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Mechanistic insights into carbonic anhydrase IX inhibition by coumarins from *Calendula officinalis*: *in vitro* and *in silico* approaches[†]

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Given the critical role of carbonic anhydrase IX (CA IX) in various pathological conditions, there is a significant demand for new inhibitors to enhance patient outcomes and clinical management. In this study, we examined the inhibitory effectiveness of five coumarins derived from Calendula officinalis against CA IX using in vitro assays and computational modeling. Among the coumarins tested, xeroboside and isobaisseoside were identified as the most potent inhibitors. Kinetic studies indicated that xeroboside and isobaisseoside exhibit a mixed inhibition mode. Molecular docking analysis showed that the tested coumarins exhibit binding affinities and extensive polar interactions with CA IX. These coumarins demonstrated significant hydrophobic interactions and occupied the same binding site as acetazolamide (AAZ). Molecular dynamics (MD) indicated that xeroboside and isobaisseoside exhibited consistent trajectories and notable energy stabilization during their interaction with CA IX. MM/PBSA calculations showed that xeroboside displayed the lowest binding free energy (-27.26 ± 2.48 kJ mol⁻¹). Potential Energy Landscape (PEL) analysis revealed distinct and stable conformational states for the CA IX-ligand complexes, with xeroboside exhibiting the most stable and lowest energy configuration. These computational findings are consistent with the experimental results, highlighting the potential efficacy of xeroboside and isobaisseoside as CA IX inhibitors. In conclusion, Calendula officinalis-derived coumarins are promising candidates as effective CA IX inhibitors.

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1. Introduction

Carbonic anhydrases (CAs) are a family of zinc metalloenzymes that catalyze the reversible hydration of carbon dioxide to bicarbonate and a proton, a reaction crucial for maintaining acid-base balance in various tissues and organs.¹ These

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enzymes are implicated in numerous physiological processes including respiration, acid-base homeostasis, and ion transport.1 Among the known isoforms in humans, CA IX is of particular interest due to its overexpression in several types of tumors, particularly under hypoxic conditions.² CA IX plays a vital role in the regulation of intracellular and extracellular pH in cancer cells, facilitating their survival, proliferation, and invasion.3 The inhibition of CA IX has thus gained significant attention as a therapeutic strategy, aiming to disrupt the pH regulatory mechanism of cancer cells, thereby inhibiting tumor growth and metastasis.⁴ CA IX is remarkably expressed in cells with high growth and glycolysis rates and cells frequently exposed to acidic and hypoxic environments, such as cancer and endothelial cells (ECs).5 In the endothelium, the role of CA IX extends beyond pH regulation to aerobic glycolysis, EC migration and network formation.6 Targeting CA IX with specific inhibitors could lead to the development of novel anticancer therapies with reduced side effects compared to conventional treatments. In addition, CA IX inhibition might influence ECs and integrity of barriers in different tissues.

Calendula officinalis (*C. officinalis*), commonly known as marigold, is a well-known medicinal plant with diverse phytochemical composition and broad spectrum of biological activities.⁷ The plant is rich in secondary metabolites, including

flavonoids, triterpenoids, carotenoids, and essential oils, which contribute to its anti-inflammatory, antioxidant, antimicrobial, and anticancer properties.8-11 Among these compounds, coumarins are notable for their therapeutic potential.¹² C. officinalis has been documented to exhibit different pharmacological effects,¹¹ making its phytoconstituents, including coumarins prime candidates for further investigation in enzyme inhibition studies. The selection of coumarins from C. officinalis for studying CA IX inhibition is justified by the plant established use in traditional medicine,11 and previous studies indicating the effectiveness of coumarins in inhibiting various enzymes and pathways involved in cancer progression.13 Therefore, exploring the inhibitory effects of these natural compounds on CA IX could provide valuable insights into novel anticancer strategies and enhance the therapeutic applications of C. officinalis.

To comprehensively assess the inhibitory activity of coumarins on CA IX, we employed an integrated approach combining *in vitro* assays and *in silico* studies. The *in vitro* assays provide direct evidence of the potency of the isolated phytoconstituents. Complementing these experiments, molecular docking and molecular dynamics (MD) simulations offer a detailed understanding of the interaction mechanisms at the molecular level.¹⁴ Docking studies help predict the binding affinity and orientation of coumarins within the active site of CA IX, while MD simulations reveal the stability and dynamic behavior of the enzyme–inhibitor complexes over time. This integrated approach not only validates the inhibitory activity observed experimentally but also elucidates the underlying molecular interactions, paving the way for the development of more effective CA IX inhibitors.

2. Materials and methods

2.1. Phytochemical investigation

2.1.1. General. NMR spectral analysis, including ¹H NMR (500 MHz) and (125 MHz) ¹³C NMR, of the isolated coumarins was performed using Bruker AM-500 spectrometer. The optical density polarimeter was Rudolph Autopol III. UV-vis spectral analysis was assessed using the Shimadzu UV–Vis 160i spectrophotometer. HREIMS EIMS spectra were obtained using the Finnigan MAT TSQ 700 mass spectrometer. FTIR spectra of the isolated compounds were obtained utilizing the Shimadzu FTIR-8400 instrument with KBr pellets.

2.1.2. Plant collection and isolation of phytochemicals. The aerial parts of *C. officinalis* were obtained from local private gardens where it is commercially grown. The plant species was identified by expert taxonomists. The powdered dried aerial parts of the plant (4.5 kg) was exhaustively extracted by cold maceration using methanol (MeOH) for three days. A portion of the obtained extract (250 g) was suspended in water and successively extracted with petroleum ether, ethyl acetate (EtOAc), and *n*-butanol (1.1 L × 3, each). The EtOAc soluble fraction (70 g) underwent chromatographic fractionations using a silica gel column, elution started with the solvent system dichloromethane (DCM)-EtOAc-toluene (7.5:1.51:1) followed by the DCM-EtOAc (8:2), and MeOH to produce 56 fractions.

These fractions were collected and combined based on their TLC profile into six main fractions (A1-A6). Fraction A3 was further chromatographed over a silica gel column eluted with dioxane-hexane (4.5:5.5) to afford five main subfractions after TLC comparison and combinations (A3.1-A3.5). Subfraction A3.1 underwent spontaneous crystallization to afford the pure form of compound 2 (26 mg). Subfraction A3.3 was subjected to chromatographic purification over Sephadex LH-20 column eluted with MeOH to produce the purified compounds 4 (31 mg) and 5 (19 mg). The fraction A4 revealed an interesting TLC profile and was subjected to fractionation using a silica gel column using chloroform-MeOH solvent (90:10) of increasing polarity. A total of six main subfractions were obtained after TLC investigation and combination (A4.1-A4.6). Subfractions A4.2 and A4.3 were purified over the Sephadex LH-20 column eluted with MeOH to afford compound 1 (28 mg) and compound 3 (32 mg), respectively. The purity of the isolated compounds was confirmed by spectroscopic analysis.

2.2. In vitro CA IX inhibition assay

The effect of coumarins on CA IX activity conducted as previously described with minor modification,15,16 using 4-nitrophenyl acetate (4-NPA) as a substrate. A buffer solution containing HEPES and Tris-HCl (20 mM) was used as the reaction medium. Acetazolamide (AAZ), a clinically used CA inhibitor, was selected as the standard drug for this study. Compounds 1-5 and AAZ were prepared in DMSO, diluted in the assay buffer to different concentrations, and then utilized in the experiments. The reaction mixture consisted of 150 µL of the prepared reaction medium, 20 µL of the aqueous CA IX solution (0.1 mg mL⁻¹ in deionized water), 20 µL of 4-NPA (0.7 mM in ethanol), and 20 µL of the tested compounds. The production of the yellow-colored 4-nitrophenolate was monitored at 400 nm and the experiment was performed in triplicate. The kinetic analysis of the CA IX inhibition was performed using varying concentrations of the tested compounds and 4-NPA. Following substrate addition, the change in absorbance was monitored every minute for 30 minutes at 25 °C. To elucidate the inhibition mechanism, a Lineweaver-Burk plot was constructed by plotting the reciprocal of the substrate concentration (1/[S]) against the reciprocal of the reaction velocity (1/ V_{max}). The inhibition constant (K_i) was determined by creating a double reciprocal plot of the inhibitor concentration ([I]) against the slope of the Lineweaver-Burk plot.

2.3. Statistical analysis

The results are presented as the mean \pm standard deviation (SD), with each data point reflecting the average of three independent experiments. Comparison of the IC₅₀ was conducted using one-way ANOVA followed by Tukey's test on GraphPad Prism 8.0. A *P* value <0.05 was considered significant.

2.4. In silico investigations

2.4.1. System preparation. The 3D crystal structure of human CA IX (PDB ID: 5FL4) was obtained from the Protein Data Bank (PDB). The enzyme was checked for missing residues

using the Swiss-PdbViewer software.¹⁷ UCSF Chimera 1.17.1 was utilized for molecular visualization, removal of nonstandard residues and water molecules, and stripping out the native ligand.¹⁸ The geometrical structures of investigated coumarins were fully optimized using the DFT approach at the B3LYP level using the 6-311G (d, p) basis set.^{19–21} Frequency calculations were also implemented to confirm the absence of imaginary frequencies for the obtained energy minima states. The Gaussian 16 suite was employed for executing all DFT calculations in this investigation.²²

2.4.2. Molecular docking analysis. The 3D structure of CA XI was prepared for docking using Autodock Tools (ADT) v1.5.6 software.²³ This preparation involves the addition of polar hydrogens and adjusting the grid box size and dimension to involve the native ligand active site amino acid residues.²⁴ The open-source AutoDock Vina v1.5.6 package was utilized to perform the molecular docking calculations of coumarins–CA IX complexes.²³

2.4.3. MD simulations. By employing molecular docking calculations, we selected the C. officinalis coumarins complexes with CA IX (PDB ID: 5FL4) that demonstrated the highest binding affinities. These complexes were then subjected to further analysis using 100 ns MD simulations conducted with GROMACS 2022.4,^{25,26} utilizing GPU acceleration to enhance performance and efficiency. Specifically, the simulations were performed on a high-performance computing cluster equipped with Intel Xeon E5-2680 v3 CPUs running at 2.5 GHz, along with NVIDIA Tesla K80 GPUs (2 GPUs per node) to speed up the calculations and running on the CentOS Linux 7 operating system. The process was initiated by extracting various coumarin molecules from their complex structures with CA IX. These structures were prepared for subsequent simulations using the pdb2gmx tool. For these simulations, all-atom CHARMM36m force field and the CHARMM-modified TIP3P water model were selected.27,28 Geometric parameters and topologies for the various coumarins were generated using the CGenFF online server (https://cgenff.com/). These parameters were then integrated into the full topology of the processed CA IX. Following the formation of the complexes, both the unbound CA IX and the inhibitor-CA IX complexes were solvated by placing them in a dodecahedron-shaped box with a final volume of 289.24 nm³. To ensure electrical neutrality, seven sodium counter ions were added.28 An energy minimization step using the steepest descent method was performed for 10 ps to eliminate any unfavorable interactions.²⁹ The system was equilibrated in two phases using both NVT (constant volume, constant temperature) and NPT (constant pressure, constant temperature) ensembles for 100 ps at 300 K.³⁰ Finally, the production phase of the MD simulation was conducted for 100 ns at a pressure of 1 bar and a temperature of 300 K.

2.4.4. MM/PBSA analysis. The Molecular Mechanics/Poisson-Boltzmann Surface Area (MM/PBSA) approach was employed to gain comprehensive insight into ligand-receptor interactions. This method integrates molecular mechanics (MM), Poisson-Boltzmann (PB) electrostatics, and Solvent Accessibility (SA) models to estimate binding energies. Renowned for its precision and effectiveness, MM/PBSA is

frequently utilized to calculate the binding free energies of noncovalently bound complexes, as corroborated by numerous studies.^{31,32} In our research, we utilized the gmx_MPBSA tool to calculate the binding free energies derived from the outputs of MD simulations.³³ This tool enabled us to assess the energetic contributions of different interactions within the enzymeinhibitor complexes, providing a detailed understanding of the binding mechanisms involved.

Results and discussion

3.1. Phytochemical study

The phytochemical investigation of the aerial parts of *C. officinalis* led to the isolation and identification of five coumarins. The chemical structures of these isolated compounds (Fig. 1) were elucidated through spectroscopic analysis, comparison of TLC profiles with known standards, and referencing existing scientific literature. Using these methods, the coumarins were identified as 6,7-di-*O*-glucopyranosyl esculetin (1),³⁴ aesculin (2),³⁵ isobaisseoside (3),³⁶ skimmin (4),³⁷ and xeroboside (5).³⁸

3.2. Inhibitory activity of *C. officinalis*-derived coumarins on CA IX

Given the involvement of CA IX in different physiological and pathological processes, including its overexpression in hypoxic tumor environments and its contribution to tumor growth and survival,^{1,2} there is a pressing need to identify effective inhibitors to modulate its activity. The dose-response curves and IC₅₀ values for the isolated coumarins and AAZ on CA IX activity are presented in Fig. 2. As shown in Fig. 2A, the compounds and AAZ showed a concentration-dependent inhibition of CA IX activity. The IC_{50} values (Fig. 2B), indicating the concentration at which 50% inhibition of CA IX activity is achieved, varied among the compounds. Xeroboside demonstrated the most potent inhibition with an IC₅₀ value of 85.34 \pm 3.01 nM, following AAZ (IC₅₀ = 42.87 \pm 1.39 nM). Isobaisseoside also showed potent inhibition with an IC₅₀ of 247.61 \pm 7.27 nM, and aesculin and skimmin displayed moderate inhibition with IC₅₀ values of 327.05 \pm 11.46 nM and 547.93 \pm 35.66 nM, respectively. The least potent inhibitor was 6,7-di-O-glucopyranosyl esculetin, with an IC_{50} value of 1.01 \pm 0.12 $\mu M.$ Statistical analysis revealed significant differences between the IC₅₀ value of compounds 1-4 when compared with either AAZ or compound 5. Also, a non-significant difference was observed between IC₅₀ value AAZ and compound 5 (Fig. 2B).

To further characterize the inhibitory effects of isobaisseoside and xeroboside on CA IX, kinetic analyses were performed, and the Michaelis–Menten (Fig. 2C and E) and Lineweaver–Burk (Fig. 2D and F) plots were constructed. The Michaelis–Menten plot for isobaisseoside (Fig. 2E) showed a decrease in the maximum reaction velocity (V_{max}) with increasing concentration ($K_i = 126$ nM), suggesting a mixed-type inhibition. The Lineweaver–Burk plot (Fig. 2F) corroborated this, as the lines intersect to the left of the *y*-axis, indicating changes in both V_{max} and K_m . Similar to isobaisseoside, xeroboside exhibited a mixed-type inhibition pattern (Fig. 2C and D). The decrease in



 V_{max} with increasing concentration was evident from the Michaelis–Menten plot ($K_i = 77.2$ nM). The Lineweaver–Burk plot showed intersecting lines, confirming the mixed inhibition. The reference inhibitor AAZ demonstrated non-competitive inhibition.

The efficacy of the tested coumarins against CA IX demonstrates the potential of these natural products as therapeutic agents. Xeroboside and isobaisseoside exhibited the highest inhibition activity among the tested coumarins and the superior activity of xeroboside can be attributed to its lower IC_{50} and higher inhibition rate compared to other coumarins. The structural features of xeroboside, including the presence of specific functional groups, may contribute to its enhanced binding affinity and inhibitory potency. The kinetic analysis revealed that isobaisseoside and xeroboside exhibited mixedtype inhibition, which suggests that they can bind to both the enzyme active site and an allosteric site, altering both V_{max} and $K_{\rm m}$. This mode of inhibition may provide a therapeutic advantage by offering multiple binding interactions that stabilize the inhibitor-enzyme complex. However, it is noteworthy to highlight that the two-site model of mixed inhibition might not be mechanistically relevant as reported by Pesaresi.39 Using a statistical approach and by combining statistical analysis of enzyme inhibition cases with a theoretical investigation of inhibition models, Pesaresi concluded that mixed inhibitors bind exclusively to the active site, ruling out allosteric involvement.³⁹ Consistently, we propose that our inhibitors interact primarily with the active site, causing a conformational change that affects substrate binding and enzyme activity. This would explain the mixed inhibition observed in our kinetic data without necessarily invoking an allosteric site. While the inhibitors do not completely block substrate binding as in pure



Fig. 2 CA IX inhibitory activity of *C. officinalis*-derived coumarins. (A) Dose–response curve of compounds **1–5** and AAZ, and (B) IC_{50} values of the inhibitory activity of the tested compounds. Data are mean \pm SD, N = 3. **P < 0.01 and ***P < 0.001 vs. AAZ. #P < 0.5, ##P < 0.01, and ###P < 0.001 vs. C5 (xeroboside). (C–F) Michaelis–Menten (C and E) and Lineweaver–Burk (D and F) plots of CA IX inhibitory activity of compounds **5** and **3**, respectively.

competitive inhibition, they still influence the enzyme activity, likely through modulation of the active site conformation. The outcomes of the computational studies conducted in this work support the idea that the inhibitors bind within or near the catalytic region of CA IX, potentially altering the active site structure and leading to mixed inhibition behavior. AAZ, a well-known CA inhibitor, displayed non-competitive inhibition, consistent with its mechanism of action that involves binding to an allosteric site, leading to a decrease in V_{max} without affecting K_{m} . The comparison with AAZ highlights the efficacy of the

natural coumarins, particularly xeroboside and isobaisseoside, as potent CA IX inhibitors.

3.3. Molecular docking analysis

The binding affinities and interacting residues of the isolated coumarins with CA IX were evaluated using molecular docking. Fig. 3, 4, and 5 represent the results of docking simulations, depicting the positioning of different ligands in the binding site of the target enzyme and showcasing residues involved in polar



Fig. 3 Binding site interactions of compounds 1 (A) and 2 (B) with CA IX.

and hydrophobic interactions. Among tested coumarins, 6,7-di-*O*-glucopyranosyl esculetin and xeroboside demonstrated the lowest binding energies $(-8.1 \text{ and } -8.3 \text{ kcal mol}^{-1})$,

respectively (Table 1). However, the remaining coumarins displayed comparable binding energies ranging from -7.3 to -7.7 kcal mol⁻¹, and AAZ had a binding energy of



Fig. 4 Binding site interactions of compounds 3 (A) and 4 (B) with CA IX.

-6.5 kcal mol⁻¹ (Table 1). These findings highlight the potential of these coumarins as effective CA IX inhibitors. The binding site of CA IX displayed a dense network of hydrophobic

interactions for all the tested coumarins. These hydrophobic interactions were crucial in ligand–enzyme complexes stability, thereby enhancing the binding affinity and inhibitory potential



Fig. 5 Binding site interactions of compound 5 (A) and acetozolamide (B) with CA IX.

of the tested compounds. The poses revealed docking of all compounds into the binding site of CA IX, which is also occupied by the reference drug AAZ. This consistent docking pattern suggested that the coumarins and the reference drug share a common binding site, highlighting the potential of these coumarins as effective CA IX inhibitors. In addition, isolated Table 1 Molecular docking analysis of the binding affinity of C. officinalis-derived coumarins towards CA IX

	Lowest binding energy (kcal mol ⁻¹)	Polar interacting residues	Hydrophobic interacting residues
6,7-di-O-glucopyranosyl esculetin	-8.1	Thr201, Thr200, His96, His94, His68, and Asn66	Val130, Leu91, Gln92, Gln71, Arg64, and Leu199
Aesculin	-7.7	Thr200, Thr201, Pro202, His86, Gln71, and His119	Trp210, His94, Val121, Leu199, Gln92, and Pro203
Isobaisseoside	-7.3	Pro202, His119, and Gln71	Thr200, His68, Thr201, Leu199, His94, Val142, Trp210, Val121, Leu91, Gln91, Trp9, and Pro203
Skimmin	-7.4	His68, Gln71, Thr201, His94, and Thr200	His96, Leu199, Val130, and Gln92
Xeroboside	-8.3	His94, Asn66, Hi68, Thr201, and Thr200	His96, Trp9, Gln71, Pro203, Leu199, Val130, and Val121
AAZ	-6.5	His94, His96, Thr200, and Thr201	His68 and Leu199

compounds showed a high extent of polar interactions, significantly contributing to the stability and efficacy of these inhibitors.

3.4. Molecular dynamics (MD) simulations

The objective of MD simulations was to elucidate the long-term stability and interactions within the coumarins-CA IX complexes, thereby illuminating the inhibition mechanisms and various binding modes. By carefully examining the simulation trajectories, we aimed to gain comprehensive insights into how the structural dynamics of the compounds influence their CA IX inhibitory potential. Understanding these molecular mechanisms is critical for the strategic design of novel CA IXtargeting therapeutics. We focused on complexes identified through initial docking analyses that exhibited the most favorable binding affinities and conducted an in-depth evaluation using 100 ns MD simulations. This rigorous analysis included key parameters such as interaction energies, MM/PBSA binding free energies, root mean square deviations (RMSD), radius of gyration (R_g) , potential energy landscape (PEL), root mean square fluctuations (RMSF), solvent accessible surface area (SASA), and hydrogen bonding profile. Our examination covered both CA IX and its complexes with coumarins, aiming at gaining understanding of their interaction characteristics and dynamic behavior. These insights are essential for elucidating the inhibitory effects of the compounds and guiding the development of effective CA IX inhibitors.

3.4.1. Structural stability and dynamics. The results of the 100 ns RMSD calculation of the CA IX backbone are depicted in Fig. 6A. The RMSD profile of investigated systems initially displayed a rising trend indicative of the system adjusting and transitioning from its initial conformation. This phase of adaptation is common as the system moves towards a more energetically favorable state. Following this initial adjustment, the RMSD profile exhibited a highly fluctuating trend, reflecting the dynamic nature of the enzyme and its interactions with the bound inhibitors. Throughout most of the simulation, these fluctuations persisted, suggesting ongoing conformational changes and adjustments within the enzyme–inhibitor complexes. It wasn't until the last quarter of the simulation that

the RMSD values began to stabilize, indicating that the system had reached a state of equilibrium. This period of equilibrium is crucial as it signifies that the conformational changes have settled, and the system is now in a stable state suitable for detailed analysis.

To investigate the inherent movement and conformational changes of the coumarin molecules independent of their interaction with CA IX, we investigated their RMSD values (Fig. 6B). The RMSD values were obtained by aligning the trajectory frames to the initial structure of each drug and calculating the deviation of the coordinates. This method enables an assessment of how much each drug deviates from its initial conformation over the 100 ns simulation period. Such an analysis is crucial in drug design research to evaluate the stability and dynamic behavior of drug molecules. As shown in Fig. 6B, the RMSD analysis of different coumarins revealed notable variations in their RMSD profiles. The compounds exhibited fluctuation patterns in their RMSD values indicating that they maintained relatively stable conformations throughout the simulation. This stability suggests the presence of strong and consistent interactions between isolated coumarins and CA IX.

The results of the R_{g} calculations for the unbound CA IX and its complexes with various coumarins are shown in Fig. 7A. The R_g profile provides insights into structural stability and compactness of the enzyme and/or its complexes over the course of the simulation.40 Throughout the majority of the simulation time, the R_g profiles of the different coumarin–CA IX complexes demonstrated a consistent and stabilized behavior, comparable to that observed for the unbound enzyme. This indicates that the binding of coumarins does not disrupt the overall compactness of the enzyme structure. Interestingly, among all the complexes studied, xeroboside (the compound with the lowest IC₅₀ value) exhibited the lowest average R_g value. This suggested that the xeroboside-CA IX complex is more compact compared to the other coumarin-CA IX complexes. A lower R_g value often correlates with a more tightly bound and stable complex, which could contribute to the enhanced inhibitory potency of xeroboside. The compact nature of the xeroboside-CA IX complex might facilitate stronger interaction



Fig. 6 MD simulation of CA IX and its complexes with C. officinalis-derived coumarins; (A) backbone RMSD of the unbound CA IX and CA IXcoumarin complexes and (B) RMSD of isolated compounds.

with the active site, thus explaining its superior inhibitory activity.

The results of the SASA calculations for the free CA IX enzyme and its complexes with various coumarins are depicted in Fig. 7B. The SASA profile provides insights into the surface exposure of the enzyme and its complexes to the solvent, which is a critical factor in understanding the interaction and stability of the complexes.¹⁴ Throughout the majority of the simulation span, the SASA profiles of different coumarin–CA IX complexes demonstrated an equilibrated behavior similar to that observed for the unbound enzyme. This indicates that the binding of coumarins does not significantly alter the overall solvent exposure of the enzyme structure. During the simulation, a noticeable decrease in SASA values for all systems was observed. This reduction in SASA values suggested that the enzyme and its complexes become more compact over time, with less surface area exposed to the solvent. Such a decrease in SASA is indicative of a stable binding interaction where the enzyme–inhibitor complex achieves a more energetically favorable conformation by minimizing its solvent-exposed surface area. The equilibrated and reduced SASA profiles across the different complexes suggested that the coumarins contribute to the stabilization of the enzyme by promoting a more compact structure. This stabilization is crucial for the inhibitory activity, as it indicates that the enzyme–inhibitor complex achieves a conformation that is less prone to destabilizing interactions with the solvent.

The SASA findings are consistent with the $R_{\rm g}$ results, reinforcing the notion that the binding of coumarins leads to



Fig. 7 MD simulation of CA IX and its complexes with *C. officinalis*-derived coumarins; (A) radius of gyration of CA IX and CA IX-coumarin complexes and (B) SASA of CA IX and various complexes.

a stabilized and compact enzyme-inhibitor complex. The observed decrease in SASA values further supports the idea that these complexes are energetically favorable and structurally stable. This stability is essential for the effective inhibition of CA IX by the coumarins, as a stable complex is less likely to dissociate and more likely to maintain its inhibitory function.

3.4.2. Hydrogen bonding dynamics and flexibility. The hydrogen bonding profile of coumarin–CA IX complexes is represented in Fig. 8A. Hydrogen bonds are critical interactions that contribute significantly to the stability and specificity of enzyme–inhibitor complexes. They play a crucial role in maintaining the structural integrity of the enzyme–ligand complex and in facilitating the correct positioning of the inhibitor within the enzyme active site.⁴¹ Our analysis revealed that the

coumarin–CA IX complexes exhibited varying extents of hydrogen bonding interactions throughout the simulation period. Notably, xeroboside showed the highest number of hydrogen bonds among the tested coumarins. This extensive hydrogen bonding network suggests a strong and stable interaction between xeroboside and CA IX, which likely contributes to its superior inhibitory activity. The formation of a high number of hydrogen bonds indicates that xeroboside is well accommodated within the enzyme active site, forming stable interactions that enhance its binding affinity. Following xeroboside, isobaisseoside also displayed a significant number of hydrogen bonds, correlating with its relatively low IC_{50} value. The robust hydrogen bonding interactions observed for



Fig. 8 MD simulation of CA IX and its complexes with *C. officinalis*-derived coumarins; (A) hydrogen bonding profile of CA IX-coumarin complexes, and (B) backbone RMSF per residue number for CA IX and CA IX-coumarin complexes.

isobaisseoside further underline its effective binding and stabilization within the CA IX active site.

The RMSF profile of the free CA IX and its complexes with different coumarins is depicted in Fig. 8B. RMSF analysis provides insights into the flexibility of individual residues within the protein structure over the course of the MD simulation. High RMSF values indicate regions of the protein that exhibit significant flexibility, while low RMSF values denote more rigid regions. The RMSF profiles of the CA IX-coumarin complexes were found to be similar to that of the free enzyme. This observation suggested that the binding of coumarins with CA IX does not induce significant conformational changes or alter the flexibility of the enzyme residues in a substantial manner. Despite the similarity in RMSF profiles, subtle differences were observed in specific regions, which may reflect localized interactions between the enzyme and coumarins. These localized interactions can stabilize certain regions of the enzyme, potentially enhancing the inhibitory effects of the coumarins without causing widespread conformational changes.

3.4.3. Interaction energy. The analysis of interaction energies highlights the significance of both electrostatic and van der Waals forces in the binding of coumarins to CA IX. The results of the Coulomb short-range (Coul-SR) and Lennard-Jones short-range (LJ-SR) interaction energies for the CA IX-coumarin complexes are presented in Fig. 9 and summarized in Table 2. These interaction energies help evaluate the binding affinities and enzyme–inhibitor complexes stability, with lower values



Coul-SR interaction energies

Fig. 9 MD simulation of CA IX and its complexes with *C. officinalis*-derived coumarins; (A) Coulomb-SR interactions energies and (B) Lennard-Jones-SR interactions energies of the enzyme amino acid residues with isolated compounds.

indicating stronger interactions and potentially higher inhibitory efficacy. Among the studied coumarins, xeroboside exhibited the lowest average Coul-SR and LJ-SR interaction energies. This finding correlates with its highest inhibitory activity, as indicated by its low IC_{50} value. The strong electrostatic (Coul-SR) and van der Waals (LJ-SR) interactions observed

Table 2 Average Coul-SR and LJ-SR interaction energies for the complexes formed between C. officinalis-derived coumarins and CA IX

	Coul-SR interaction energy		LJ-SR interaction energy		
	Average (kJ mol ^{-1})	RMSD (nm)	Average (kJ mol ^{-1})	RMSD (nm)	
6,7-di-O-glucopyranosyl esculetin	-46.27 ± 4.7	45.69	-58.32 ± 3.1	37.69	
Aesculin	-37.90 ± 3.0	7.80	-95.40 ± 1.9	17.74	
Isobaisseoside	-72.47 ± 3.8	36.79	-96.67 ± 1.73	12.66	
Skimmin	-42.43 ± 3.1	32.14	-68.74 ± 2.5	11.46	
Xeroboside	-94.09 ± 3.6	32.45	-115.83 ± 1.7	11.20	

Table 3 The results of MM/PBSA calculations (kJ mol⁻¹)

	$\Delta E_{\rm yrdyr}$	ΔE_{ele}	$\Delta G_{ m solv}$	ΔG_{max}	ΔG_{total}
	vuw	eie	- 5019	- gas	- total
6,7-di-O-glucopyranosyl esculetin	-11.71 ± 0.79	-13.42 ± 1.93	19.46 ± 1.31	-25.13 ± 2.11	-5.67 ± 2.35
Aesculin	-25.08 ± 0.64	-7.94 ± 1.64	20.86 ± 1.11	-33.02 ± 1.77	-12.16 ± 2.06
Isobaisseoside	-25.13 ± 1.76	-15.16 ± 1.38	26.30 ± 1.30	-40.29 ± 2.20	-14.00 ± 2.67
Skimmin	-17.96 ± 1.65	-11.66 ± 1.53	18.76 ± 3.19	-29.62 ± 2.25	-10.86 ± 3.72
Xeroboside	-34.62 ± 1.78	-31.58 ± 1.96	$\textbf{38.94} \pm \textbf{0.19}$	-66.20 ± 2.63	-27.26 ± 2.48

for xeroboside suggested a robust binding affinity to the active site of CA IX, contributing to its effective inhibition. Isobaisseoside, the coumarin with the second lowest IC_{50} value, also demonstrated favorable interaction energies, with average Coul-SR and LJ-SR values slightly higher than those of xeroboside. This trend further supported the correlation between low interaction energies and high inhibitory activity. Aesculin, although not as potent as xeroboside or isobaisseoside, showed

LJ-SR interaction energies comparable to those of isobaisseoside, indicating a similar degree of van der Waals interactions.

3.4.4. MM/PBSA analysis. The MM/PBSA calculations provide an in-depth quantitative analysis of the binding free energies of different coumarins to CA IX. The results summarized in Table 3 highlight the contributions of van der Waals (ΔE_{vdw}) , electrostatic (ΔE_{ele}) , solvation (ΔG_{solv}) , and gas phase (ΔG_{gas}) interactions to the total binding free energy (ΔG_{total}) for each complex. Xeroboside exhibited the lowest total binding



Fig. 10 The potential energy landscape (PEL) analysis of free CA IX, 6,7-di-O-glucopyranosyl esculetin, and aesculin. (A) The 2D heat map of PEL, (B) the lowest energy conformation of investigated coumarins within the binding site of CA IX, and the key residues involved in the binding mechanism. (C) A 3D representation of PEL.



Fig. 11 The potential energy landscape (PEL) analysis of free isobaisseoside, skimmin, and xeroboside. (A) The 2D heat map of PEL, (B) the lowest energy conformation of investigated coumarins within the binding site of CA IX, and the key residues involved in the binding mechanism. (C) A 3D representation of PEL.

free energy, indicating the strongest binding affinity among the studied coumarins. This strong binding is reflected in both its substantial van der Waals and electrostatic interaction energies, which are the most favorable among all the compounds. This aligns with the experimental data where xeroboside displayed the lowest IC₅₀ value, underscoring its potent inhibitory effect. Isobaisseoside also demonstrated a strong binding affinity. Similar to xeroboside, the binding of isobaisseoside is characterized by favorable van der Waals and electrostatic interactions, though these interactions are less pronounced than those observed for xeroboside. The strong binding affinity of isobaisseoside is also corroborated by its low IC₅₀ value, making it the second most potent inhibitor among the tested coumarins. The solvation energies (ΔG_{solv}) for all complexes are positive, indicating an unfavorable desolvation process upon binding. However, the strong negative contributions from van der Waals and electrostatic interactions outweigh these unfavorable solvation effects, resulting in overall negative total binding free energies. Overall, the MM/PBSA analysis underscores

xeroboside superior binding affinity towards CA IX, followed by isobaisseoside and aesculin. The strong van der Waals and electrostatic interactions primarily drive the binding affinities of these coumarins. These findings align with the inhibitory activities observed in experimental assays.

3.4.5. Potential energy landscape. Principal component analysis (PCA) was implemented to assess the conformational dynamics of the unbound CA IX and its complexes with target coumarins. The potential energy landscape (PEL) of the unbound CA IX and its complexes with various coumarins is depicted in Fig. 10 and 11. Panel A in both figures illustrates the 2D PEL of free CA IX and different complexes, showcasing distinct energy minima, which indicate stable conformational states throughout the simulation trajectory. The lowest energy basins in the PEL correspond to the most stable conformations of the enzyme, which are critical for understanding its inherent stability and functional states. Panel B highlights the minimum energy conformations of CA IX in complexes with different coumarins. The CA IX structure serves as a reference for

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comparing the structural adaptations upon ligand binding. For 6.7-di-O-glucopyranosyl esculetin and aesculin, the snapshots show significant interactions within the enzyme active site, including hydrogen bonding and hydrophobic interactions, which likely contribute to their inhibitory effects. The binding conformations revealed how these compounds fit into the active site, inducing minor or major conformational changes to optimize binding. Panel C presents the three-dimensional potential energy surfaces for these complexes. The stable binding interactions and minimal energy fluctuations throughout the simulations indicated that coumarin-CA IX complexes achieve stable binding conformations essential for their inhibitory function. Notably, the PELs of xeroboside displayed more pronounced energy basins, indicating highly stable and energetically favorable binding conformations. This suggested a robust interaction and a potent inhibitory effect on CA IX. Overall, the PEL analysis underscores the importance of stable and energetically favorable binding conformations in the inhibitory activity of these coumarins against CA IX. The distinct energy minima and stable binding interactions highlight their potential as effective CA IX inhibitors, paving the way for further development and optimization of these compounds for therapeutic applications.

4. Conclusion

This study examined the inhibitory capabilities of C. officinalisderived coumarins against CA IX using an integrated approach, including in vitro and in silico investigations. The results identified xeroboside as the most effective CA IX inhibitor followed by isobaisseoside and both compounds displayed a mixed inhibition mode. Molecular docking revealed polar and hydrophobic interactions between the isolated coumarins and CA IX. MD simulations offered insights into the dynamic behavior of the coumarins within the CA IX binding site. Xeroboside and isobaisseoside exhibited consistent trajectories, with lower Coul-SR and LJ-SR interaction energy values, indicating favorable electrostatic and van der Waals interactions. Xeroboside showed extensive hydrogen bonding with CA IX, affirming its high inhibitory potential. Additionally, xeroboside exhibited lower average R_g and SASA values, suggesting a more compact structure and reduced solvent exposure of the xeroboside-CA IX complex. The MM/PBSA analysis indicated that xeroboside had the most favorable binding free energies when interacting with CA IX. Furthermore, the PEL analysis showed distinct and stable conformational states for the CA IX-ligand complexes, with xeroboside presenting the most stable and lowest energy configuration. These computational results are consistent with our experimental findings, highlighting the potential of xeroboside and isobaisseoside as potent CA IX inhibitors. Therefore, C. officinalis-derived coumarins are promising CA IX inhibitors, pending in vitro, in vivo, and clinical investigations to determine their exact mechanism(s) of action.

Data availability

The manuscript and ESI[†] contain all data supporting the reported results.

Conflicts of interest

All authors declare no conflicts of interest in relation to the manuscript.

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