimethyl-2-oxo-1,2-dihydropyridine-3-carbonitrile (**2**)<sup>22</sup> in absolute ethanol using DBU as a base (Scheme 1). The structure of the obtained product was established through spectroscopic analyses combined with theoretical calculations. Formation of the condensation product **3** was excluded, since the carbonyl group is generally unreactive toward condensation reactions.<sup>18</sup> Instead, the most plausible pathway involves nucleophilic attack at the C-2 position, which is activated by both the mesomeric electron-withdrawing effect of the carbonyl group and the inductive effect of the adjacent oxygen atom.<sup>19</sup> Following the ring opening of substrate **1**, two possible conformers could be generated: **4** (*E*-isomer) and **5** (*Z*-isomer). Among these, the *Z*-conformer **5** is favored due to intramolecular hydrogen bonding between the carbonyl oxygen and the NH proton. The predominance of structure **5** was further validated through spectral evidence and theoretical calculations, both of which play a crucial role in confirming its identity as the *Z*-conformer. In this context, DFT calculations were



Scheme 1 Reaction of 2-methylchromone (**1**) and aminopyridine derivative **2**.



employed as a complementary tool to support the experimental assignment and to rationalize the relative stability of the *Z* and *E* isomers, rather than to replace NMR-based structural elucidation (*vide infra*).

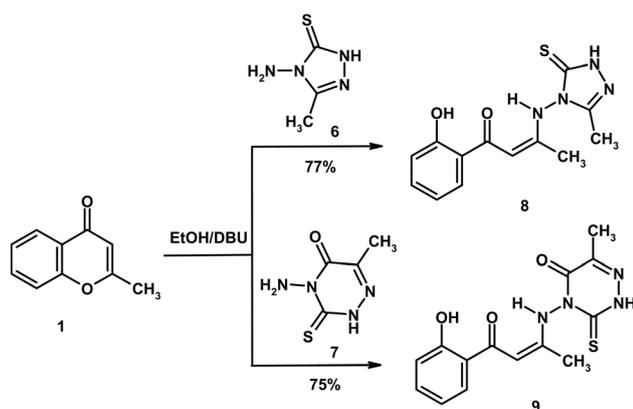
The *Z*-configuration of compound **5** was confirmed by spectroscopic evidence. In the IR spectrum, the C=O stretching vibration was observed at a relatively low frequency (1642 cm<sup>-1</sup>), indicative of intramolecular hydrogen bonding with the NH proton. This hydrogen-bonding interaction is further supported by the pronounced downfield shift of the NH signal in the <sup>1</sup>H NMR spectrum, appearing at  $\delta$  9.80 and reflecting deshielding associated with hydrogen-bond-induced bond elongation. Such stabilization effectively clarifies for the preferential formation of the *Z*-isomer over the *E*-isomer. Further, in the <sup>1</sup>H NMR spectrum, three singlet signals were recorded in the upfield region at  $\delta$  2.10, 2.31, and 2.39 correspond to the three methyl groups. Two additional singlets at  $\delta$  6.20 and 6.34 were assigned to H- $\alpha$  (olefinic) and H-5 (pyridine), respectively, while the aromatic protons resonated as expected at  $\delta$  7.38–8.12. The <sup>13</sup>C NMR spectrum exhibited three methyl carbons at  $\delta$  15.9, 17.5, and 19.4. The olefinic  $\alpha$ -carbon appeared at  $\delta$  94.2, shifted upfield by shielding by the electron donating mesomeric effect of the NH group, while the  $\beta$ -carbon resonated at  $\delta$  148.2 due to deshielding by electron withdrawing mesomeric effect of the carbonyl group and the inductive effect adjacent nitrogen atoms. The mass spectrum confirmed the proposed structure, showing a molecular ion peak at *m/z* 323 (11%) and a base peak at *m/z* 176 (100%), the latter resulting from facile cleavage of the N–N bond.

Similarly, treatment of 2-methylchromone (**1**) with 4-amino-5-methyl-2,4-dihydro-3*H*-1,2,4-triazole-3-thione (**6**)<sup>23</sup> and 4-

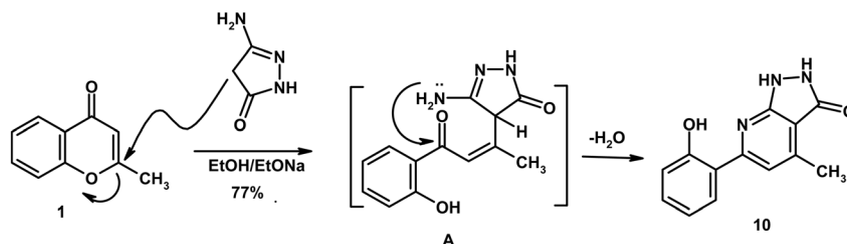
amino-6-methyl-3-thioxo-3,4-dihydro-1,2,4-triazin-5(2*H*)-one (**7**)<sup>24</sup> in absolute ethanol containing DBU afforded the enaminone derivatives **8** and **9**, respectively (Scheme 2). Both compounds gave a red coloration with FeCl<sub>3</sub> solution, indicating the presence of a free phenolic OH group and supporting the pathway of ring-opening. Their mass spectra exhibited molecular ion peaks at *m/z* 290 (21%) and 318 (15%) supporting the molecular formula of C<sub>13</sub>H<sub>14</sub>N<sub>4</sub>O<sub>2</sub>S (290.34) and C<sub>14</sub>H<sub>14</sub>N<sub>4</sub>O<sub>3</sub>S (318.35), that consistent with the proposed structures. The IR spectra of **8** and **9** displayed characteristic absorptions for C=O<sub>enaminone</sub> at 1646 and 1644 cm<sup>-1</sup>, respectively, while compound **9** also showed a distinct absorption band due to C=O<sub>triazine</sub> at 1692 cm<sup>-1</sup>. The <sup>1</sup>H NMR spectra of compounds **8** and **9** revealed singlets for the olefinic proton at  $\delta$  6.32 and 6.21, respectively. Signals for NH ( $\delta$  9.94/9.81) and OH ( $\delta$  11.81/12.36) were also observed, both shifted downfield due to hydrogen bonding with carbonyl groups. In the <sup>13</sup>C NMR spectrum of compound **9**, the olefinic  $\alpha$ -carbon resonated at  $\delta$  95.9, whereas the  $\beta$ -carbon resonated at  $\delta$  148.6.

On the other hand, reaction of 2-methylchromone (**1**) with 5-amino-2,4-dihydro-3*H*-pyrazol-3-one in sodium ethoxide solution afforded pyrazolo[3,4-*b*]pyridine **10**. Herein, sodium ethoxide was employed to generate enolate species from active methylene compounds and to ensure effective ring opening of the  $\gamma$ -pyrone system. The mechanism proceeds *via*  $\gamma$ -pyrone ring opening by the deprotonated CH<sub>2</sub> group (intermediate **A**), followed by intramolecular heterocyclization (Scheme 3). The mass spectrum of compound **10** exhibited a molecular ion peak at *m/z* 241, that consistent with the molecular formula C<sub>13</sub>H<sub>11</sub>N<sub>3</sub>O<sub>2</sub>. The IR spectrum displayed absorption bands at 3405 (OH), 3291 (2NH), 1659 (C=O<sub>pyrazole</sub>), and 1609 cm<sup>-1</sup> (C=N). The <sup>1</sup>H NMR spectrum revealed singlets at  $\delta$  2.09 and 8.37, assigned to CH<sub>3</sub> and H-3<sub>pyridine</sub>, respectively. D<sub>2</sub>O-exchangeable signals due to 2NH and OH were recorded at  $\delta$  9.84, 10.72 and 11.90, respectively.

Similarly, reaction of 2-methylchromone (**1**) with 3-amino-6-methyl-1,2,4-triazin-5(4*H*)-one (**11**),<sup>24</sup> in sodium ethoxide solution under reflux, afforded pyrimido[2,1-*c*][1,2,4]triazine **12**; *via*  $\gamma$ -pyrone ring opening (intermediate **B**) followed by heterocyclization (Scheme 4). The mass spectrum displayed a molecular ion peak, as the base peak, at *m/z* 268 (C<sub>14</sub>H<sub>12</sub>N<sub>4</sub>O<sub>2</sub>). The IR spectrum showed characteristic absorption bands at 3405 (OH), 1655 (C=O<sub>triazine</sub>) and 1608 (C=N) cm<sup>-1</sup>. The <sup>1</sup>H NMR spectrum exhibited two singlets at  $\delta$  2.15 and 2.29, attributed to the methyl groups, along with a distinctive singlet at  $\delta$  8.24 corresponding to H-5 of the pyrimidine ring.

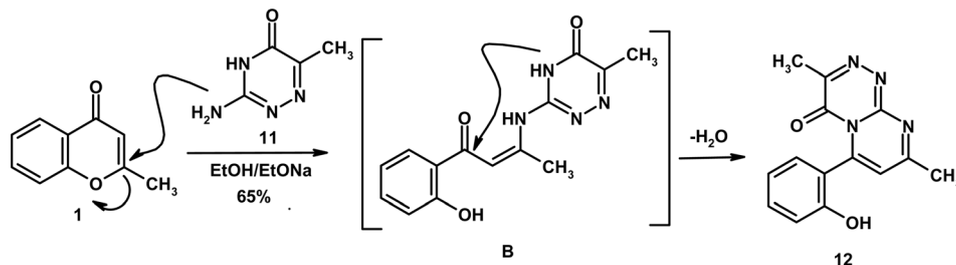


Scheme 2 Formation of enaminone derivatives **8** and **9**.

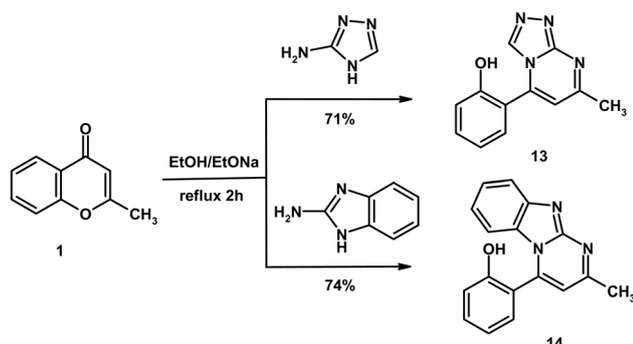


Scheme 3 Formation of pyrazolo[3,4-*b*]pyridine derivative **10**.





Scheme 4 Formation of pyrimido[2,1-c][1,2,4] triazine 12.



Scheme 5 Formation of triazolo[4,3-a]pyrimidine 13 and pyrimido[1,2-a]benzimidazole 14.

Finally, reaction of substrate 1 with 3-amino-1,2,4-triazole and 2-aminobenzimidazole in ethanolic sodium ethoxide solution afforded the novel heterocycles triazolo[4,3-*a*]pyrimidine 13 and pyrimido[1,2-*a*]benzimidazole 14, respectively

(Scheme 5). Both compounds produced a red coloration with  $\text{FeCl}_3$  solution, indicating the presence of a free OH group arising from  $\gamma$ -pyrone ring opening. The IR spectra of products 13 and 14 confirmed the absence of the carbonyl absorption band ( $\text{C}=\text{O}_{\gamma\text{-pyrone}}$ ) and exhibited new absorption bands due to  $\text{C}=\text{N}$  at 1608 and 1615  $\text{cm}^{-1}$ , respectively. Their mass spectra supported the proposed structures, displaying molecular ion peaks at  $m/z$  226 ( $\text{C}_{12}\text{H}_{10}\text{N}_4\text{O}$ , 226.23) for compound 13 and  $m/z$  275 ( $\text{C}_{17}\text{H}_{13}\text{N}_3\text{O}$ , 275.30) for compound 14. In the  $^1\text{H}$  NMR spectra, both compounds showed characteristic singlets for H-5 of the pyrimidine ring at  $\delta$  8.38 (compound 13) and 8.37 (compound 14). Additionally, compound 13 displayed a further singlet at  $\delta$  9.12, assigned to H-3 of the triazole moiety.

### 3.2. Theoretical studies

The primary objective of the theoretical investigation was to complement experimental studies. Accordingly, DFT calculations were carried out to rationalize the experimentally obtained products, providing molecular-level insight into charge

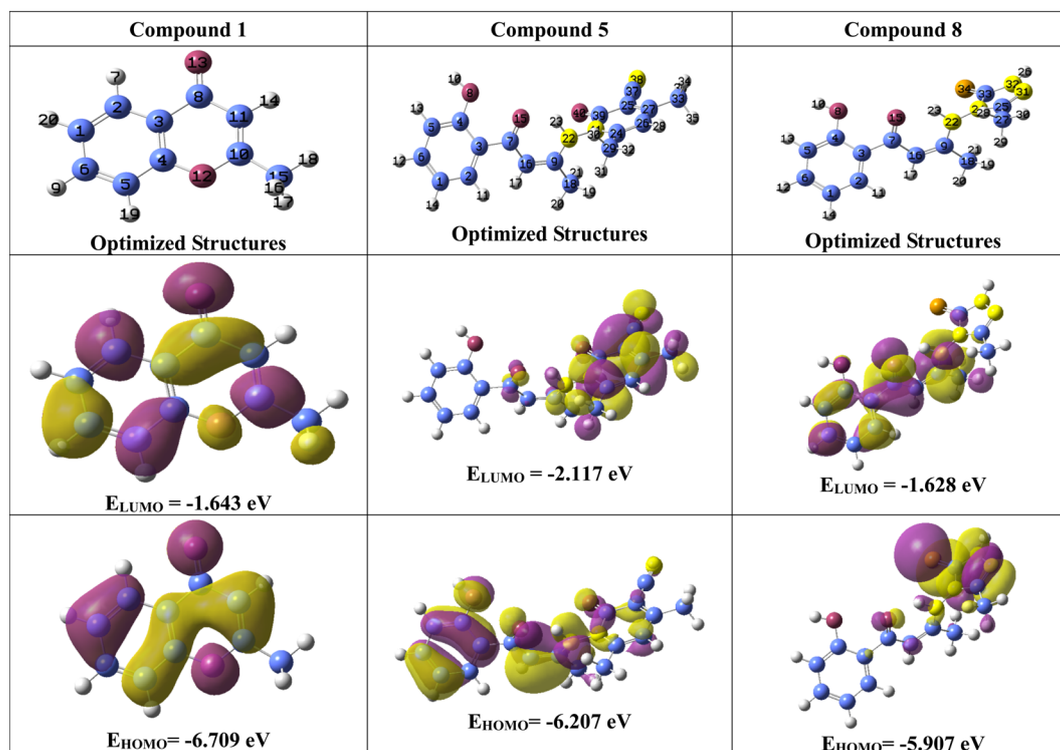


Fig. 1 Molecular modeling and the electron density of HOMO and LUMO of compounds 1, 5 and 8.



distribution, preferred reaction pathways, and the relative stability of the resulting enaminone and heterocyclic systems, rather than to predict new reactions.

**3.2.1. Frontier molecular orbital energies and chemical reactivity.** The molecular structures of the synthesized compounds were optimized using DFT at the B3LYP/6-311+G(d,p) level. The frontier molecular orbital (FMO) analysis (Fig. 1, S1 and S2) revealed extensive  $\pi$ -electron delocalization across the molecular frameworks, consistent with the conjugated enaminone and fused heterocyclic systems.<sup>43</sup> The HOMO–LUMO gap provides a useful indicator of electronic mobility and chemical reactivity: smaller gaps correspond to higher reactivity and lower kinetic stability, while larger gaps reflect greater hardness and reduced reactivity.<sup>44</sup> In this series, compound **1** exhibits the widest gap (5.067 eV), indicating lower reactivity, whereas compound **9** has the narrowest gap (3.666 eV), consistent with enhanced electron-transfer potential and observed experimental reactivity trends (Table 1).<sup>45</sup>

Selected global descriptors [eqn (1)–(9)] derived from the FMO energies further clarify these trends. Compound **1** shows the highest hardness ( $\eta$ ), while compound **9** is the softest ( $S$ ) and most reactive. Electrophilicity ( $\omega$ ) and maximum charge-transfer capacity ( $\Delta N_{\max}$ ) highlight the electron-accepting ability of the molecules, with compound **9** exhibiting the largest values ( $\omega = 4.903$  eV,  $\Delta N_{\max} = 2.313$ ), consistent with its pronounced electrophilic character. Compound **12** displays the highest electronegativity ( $\chi$ ), indicating strong electron affinity. Fig. 2 summarizes the calculated global reactivity parameters for compounds **1**, **5**, **8–10**, and **12–14**.<sup>46–48</sup>

$$I = -E_{\text{HOMO}} \quad (1)$$

$$Y = -E_{\text{LUMO}} \quad (2)$$

$$\chi = \frac{I + Y}{2} \quad (3)$$

$$\eta = \frac{I - Y}{2} \quad (4)$$

$$S = \frac{1}{\eta} \quad (5)$$

$$\omega = \frac{\chi^2}{2\eta} \quad (6)$$

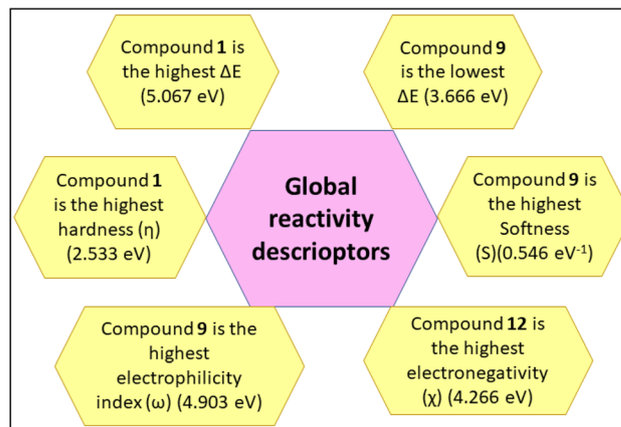


Fig. 2 A graphical representation of the global reactivity indices.

$$\varepsilon = \frac{1}{\omega} \quad (7)$$

$$\mu = -\chi \quad (8)$$

$$\Delta N = -\frac{\mu}{\eta} \quad (9)$$

Finally, in quantum chemical investigations, the total energy of a molecule represents the sum of all electronic and nuclear interactions within the system and serves as a fundamental criterion for evaluating molecular stability.<sup>49</sup> Generally, lower total energy values indicate a more stable configuration, as the system requires less energy to maintain its optimized geometry. In the present theoretical study, the calculated total energies of compounds **3–5** were found to be  $-1101.23$ ,  $-1164.44$  and  $-1873.59$  au, respectively.<sup>50</sup> Comparison of these values shows that compound **5** is significantly more stable than compound **3**, with an energy difference of 772.36 au, and more stable than compound **4**, with a difference of 709.15 au. These theoretical results are consistent with the experimental findings, further confirming the higher stability of compound **5**.<sup>51–53</sup>

**3.2.2. Molecular electrostatic potential (MEP).** Molecular electrostatic potential (MEP) analysis was employed to visualize charge distribution relevant to chemical reactivity and experimental observations, enabling identification of regions

Table 1 Calculated chemical reactivity descriptors of the studied compounds

Compound no.	$E_T$ (au)	HOMO (au)	LUMO (au)	$E$ (HOMO)	$E$ (LUMO)	$I$ (eV)	$A$ (eV)	$\Delta E$ (eV)	$\chi$ (eV)	Pi (eV)	$\eta$ (eV)	$S$ (eV <sup>-1</sup> )	$\omega$ (eV)	$\varepsilon$ (eV <sup>-1</sup> )	$\Delta N$
<b>1</b>	-536.5	-0.247	-0.060	-6.709	-1.643	6.709	1.643	5.067	4.176	-4.176	2.533	0.395	3.442	0.291	1.648
<b>5</b>	-1086.3	-0.228	-0.078	-6.207	-2.117	6.207	2.117	4.090	4.162	-4.162	2.045	0.489	4.235	0.236	2.035
<b>8</b>	-1271.7	-0.217	-0.060	-5.907	-1.628	5.907	1.628	4.279	3.767	-3.767	2.140	0.467	3.317	0.302	1.761
<b>9</b>	-1385.0	-0.223	-0.088	-6.072	-2.407	6.072	2.407	3.666	4.240	-4.240	1.833	0.546	4.903	0.204	2.313
<b>10</b>	-816.9	-0.225	-0.066	-6.125	-1.808	6.125	1.808	4.316	3.966	-3.966	2.158	0.463	3.645	0.274	1.838
<b>12</b>	-910.4	-0.229	-0.084	-6.236	-2.295	6.236	2.295	3.941	4.266	-4.266	1.971	0.508	4.617	0.217	2.165
<b>13</b>	-757.7	-0.233	-0.075	-6.350	-2.046	6.350	2.046	4.304	4.198	-4.198	2.152	0.465	4.095	0.244	1.951
<b>14</b>	-895.4	-0.219	-0.076	-5.965	-2.074	5.965	2.074	3.891	4.020	-4.020	1.945	0.514	4.153	0.241	2.066



susceptible to electrophilic or nucleophilic attack and providing insight into potential intermolecular interactions.<sup>54</sup> The electrostatic potential is represented using a color scale, where positive regions (blue) indicate electron-deficient sites and negative regions (red) correspond to electron-rich areas.<sup>55</sup>

Herein, the MEP maps were used to rationalize the preferred site of nucleophilic attack in the starting chromone derivative and to interpret the electronic features of the resulting enaminone products. For the parent compound **1**, the MEP surface (Fig. 3) shows a pronounced positive potential at the C-2 position,

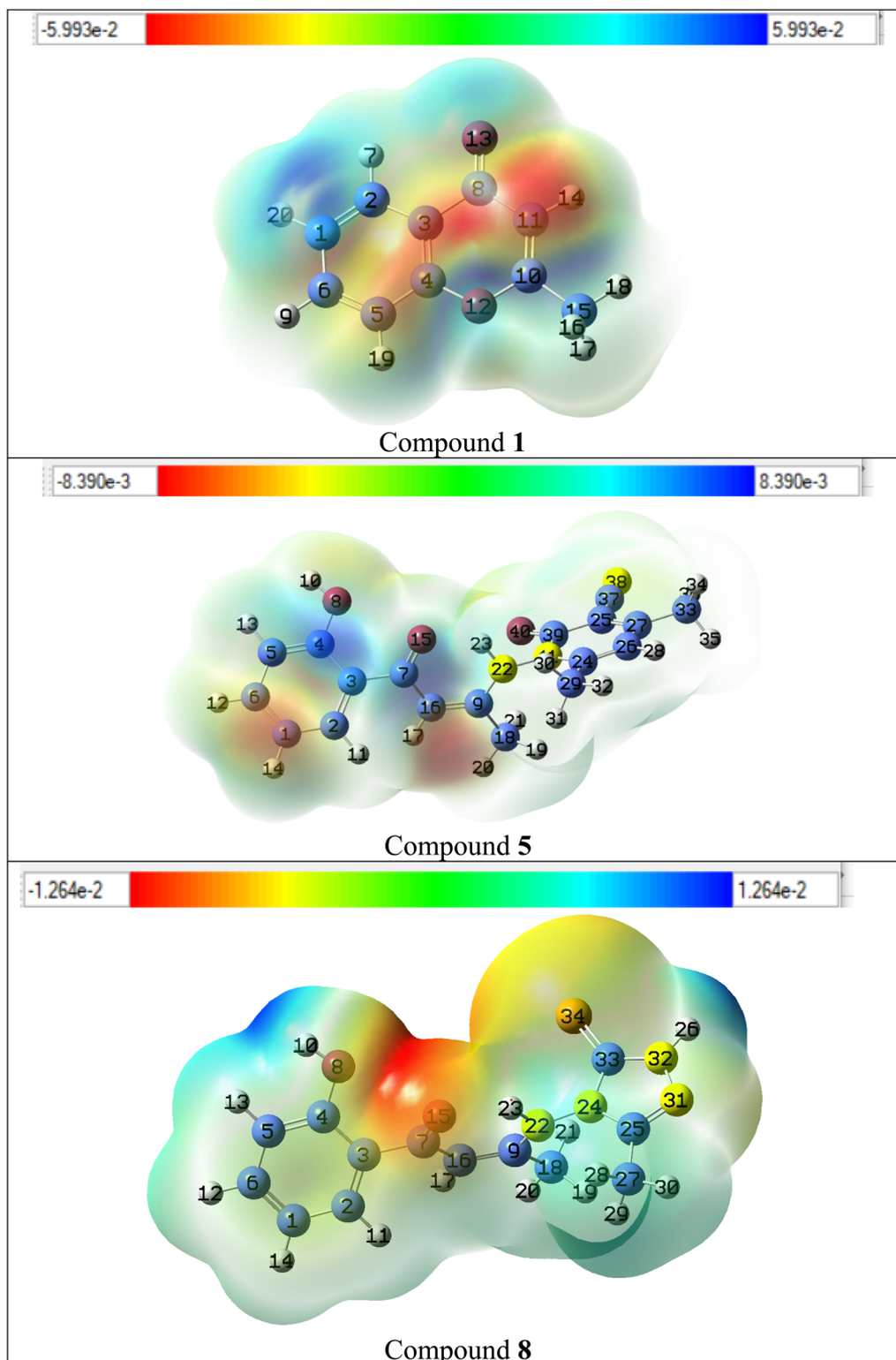


Fig. 3 Molecular electrostatic potential of compounds **1**, **5** and **8**.



indicating its electrophilic character and enhanced susceptibility toward nucleophilic attack. In contrast, the carbonyl carbon at C-4 exhibits a negative potential, consistent with higher electron density at this site. This clear differentiation supports the experimentally observed preference for nucleophilic attack at C-2 followed by ring opening, rather than condensation at the carbonyl group; supporting the experimental results.

In the MEP maps of the representative products **5**, **8**, and **9** (Fig. 3, S3 and S4), the charge distribution is dominated by the enaminone fragment. Electron-rich regions are localized at the C- $\alpha$  positions, while the corresponding C- $\beta$  carbons exhibit electron-deficient character. This polarization of the enaminone system is consistent with conjugation and intramolecular charge delocalization. Importantly, these features correlate well with the experimental NMR data, where C- $\alpha$  carbons resonate upfield relative to the more deshielded C- $\beta$  carbons. As a result, the MEP analysis directly supports the proposed reaction mechanism and provides a coherent electronic explanation for the observed regioselectivity and NMR characteristics.

**3.2.3. FT-IR vibrational analysis.** Both experimental and theoretical approaches were employed to compare the vibrational properties of the synthesized compounds. Theoretical IR spectra were obtained using DFT calculations at the B3LYP/6-311++G(d,p) level of theory.<sup>56</sup> A comparison of experimental and computed FT-IR spectra is shown in Fig. S14, S18, S22, S26, S30, S34 and S38. Because the theoretical calculations were performed in the gas phase, while the experimental measurements were carried out in the solid state, a scaling factor of 0.99 was applied to the calculated frequency values. The experimental and theoretical wavenumbers and the corresponding functional groups assignments are summarized in Tables 2 and 3.

The IR spectra of the synthesized compounds confirmed the presence of characteristic absorption bands consistent with theoretical predictions. For the free OH group, the calculated values appeared at 3410–3433  $\text{cm}^{-1}$ , while the experimental bands were observed at 3402–3416  $\text{cm}^{-1}$ . Similarly, the NH groups showed theoretical absorptions within 3217–3320  $\text{cm}^{-1}$ , compared with experimental values of 3198–3291  $\text{cm}^{-1}$ . The C=O stretching of the enaminone moiety was detected experimentally at 1642–1646  $\text{cm}^{-1}$ , in close agreement with the calculated range of 1650–1660  $\text{cm}^{-1}$ . For the heterocyclic carbonyl groups in compounds **5** and **9–11**, the experimental absorption bands were observed at 1655–1692  $\text{cm}^{-1}$ , whereas the corresponding theoretical values ranged from 1664 to 1705  $\text{cm}^{-1}$ . The C=N group exhibited theoretical stretching frequencies of 1620–1624  $\text{cm}^{-1}$ , slightly higher than the experimental bands at 1608–1615  $\text{cm}^{-1}$ . In compound **8**, the stretching vibration cyano function (C $\equiv$ N) appeared experimentally at 2217  $\text{cm}^{-1}$ , compared to a computed value of 2233  $\text{cm}^{-1}$ . Additionally, the C=C stretching bands were observed in the experimental range of 1566–1587  $\text{cm}^{-1}$ , close to the theoretical values of 1570–1608  $\text{cm}^{-1}$ . The correlation between the experimental and theoretical wavenumbers for these functional groups, illustrated in Fig. 4 and S5–S7, was excellent with a correlation coefficient ( $R^2$ ) of 0.99.

**3.2.4.  $^1\text{H}$  NMR and  $^{13}\text{C}$  NMR spectroscopy.** In recent years, theoretical calculations of  $^1\text{H}$  and  $^{13}\text{C}$  NMR chemical shifts have received considerable attention as a means of verifying the structural and functional features of newly synthesized compounds.<sup>57</sup> Among the available methods, the gauge-including atomic orbital (GIAO) approach is the most widely employed DFT-based technique for NMR shift predictions.<sup>58</sup> In

**Table 2** Experimental and theoretical frequencies and corresponding vibrational assignments of compounds **5**, **8** and **9** at the B3LYP/6-311++G(d,p)

Compound 5			Compound 8			Compound 9		
$\nu_{\text{exp.}}$ ( $\text{cm}^{-1}$ )	$\nu_{\text{the.}}$ ( $\text{cm}^{-1}$ )	Assignment	$\nu_{\text{exp.}}$ ( $\text{cm}^{-1}$ )	$\nu_{\text{the.}}$ ( $\text{cm}^{-1}$ )	Assignment	$\nu_{\text{exp.}}$ ( $\text{cm}^{-1}$ )	$\nu_{\text{the.}}$ ( $\text{cm}^{-1}$ )	Assignment
3406	3415	OH	3409	3424	OH	3402	3410	OH
3279	3283	NH	3248, 3211	3289, 3220	2NH	3237, 3198	3280, 3217	2NH
2217	2233	C $\equiv$ N	1644	1651	C=O <sub>enaminone</sub>	1692	1705	C=O <sub>triazine</sub>
1676	1680	C=O <sub>pyridone</sub>	1613	1620	C=N	1646	1660	C=O <sub>enaminone</sub>
1642	1650	C=O <sub>enaminone</sub>	1587	1595	C=C	1570	1600	C=C
1588	1607	C=C	1223	1244	C=S	1226	1244	C=S

**Table 3** Experimental and theoretical frequencies and corresponding vibrational assignments of compounds **10–13** at the B3LYP/6-311++G(d,p)

Compound 10			Compound 12			Compound 13			Compound 14		
$\nu_{\text{exp.}}$ ( $\text{cm}^{-1}$ )	$\nu_{\text{the.}}$ ( $\text{cm}^{-1}$ )	Assignment	$\nu_{\text{exp.}}$ ( $\text{cm}^{-1}$ )	$\nu_{\text{the.}}$ ( $\text{cm}^{-1}$ )	Assignment	$\nu_{\text{exp.}}$ ( $\text{cm}^{-1}$ )	$\nu_{\text{the.}}$ ( $\text{cm}^{-1}$ )	Assignment	$\nu_{\text{exp.}}$ ( $\text{cm}^{-1}$ )	$\nu_{\text{the.}}$ ( $\text{cm}^{-1}$ )	Assignment
3405	3416	OH	3405	3420	OH	3416	3433	OH	3410	3424	OH
3291	3320, 3304	2NH	1655	1664	C=O <sub>triazine</sub>	1608	1620	C=N	1615	1624	C=N
1659	1672	C=O <sub>pyrazole</sub>	1608	1620	C=N	1587	1592	C=C	1583	1600	C=C
1609	1620	C=N	1566	1570	C=C						
1577	1608	C=C									



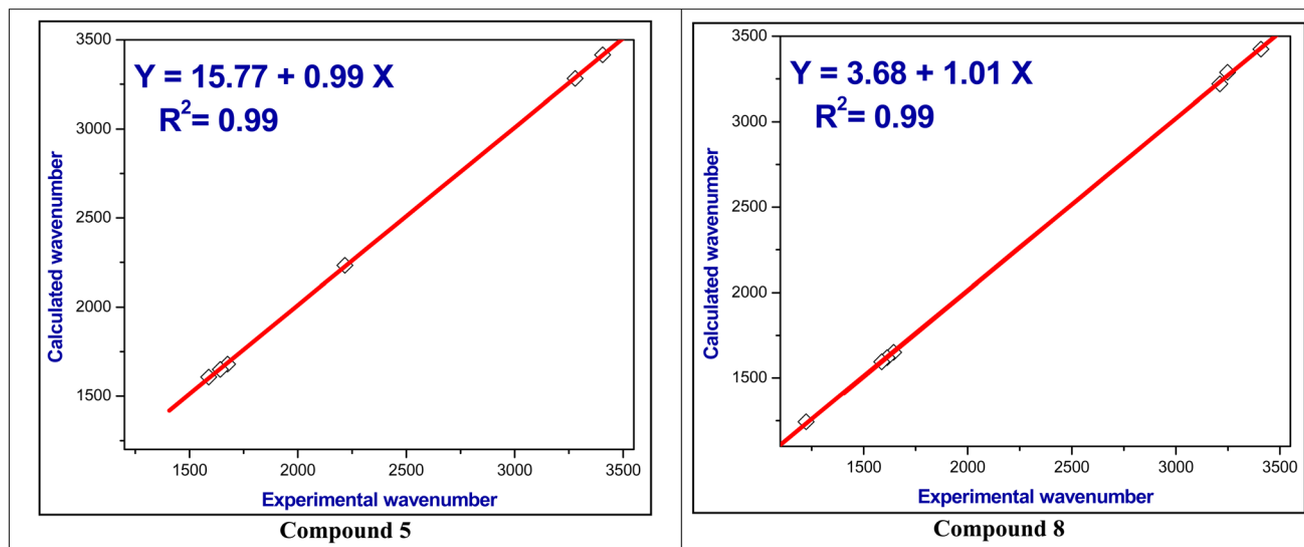


Fig. 4 The correlation relationships of the experimental *versus* calculated IR wavenumbers of compounds 5 and 8.

this study, the chemical shift calculations were performed using Gaussian 09 software at the B3LYP/6-311++G(d,p) level of theory. The computed values were then compared with the experimental NMR data, which were obtained in deuterated DMSO using TMS as the internal standard.

The observed and calculated  $^1\text{H}$  NMR chemical shifts for compounds 5, 8–10, and 12–14 are summarized in Tables 4 and 5, while the spectra (experimental and theoretical) are shown in Fig. S15, S19, S23, S27, S31, S35 and S39. In the upfield region, the  $\text{CH}_3$  protons were experimentally observed as singlets at  $\delta$  2.09–2.40 ppm, whereas the corresponding calculated signals appeared at  $\delta$  1.60–2.87 ppm. Theoretically, three distinct signals were predicted for each methyl group, which may be attributed to differences in the spatial orientation of the protons, leading to slightly varied electronic environments and thus different chemical shifts.

For enaminone derivatives (compounds 5, 8, and 9), the experimental H- $\alpha$  (olefinic) appeared at  $\delta$  6.20–6.32 ppm, closely matching the computed range of  $\delta$  6.35–6.78 ppm. The aromatic protons of the synthesized compounds were recorded at  $\delta$  7.27–8.14 ppm, in good agreement with the calculated range of  $\delta$  6.82–8.09 ppm. In compounds 12–14, the H-5 of the pyrimidine ring was calculated at  $\delta$  8.18–8.83 ppm and experimentally detected at  $\delta$  8.24–8.38 ppm. Similarly, the H-3 of pyridine and triazole moieties in compounds 10 and 13 showed calculated signals at  $\delta$  8.17 and  $\delta$  9.12 ppm, compared with experimental values of  $\delta$  8.37 and  $\delta$  8.18 ppm, respectively. For heteroatom-bound protons, the NH and OH signals were theoretically predicted at  $\delta$  9.21–12.35 and  $\delta$  11.24–12.55 ppm, respectively, while the experimental values were observed at  $\delta$  9.80–11.64 and  $\delta$  11.04–12.36 ppm. The relatively downfield chemical shift of the triazole proton can be attributed to the strong electron-

Table 4 Calculated and experimental  $^1\text{H}$  NMR chemical shifts of compounds 5, 8 and 9 on B3LYP/6-311++G(d,p) basis set

Compound 5			Compound 8			Compound 9		
Atoms	Calculated	Experimental	Atoms	Calculated	Experimental	Atoms	Calculated	Experimental
20-H	1.652009	2.10	20-H	1.641953	2.27	20-H	1.601549	2.18
19-H	1.746833	2.10	19-H	1.757529	2.27	21-H	1.822439	2.18
32-H	2.054319	2.31	21-H	2.144781	2.27	19-H	1.828355	2.18
21-H	2.096794	2.10	29-H	2.180696	2.37	31-H	1.959994	2.40
35-H	2.309049	2.39	30-H	2.241001	2.37	33-H	2.272638	2.40
34-H	2.318255	2.39	28-H	2.264705	2.37	32-H	2.329728	2.40
36-H	2.332206	2.39	17-H	6.778829	6.32	17-H	6.67227	6.21
31-H	2.353999	2.31	13-H	6.835984	7.54	13-H	6.823773	7.4
30-H	2.521359	2.31	14-H	7.040542	7.72	14-H	7.031603	7.55
17-H	6.347872	6.20	12-H	7.470823	7.82	12-H	7.452422	7.84
28-H	6.515025	6.34	11-H	7.551808	8.14	11-H	7.550952	8.06
13-H	6.82574	7.38	26-H	9.40444	9.94	23-H	9.287014	9.81
14-H	7.02641	7.61	23-H	11.06389	10.74	26-H	12.34601	11.65
12-H	7.443389	7.81	10-H	12.5549	11.81	10-H	12.48793	12.36
11-H	7.550339	8.12						
23-H	9.218664	9.80						
10-H	12.52881	12.22						



Table 5 Calculated and experimental  $^1\text{H}$  NMR chemical shifts of compounds 10–13 on B3LYP/6-311++G(d,p) basis set

Compound 10			Compound 12			Compound 13			Compound 14		
Atoms	Calculated	Experimental									
26-H	2.209143	2.09	26-H	2.330597	2.29	27-H	2.303778	2.25	31-H	2.344838	2.26
24-H	2.774632	2.09	20-H	2.409817	2.15	25-H	2.835709	2.25	33-H	2.855805	2.26
25-H	2.847352	2.09	18-H	2.447843	2.15	26-H	2.851774	2.25	32-H	2.869418	2.26
21-H	7.249949	7.27	19-H	2.501868	2.15	22-H	7.139134	7.53	19-H	7.133265	7.29
28-H	7.470519	7.42	24-H	2.782585	2.29	20-H	7.267872	7.35	23-H	7.269587	7.33
20-H	7.680868	7.76	25-H	2.820074	2.29	21-H	7.666869	7.69	25-H	7.302267	7.38
19-H	8.088426	8.00	30-H	6.819517	7.38	19-H	7.711719	7.99	27-H	7.411334	7.40
22-H	8.167953	8.37	28-H	7.198545	7.58	23-H	8.179033	8.38	24-H	7.47455	7.54
14-H	9.845704	9.84	27-H	7.502773	7.82	17-H	8.698514	9.12	22-H	7.671507	7.74
27-H	10.33798	10.72	29-H	7.579888	8.01	18-H	11.75176	11.75	26-H	7.825796	7.83
18-H	12.17287	11.90	32-H	8.7873	8.24				18-H	8.026008	8.08
			31-H	11.83166	11.04				29-H	8.833845	8.37
									28-H	11.23864	11.39

withdrawing inductive and mesomeric effects of the adjacent nitrogen atoms, which increase deshielding.

The  $^{13}\text{C}$  NMR spectra of compounds 5, 8–10, and 12–14 are presented in Fig. S16, S20, S24, S28, S32, S36 and S40, while the theoretical and experimental chemical shift values are summarized in Tables 6 and 7.

As expected, the methyl carbons appeared in the upfield region, with theoretical signals in the range  $\delta$  11.6–26.7 ppm and experimental values between  $\delta$  15.6–19.4 ppm. The aromatic carbons of the benzene rings were observed computationally at  $\delta$  118.4–133.9 ppm, closely matching the experimental range of  $\delta$  121.7–131.5 ppm. While, the C–OH carbons were calculated at  $\delta$  150.0–156.1 ppm, in good agreement with the experimental values at  $\delta$  150.3–152.1 ppm. For compounds 8 and 9, the C=S signals were predicted at  $\delta$  198.6–200.7 ppm, consistent with the experimental resonances at  $\delta$  194.5–195.8 ppm. C- $\alpha$  C- $\beta$ .

In the case of enaminone derivatives (compounds 5, 8, and 9), the C- $\alpha$  resonate theoretically at  $\delta$  101.2–106.7 ppm,

compared to experimental values at  $\delta$  94.2–96.1 ppm, while the C- $\beta$  appeared computationally at  $\delta$  150.2–151.3 ppm, in agreement with experimental signals at  $\delta$  148.1–148.6 ppm. The carbonyl carbons of the enaminone moieties were calculated at  $\delta$  186.3–196.1 ppm, consistent with the experimental values at  $\delta$  182.3–182.7 ppm. The relatively high chemical shift of C- $\beta$  may be attributed to the combined electron-withdrawing mesomeric effect of the carbonyl group and the inductive effect of the adjacent nitrogen atom. Conversely, the lower chemical shift of C- $\alpha$  may result from the electron-donating influence of the NH group, which increases the electron density at these carbons, leading to shielding and reduced  $\delta$  values.

To evaluate the reliability of the theoretical calculations, correlation plots of experimental *versus* calculated chemical shifts for both  $^1\text{H}$  and  $^{13}\text{C}$  NMR are provided in Fig. 5, 6 and S8–S13. The excellent correlation coefficients ( $R^2 = 0.98$ – $0.99$  for both  $^1\text{H}$  and  $^{13}\text{C}$  NMR) confirm the strong agreement between the experimental and theoretical chemical shift values.

Table 6 Calculated and experimental  $^{13}\text{C}$  NMR chemical shifts of compounds 5, 8 and 9 on B3LYP/6-311++G(d,p) basis set

Compound 5			Compound 8			Compound 9		
Atoms	Calculated	Experimental	Atoms	Calculated	Experimental	Atoms	Calculated	Experimental
18-C	18.31434	15.9	27-C	11.65242	15.6	18-C	17.6735	15.7
33-C	21.43412	17.5	18-C	18.83939	16.4	30-C	18.96897	17.5
29-C	21.57802	19.4	9-C	106.7458	96.1	9-C	103.5993	95.9
9-C	101.2112	94.2	5-C	118.593	122.6	5-C	118.4184	122.5
25-C	101.9339	106.2	1-C	122.5349	123.3	1-C	122.5228	124.7
26-C	111.3303	108.3	2-C	125.011	125.1	3-C	128.5142	126.3
37-C	119.7519	116.4	3-C	128.0264	126.4	2-C	130.9209	128.6
5-C	120.4737	122.2	6-C	130.3065	128.6	6-C	135.9791	130.5
1-C	122.4745	124.9	25-C	147.2659	144.3	25-C	148.5132	144.8
3-C	130.063	126.8	16-C	149.7669	148.1	16-C	150.2136	148.6
2-C	131.8666	128.4	4-C	151.2982	150.8	4-C	155.9602	151.7
6-C	132.1563	129.5	7-C	191.1075	182.3	27-C	167.8004	166.5
24-C	133.3026	132.8	33-C	200.7159	194.5	7-C	186.2596	182.7
27-C	144.9511	142.3				24-C	198.6326	195.8
16-C	150.7324	148.2						
4-C	156.1305	152.1						
39-C	163.9764	163.4						
7-C	196.1162	182.6						



Table 7 Calculated and experimental  $^{13}\text{C}$  NMR chemical shifts of compounds 10 and 12–14 on B3LYP/6-311++G(d,p) basis set

Compound 10			Compound 12			Compound 13			Compound 14		
Atoms	Calculated	Experimental									
23-C	19.75128	15.6	17-C	20.36131	16.7	24-C	26.68692	18.4	30-C	20.87679	17.5
13-C	110.2318	112.2	23-C	21.06106	18.5	5-C	119.3801	121.7	17-C	118.3667	122.7
8-C	123.9155	122.3	8-C	109.0970	108.7	3-C	121.6778	123.5	5-C	120.1977	123.2
5-C	124.8377	123.7	5-C	120.3025	122.7	1-C	124.1116	125.9	3-C	122.6261	124.1
1-C	125.7894	125.4	1-C	123.2929	124.3	2-C	128.0574	127.3	16-C	123.383	125.3
3-C	128.5005	126.4	3-C	127.5388	126.4	6-C	129.6411	128.7	1-C	123.992	126.2
6-C	130.9126	128.7	2-C	130.6697	127.2	8-C	133.1513	132.3	20-C	124.5535	126.7
2-C	132.4519	129.9	6-C	132.0601	128.6	10-C	137.9544	140.4	21-C	129.4937	127.2
10-C	145.6349	142.2	7-C	143.1455	142.3	7-C	138.143	140.9	13-C	130.0049	128.5
7-C	146.0137	144.5	14-C	146.4146	144.3	14-C	142.4465	141.2	2-C	131.3841	129.1
11-C	148.2719	146.3	10-C	147.9013	145.8	11-C	144.9029	142.5	6-C	132.465	129.7
4-C	155.9500	151.6	11-C	153.2130	147.4	4-C	155.7184	150.3	15-C	132.6225	130.4
17-C	171.5148	165.4	4-C	154.8726	151.6				8-C	133.2795	131.5
			21-C	172.2610	170.5				10-C	141.4853	140.1
									7-C	142.8493	140.9
									11-C	143.8656	142.2
									4-C	150.0403	150.6

**3.2.5. Nonlinear optical (NLO) properties.** As a complementary outcome of the electronic structure calculations, the nonlinear optical (NLO) properties of the synthesized compounds were evaluated. Organic  $\pi$ -conjugated systems with donor–acceptor characteristics are known to facilitate intramolecular charge transfer, which can give rise to NLO responses.<sup>59,60</sup>

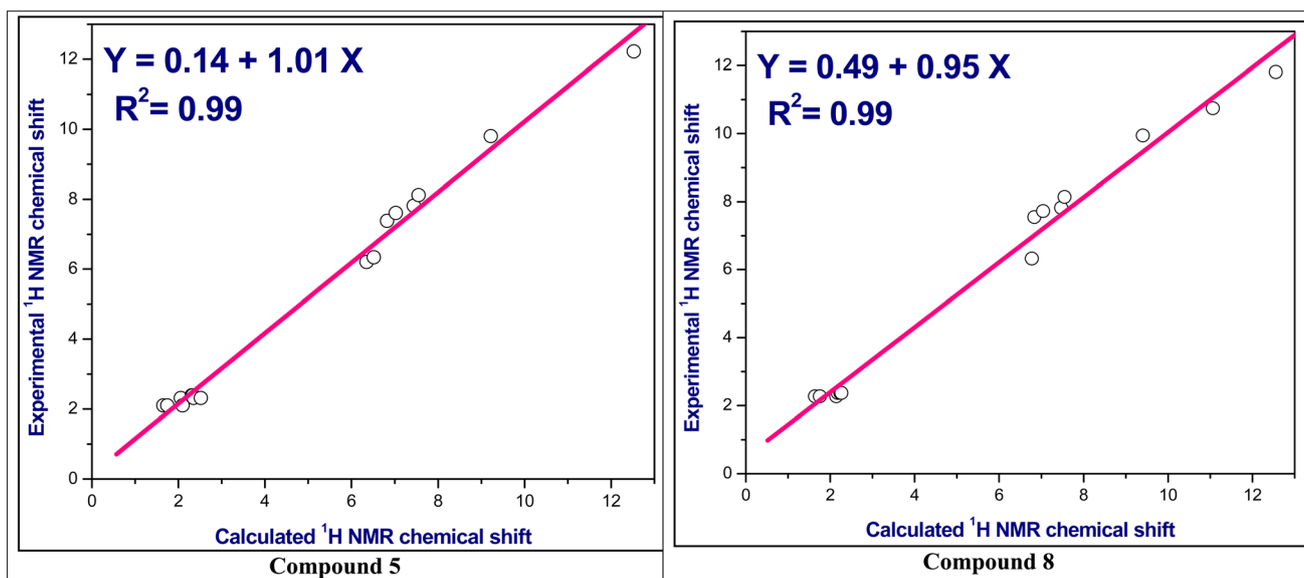
The key NLO parameters, including dipole moment ( $\mu$ ), polarizability ( $\alpha$ ), polarizability anisotropy ( $\Delta\alpha$ ), and total hyperpolarizability ( $\beta_{\text{tot}}$ ), were calculated in the gas phase at the B3LYP/6-311+G(d,p) level and are summarized in Table 8.<sup>61,62</sup> The dipole moments range from 2.884 to 9.792 D, while the polarizabilities fall between  $0.996 \times 10^{-23}$  and  $1.969 \times 10^{-23}$

esu. The calculated  $\beta_{\text{tot}}$  values ( $0.398 \times 10^{-30}$ – $2.358 \times 10^{-30}$  esu) exceed that of urea, a commonly used reference compound, by approximately 1.1–6.3 times. These findings suggest that the synthesized compounds possess promising potential as candidates for NLO material development.<sup>63,64</sup> The principal NLO parameters are evaluated using the following equations:

$$\mu = (\mu_x^2 + \mu_y^2 + \mu_z^2)^{1/2} \quad (10)$$

$$\alpha = (\alpha_{xx} + \alpha_{yy} + \alpha_{zz})/3 \quad (11)$$

$$\Delta\alpha = (2)^{-0.5}[(\alpha_{xx} - \alpha_{yy})^2 + (\alpha_{yy} - \alpha_{zz})^2 + (\alpha_{zz} - \alpha_{xx})^2 + 6(\alpha_{yz})^2 + 6(\alpha_{xy})^2 + 6(\alpha_{xz})^2]^{0.5} \quad (12)$$

Fig. 5 The correlation relationships of the calculated versus experimental  $^1\text{H}$  NMR chemical shifts of compounds 5 and 8.

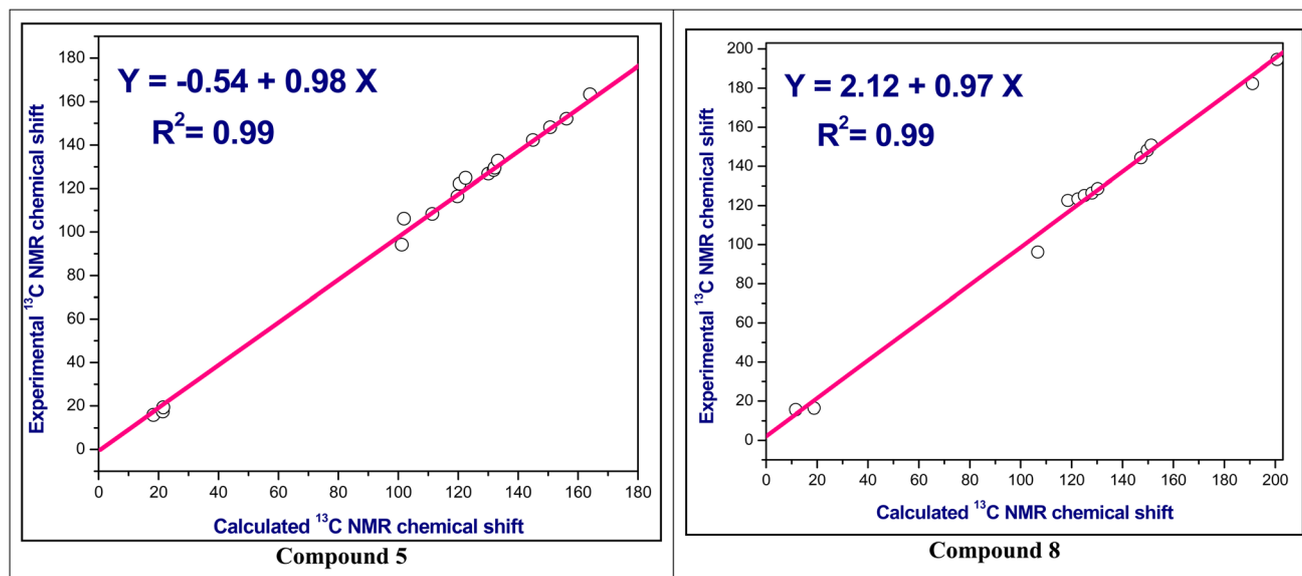


Fig. 6 The correlation relationships of the calculated *versus* experimental  $^{13}\text{C}$  NMR chemical shifts of compounds 5 and 8.

$$\beta_{\text{tot}} = [(\beta_{xxx} + \beta_{xyy} + \beta_{xzz})^2 + (\beta_{yyy} + \beta_{yzz} + \beta_{yxx})^2 + (\beta_{zzz} + \beta_{zxx} + \beta_{zyy})^2]^{0.5} \quad (13)$$

### 3.3. Biological evaluation

**3.3.1. Vitro anticancer assay.** The newly synthesized compounds were evaluated for their anticancer using the MTT colorimetric assay against hepatocellular carcinoma (HepG-2) cell lines after 24 hours of incubation.<sup>32–34</sup> The results demonstrated that all of the tested compounds exhibited growth inhibitory effects on HepG-2 cells (Fig. 7). The cytotoxic activity was quantified in terms of  $\text{IC}_{50}$  values, defined as the concentration ( $\mu\text{M}$ ) required to inhibit 50% of cell growth (Fig. 8, and Table 9). Cisplatin, a clinically established anticancer drug, was used as the reference standard. Overall, the synthesized derivatives displayed a broad spectrum of cytotoxic activity, with  $\text{IC}_{50}$  values ranging from 7.88 to 32.43  $\mu\text{M}$ , indicating activities that varied from strong to moderate potency.

Among the synthesized product, compound 9 displayed the strongest activity ( $\text{IC}_{50} = 7.88 \mu\text{M}$ ), while compounds 5 and 8 showed comparable or slightly better activity than cisplatin

(Table 9). Compounds 12 and 14 also demonstrated notable activity ( $\text{IC}_{50} = 10.66$  and  $8.76 \mu\text{M}$ , respectively). Moderate cytotoxicity was observed for compounds 1, 10, and 13. The observed activity may be influenced by structural features such as conjugated enaminone groups, thioxo substituents, and fused heterocycles.

These results provide initial insight into the cytotoxic effects of the synthesized compounds; however, they are limited to a single cancer cell line (HepG2), and no selectivity data against normal cells were assessed. Therefore, while the compounds show promising growth-inhibitory activity, further studies across multiple cancer and non-cancer cell lines are required to fully evaluate their anticancer potential and therapeutic relevance.

A preliminary linear regression analysis was conducted using the experimental  $\text{IC}_{50}$  values as the dependent variable and selected DFT-derived global reactivity descriptors in the gas phase as independent variables (Table 1). The analysis suggested the following trends:

(1) A reduction in the energy gap ( $E_{\text{gap}}$   $\text{eV}^{-1}$ ) of the studied compounds enhanced their anticancer activity, as reflected by the positive slope of  $\text{IC}_{50}$  *versus*  $E_{\text{gap}}$ , expressed as  $\text{IC}_{50} = -64.55$

Table 8 The dipole moment ( $\mu$ ), mean polarizability ( $\alpha$ ), anisotropy of the polarizability ( $\Delta\alpha$ ) and first-order hyperpolarizability ( $\beta$ ) for the studied compounds

Compound no.	$\mu_x$	$\mu_y$	$\mu_z$	$\mu_{\text{total}}$	$\langle\alpha\rangle$ (au)	$\langle\alpha\rangle$ (esu) $\times 10^{-23}$	$\Delta\alpha$ (au)	$\Delta\alpha$ (esu) $\times 10^{-24}$	$\beta_{\text{total}}$ (au)	$\beta_{\text{total}}$ (esu) $\times 10^{-30}$
1	0.510	-3.885	0.0002	3.919	67.22	0.996	20.77	3.078	46.12	0.398
5	1.304	-6.600	-0.632	6.757	132.86	1.969	66.12	9.799	272.91	2.358
8	-1.164	-3.577	-0.429	3.786	115.61	1.713	32.69	4.844	122.19	1.056
9	-1.349	1.456	2.092	2.884	122.88	1.821	36.01	5.337	133.62	1.154
10	-3.983	-0.047	-0.594	4.027	98.730	1.463	30.58	4.532	131.43	1.136
12	-6.134	2.395	0.982	6.658	112.92	1.674	25.69	3.808	149.79	1.294
13	8.257	5.230	0.591	9.792	98.90	1.466	24.03	3.561	229.73	1.985
14	4.450	-0.796	0.743	4.581	119.79	1.775	20.41	3.025	86.15	0.744



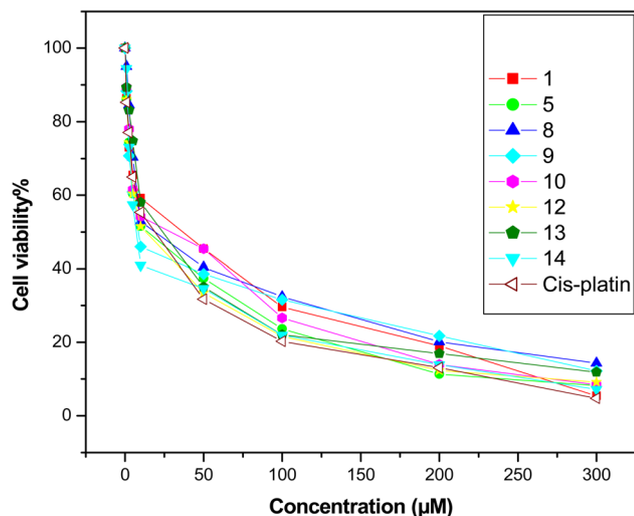


Fig. 7 Relation between cell viability and concentration of all synthesized compounds on the proliferation of HepG2 cell line (cis-platin is the standard drug).

+ 19.53  $E_{\text{gap}}$  eV<sup>-1</sup>,  $r = 0.94$ ,  $n = 8$ . This positive slope suggests that decreasing the energy gap reduces IC<sub>50</sub> values, thereby improving anticancer potential (Fig. 9).

(2) The stability of the compounds, represented by hardness, showed the relation  $\text{IC}_{50} = -64.55 + 39.07 \eta$  eV<sup>-1</sup>,  $r = 0.94$ ,  $n = 8$ . The positive slope indicates that lowering hardness decreases IC<sub>50</sub>, which enhances inhibitory activity (Fig. 9). Conversely, softness displayed an inverse relationship with IC<sub>50</sub>. This was confirmed by the negative slope of softness against IC<sub>50</sub>, expressed as  $\text{IC}_{50} = 105.73 - 183.75 S$  eV<sup>-1</sup>,  $r = 0.94$ ,  $n = 8$  (Fig. 9).

These observations suggest that smaller energy gaps and lower hardness may correlate with higher cytotoxic activity. However, the dataset is limited ( $n = 8$ ), and therefore, the statistical reliability of these correlations is low. Consequently, these trends should be considered exploratory and interpreted cautiously rather than as definitive predictors of biological activity.

Table 9 IC<sub>50</sub> values of the prepared compounds against hepatocellular carcinoma cells lines (HepG-2) for 24 hours

Compounds no.	(HepG-2 cells) IC <sub>50</sub> (µM L <sup>-1</sup> )
1	32.43 ± 1.44
5	14.65 ± 0.88
8	17.17 ± 0.99
9	7.88 ± 0.52
10	26.60 ± 1.17
12	10.66 ± 0.79
13	20.94 ± 1.04
14	8.76 ± 0.67
Cisplatin	17.86 ± 0.93

### 3.3.2. Drug-likeness and *in silico* ADME anticipation.

SwissADME is a widely used computational tool for predicting absorption, distribution, metabolism, and excretion (ADME) parameters, as well as evaluating drug-likeness based on physicochemical properties.<sup>65</sup> It provides insight into whether a compound is likely to exhibit favorable oral bioavailability. Compounds that violate more than one of the commonly applied rules—Lipinski's,<sup>66</sup> Veber's,<sup>67</sup> Egan's,<sup>68</sup> or Ghose's<sup>69</sup>—are generally considered less suitable for oral administration.

The results indicated that all studied compounds complied with Lipinski's rule of five, suggesting favorable oral bioavailability. Specifically, molecular weights ranged from 160 to 323 g mol<sup>-1</sup>,  $M \log P$  values from 0.51 to 2.85, hydrogen bond donors between 0 and 3, and hydrogen bond acceptors within the acceptable range (Table 10).

Veber's rule, which emphasizes total polar surface area (TPSA ≤ 140 Å<sup>2</sup>) and rotatable bonds (<10), was satisfied, with TPSA values of 30–132 Å<sup>2</sup> and an acceptable number of rotatable bonds. Egan's rule (TPSA ≤ 131.6 Å<sup>2</sup> and WLOGP ≤ 5.88) and the Ghose filter (molecular weight 160–323 g mol<sup>-1</sup>, molar refractivity 47–91, WLOGP 1.45–3.56, and total atom count 20–70) were also fully satisfied (Table 10). Collectively, compliance with these rules indicates favorable physicochemical properties and drug-likeness, supporting the potential for oral activity.

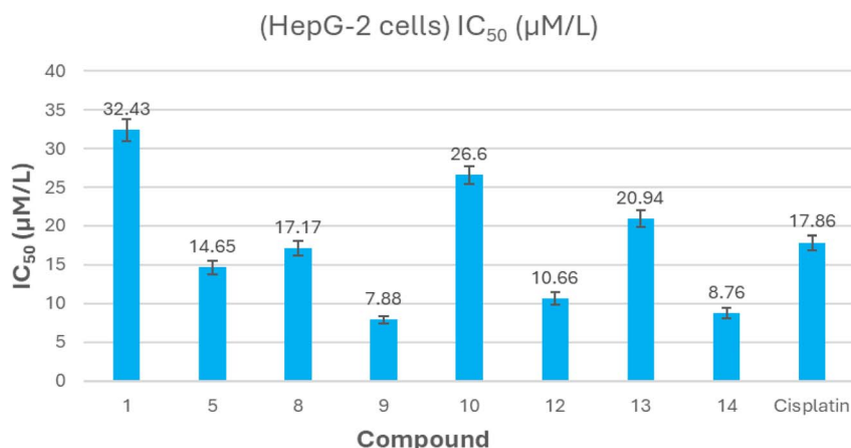


Fig. 8 Virtual IC<sub>50</sub> values of the target compounds and cis-platin as a standard drug against HepG-2 cell lines.



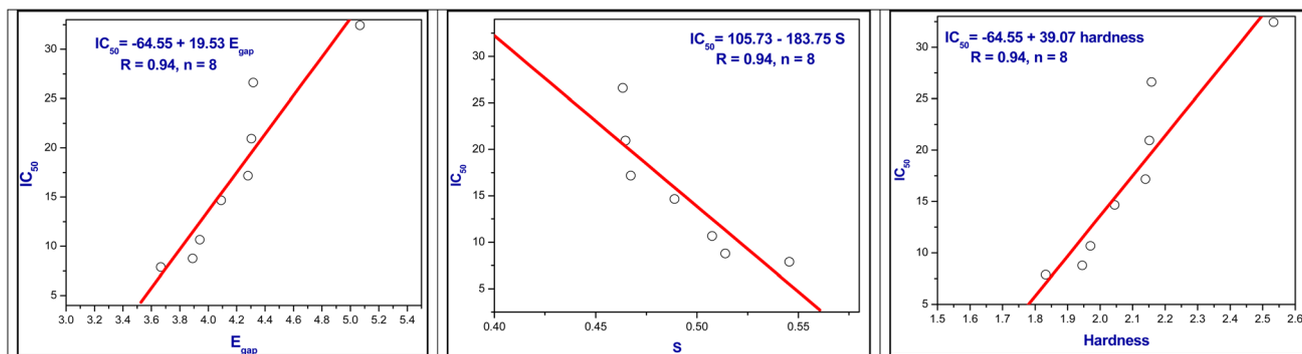


Fig. 9 Linear correlation between  $IC_{50}$  versus energy gap, hardness and softness.

The bioavailability radar provides a visual assessment of six key physicochemical parameters: lipophilicity (XLOGP3  $-0.7$  to  $+5.0$ ), size (MW  $150$ – $500$  g mol $^{-1}$ ), polarity (TPSA  $20$ – $130$  Å $^2$ , except compound 9), solubility ( $\log S \leq 6$ ), saturation (fraction Csp $^3$   $0.25$ – $1$ ), and flexibility ( $\leq 9$  rotatable bonds). As shown in Fig. 10 and S42, all compounds met these criteria, with minor deviations observed for saturation (INSATU). Overall, the *in silico* ADME and drug-likeness results support the potential oral bioavailability of the studied compounds at a screening level. However, these predictions are preliminary and should be interpreted with caution, as they do not replace experimental pharmacokinetic or metabolism studies.

**3.3.3. Molecular docking.** Docking simulations are a widely used tool in drug discovery to predict interactions between small molecules and target proteins, providing insight into potential mechanisms of action.<sup>70–73</sup> In this study, cyclin-dependent kinase 1 (CDK1) was selected as a relevant anti-cancer target because it plays a central role in regulating the G2/M phase of the cell cycle and is often overexpressed in hepatocellular carcinoma and other cancers.<sup>38,39</sup>

The docking protocol was validated by re-docking the co-crystallized ligand, [(2,6-difluorophenyl)carbonyl]amino)-*N*-(4-fluorophenyl)-1*H*-pyrazole-3-carboxamide, into the CDK1 active site (PDB ID: 4Y72). The re-docked pose closely reproduced the

Table 10 Estimations of ADME and physicochemical properties of the studied compounds

Compound no.	MW	MLOGP	HBA	HBD	TPSA	nRotB	WLOGP	XLOGP3	Log <i>S</i>	Csp3	Fraction MR	Lipinski violations	Veber violations	Egan violations	Ghose violations
1	160.17	1.14	2	0	30.21	0	2.10	2.27	$-2.54$	0.10	47.45	0	0	0	0
5	323.35	1.49	4	2	95.12	4	2.18	2.98	$-4.64$	0.17	91.27	0	0	0	0
8	290.34	0.75	3	3	115.03	4	2.09	2.35	$-4.41$	0.15	78.30	0	0	0	0
9	318.35	0.51	4	3	132.10	4	1.45	2.23	$-4.64$	0.14	84.57	0	0	1	0
10	241.25	1.80	3	3	81.77	1	1.93	2.09	$-3.44$	0.08	69.14	0	0	0	0
12	268.27	1.62	5	1	80.38	1	1.47	0.85	$-2.12$	0.14	74.44	0	0	0	0
13	226.23	1.60	4	1	63.31	1	1.81	2.23	$-3.19$	0.08	63.21	0	0	0	0
14	275.30	2.85	3	1	50.42	1	3.56	3.43	$-4.17$	0.06	82.92	0	0	0	0

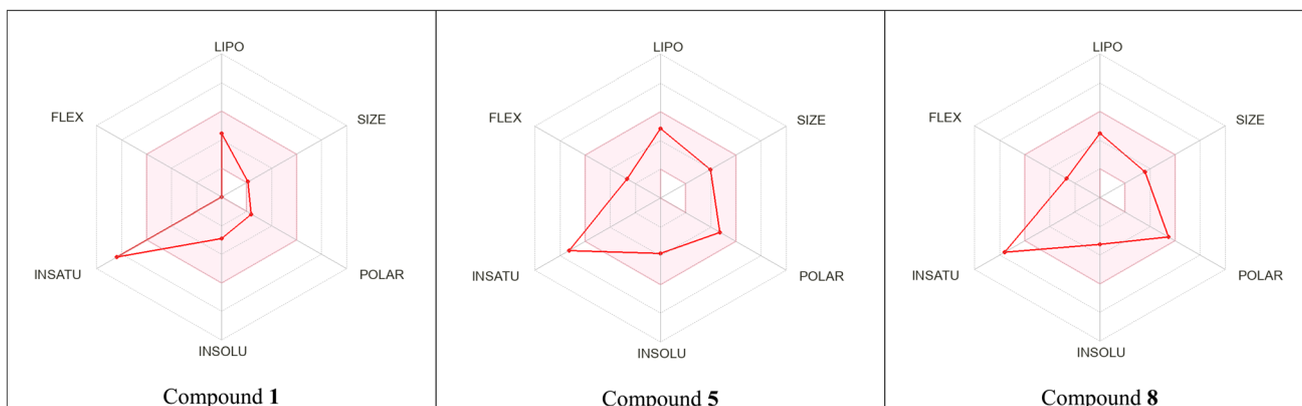


Fig. 10 The bioavailability radars of compounds 1, 5 and 8.



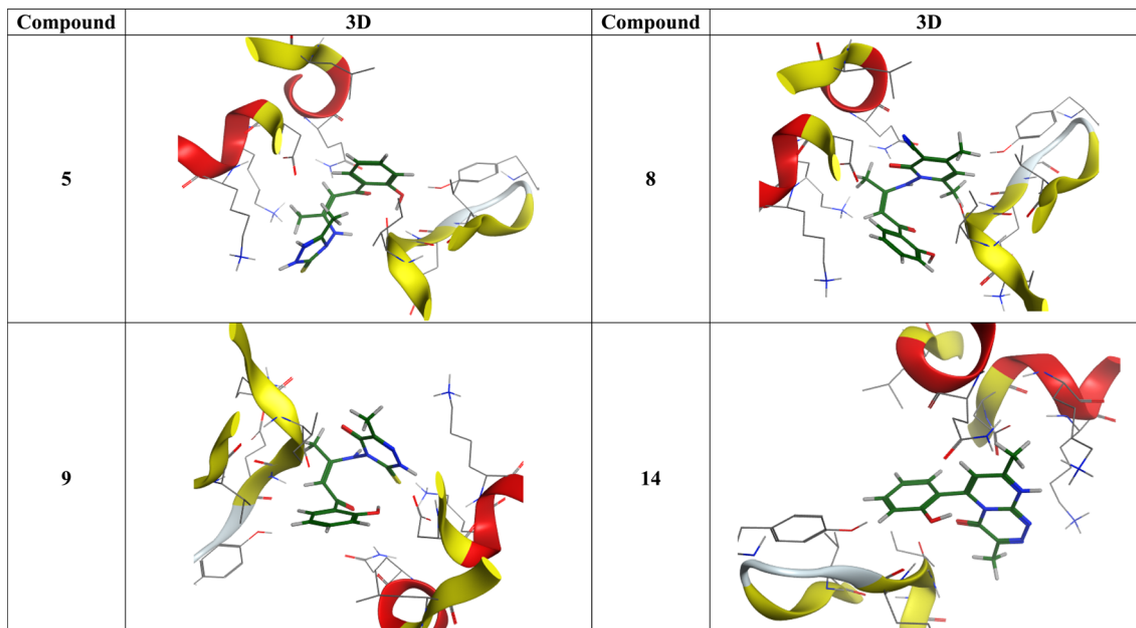


Fig. 11 3D representation of the hydrogen bonding between the studied compounds (5, 8, 9 and 14) with the amino acids residues of the target protein (PDB ID: 4Y72).

Table 11 Various interactions between the studied compounds 5, 8, 9, and 14 with the amino acids residues of the target protein (PDB ID: 4Y72)

Receptor	Distance	Interaction type	Receptor	Distance	Interaction type
<b>Compound 5 (binding energy <math>-8.3 \text{ kcal mol}^{-1}</math>)</b>			<b>Compound 8 (binding energy <math>-8.1 \text{ kcal mol}^{-1}</math>)</b>		
TYR15	1.94	H-bond	VAL18	3.82	Alkyl
LYS88	4.36	Alkyl	LYS88	3.38	Alkyl
<b>Compound 9 (binding energy <math>-8.6 \text{ kcal mol}^{-1}</math>)</b>			<b>Compound 14 (binding energy <math>-8.5 \text{ kcal mol}^{-1}</math>)</b>		
ASP86	2.18	H-bond	GLU12	2.33	H-bond
LYS89	5.31	Pi-alkyl	ASP86	3.85	Pi-anion
			GLY11	3.93	Pi-Sigma
			VAL18	5.05	Pi-alkyl

crystallographic binding mode (Fig. S43), confirming the reliability of the docking methodology.<sup>74</sup>

The synthesized compounds (1, 5, 8–10 and 12–14) were docked into the CDK1 active site, and their interactions were analyzed (Fig. 11, S44, Tables 11, and S1). Across the series, recurring interaction motifs were observed, including hydrogen bonding with GLU12, ASP86, or TYR15, and Pi-anion or Pi-alkyl interactions with residues such as LYS88 and LYS89. Compounds showing the most favorable binding energies (*e.g.*, 9, 14, and 5) generally engaged in multiple hydrogen bonds and Pi-anion interactions, suggesting these contacts are key contributors to binding affinity.

A comparison of docking results with *in vitro* cytotoxicity (HepG2  $IC_{50}$  values) suggests a general trend in which stronger binding energies correspond to lower  $IC_{50}$  values. For example, compounds 9 and 14, which exhibited the most favorable binding energies ( $-8.6$  and  $-8.5 \text{ kcal mol}^{-1}$ , respectively), also displayed the highest cytotoxic activity ( $IC_{50} = 7.88$  and  $8.76 \mu\text{M}$ , respectively). Conversely, compounds with weaker binding

energies, such as 1 ( $-7.6 \text{ kcal mol}^{-1}$ ), showed higher  $IC_{50}$  values ( $32.43 \mu\text{M}$ ). Although this trend is consistent with the proposed mode of action, it should be interpreted cautiously, as *in vitro* activity may also be influenced by other pharmacokinetic or cellular factors not captured in docking simulations.

Overall, these results indicate that hydrogen bonding and Pi-anion interactions are recurring motifs likely contributing to the observed cytotoxicity, supporting CDK1 as a plausible target. The docking data complement the experimental findings but are presented as a preliminary theoretical evaluation rather than definitive proof of mechanism.

## 4. Conclusions

In conclusion, the chemical reactivity of 2-methylchromone (1) with different nitrogen-based nucleophiles enabled the synthesis of *Z*-configured enaminones and a series of novel pyrimidine fused heterocycles, highlighting its utility as a versatile precursor in heterocyclic construction. In addition,



computational studies at the B3LYP/6-311++G(d,p) level revealed strong agreement between calculated and experimental IR vibrational frequencies and NMR ( $^1\text{H}$  and  $^{13}\text{C}$ ) chemical shifts, with correlation coefficients ( $R^2$ ) = 0.99. FMO analysis and global reactivity descriptors indicated that compound **9** possesses the highest softness ( $S = 0.546$  eV), compound **12** showed the greatest electronegativity ( $\chi = 4.266$  eV), and compound **9** exhibited the strongest electrophilicity index ( $\omega = 4.903$  eV). The MEP mapping identified negative potential regions around sulphur, oxygen and nitrogen atoms, favoring electrophilic attack, and positive regions around hydrogen and carbon atoms, favoring nucleophilic attack. The first hyperpolarizability values ( $0.398 \times 10^{-30}$ – $2.358 \times 10^{-30}$  esu) suggest promising non-linear optical (NLO) activity. Biologically, compounds **9** and **14** displayed notable anticancer activity compared with cisplatin. Moreover, the binding energy computations have confirmed the experimental data in anticancer evaluation. Further, drug-likeness evaluation showed that all synthesized compounds comply with Lipinski's, Ghose's, Egan's, and Veber's rules, supporting their potential for oral administration.

## Conflicts of interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

## Data availability

Data are available upon request from the authors.

Supplementary information (SI) is available. See DOI: <https://doi.org/10.1039/d5ra09088a>.

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