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Design, synthesis, molecular modelling investigation and biological studies of novel human carbonic anhydrase inhibitors based on coumalic acid†

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The present study provides the design, synthesis, molecular modelling investigation and biological evaluation of a series of coumalic acid-based selective inhibitors of hCA IX and XII. Based on the previously obtained results with carboxamide analogues of coumalic acid, here we explored some modifications of the original scaffold with the aim to expand our knowledge about these compounds and to promote structure–activity relationship (SAR) studies. Structural modifications as lactone-to-lactam conversion, repositioning of the carboxylic acid moiety, and the introduction of ester or amidic linkers, as well as triazole-based elongation *via* click chemistry have been performed. The synthesised compounds were tested for their inhibitory activities against hCA I, II, IX, and XII showing the lactone moiety to be crucial for inhibitory potency. Particularly, ester derivatives demonstrated selectivity for hCA IX and XII, with certain compounds exhibiting micromolar affinities. Notably, compound 8, featuring a bromine substitution, displayed the highest selectivity and potency against hCA IX and XII. Molecular docking studies further elucidated the binding mechanisms, revealing that the lactone ring hydrolysis plays a significant role in the inhibition process. These results offer valuable insights into the structure–activity relationships (SAR) of coumalic acid analogues and support their further biological investigation for cancer therapy by targeting hCA IX and XII.

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1. Introduction

Homeostasis is the crucial state required for the optimal physical, chemical, and internal functioning of all living systems. Consequently, different variables are constantly controlled such as fluid balance, ion concentration, and pH equilibrium between intra- and extra-cellular environments.² Among the different homeostatic mechanisms, carbonic anhydrases (CAs, EC 4.2.1.1) play an important role as pH regulators.^{3,4} These ubiquitous metalloenzymes catalyse the reversible hydration of CO₂ in HCO₃, an important reaction involved in a plethora of fundamental processes.^{5,6} To date, eight distinct genetic families have been defined α -, β -, γ -, δ -, η-, ζ-, θ-, and ι-, with the latter having been very recently discovered. The α -CAs are the best-characterised enzymes as they are expressed in vertebrates, fungi, protozoa, plants, algae, diatoms, archaea, and bacteria.12 The fifteen human carbonic anhydrase isoforms (hCAs) belong to the α -family

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and cover a pivotal role in a multitude of physiological functions such as respiration, transport of carbon dioxide between metabolising tissues and lungs, pH homeostasis, electrolyte secretion in various tissues/organs, as well as biosynthetic reactions. However, these enzymes may also contribute to pathological processes when their dysregulated expression and/or abnormal activity occurs (*e.g.*, in glaucoma, oedema, epilepsy, altitude sickness, and cancer). For this reason, different libraries of compounds have been explored as CA inhibitors (CAIs) and activators. T-22

The transmembrane tumour-related isoforms CA IX and XII are overexpressed in solid hypoxic tumours due to their activation by the HIF pathway. 23-25 Their mechanism of action is associated with intra- and extra-cellular pH regulation which contributes to tumour proliferation and invasiveness.^{26,27} Accordingly, their inhibition could be considered as a prognostic tool and a therapeutic target for tumour treatment.²⁸⁻³⁰ Among the CAIs developed so far, the coumarin-based ones occupy a prominent place due to their selective inhibition mechanism against hCA isoforms IX and XII. 21,31-33 The lactone portion of the coumarin core undergoes hydrolysis by the Zn(II) bound hydroxide ion, leading to the cinnamic acid product, or the open form, considered the real inhibitor of the enzyme.34 The ability of the latter to bind the entrance of the active site cavity, where the differences among the isoforms are higher, results in increased selectivity. In 2019 Cornelio and colleagues performed the molecular simplification of the saccharin scaffold to obtain potent hCAs inhibitors based on the 5-arylisothiazol-3(2H)-one core (as mono or dioxides at the sulphur atom).³⁵ Recently, we employed a similar approach on coumarin nucleus to design a series of potent and selective hCA IX and XII inhibitors provided of the 2H-pyran-2-one core 1).36 These compounds, obtained from the commercially available coumalic acid, retained the lactone moiety that should represent the molecular attribute responsible for the inhibition of hCAs, through (i) its putative hydrolysis provoked by the enzyme and successive binding in the entrance of the active site, or (ii) anchoring the zinc-bound water in its hydrolysed/nothydrolysed form. Position 5 of the 2H-pyran-2-one core was provided with an amide linker group bearing different (un)substituted aromatic or (hetero)(cyclo)aliphatic residues.36 With the aim to expand our knowledge about these compounds and to promote structure-activity relationship (SAR) studies, we explored some modifications of the original scaffold (Fig. 1).

The reasons behind these structural changes can be resumed as follows:

- The docking studies performed in the previous work underlined the importance of the lactone moiety in obtaining potent hCA IX and XII inhibitors.³⁶ To challenge this claim, we evaluated the effects of lactone to lactam replacement on hCA inhibition (compounds 1–2).
- With the aim of investigating the importance of the carboxylic acid moiety position, we tried to move this fragment from position 5 of the original coumalic acid (2-oxo-2*H*-pyran-5-carboxylic acid) to position 6, as in the compound 3 (2-oxo-2*H*-pyran-6-carboxylic acid, Fig. 1).
- Novel coumalic acid analogues provided with ester moiety in spite of amidic of the formers already published, have been designed to investigate the influence of nitrogen

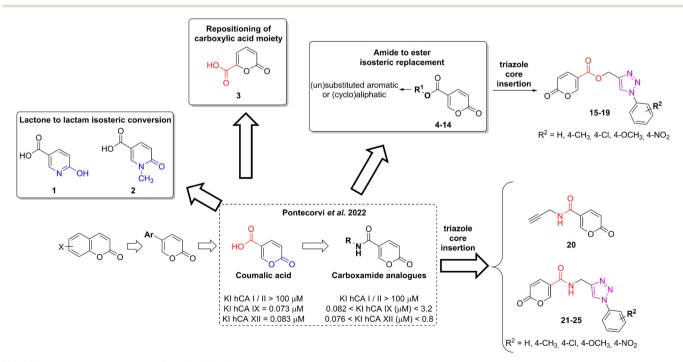


Fig. 1 Design of the novel coumalic acid derivatives.

to oxygen isosteric substitution at the linker moiety (compounds 4-14).

• The insertion of the triazole ring on the coumarin core has been extensively investigated to obtain potent and selective hCA IX and XII inhibitors. 37-39 Taking into account this evidence, we also evaluated the effect of molecular "elongation" obtained by exploiting click chemistry to insert triazole core between the amide (or ester) linker, and the (un) substituted aromatic residues (compounds 16–24).⁴⁰

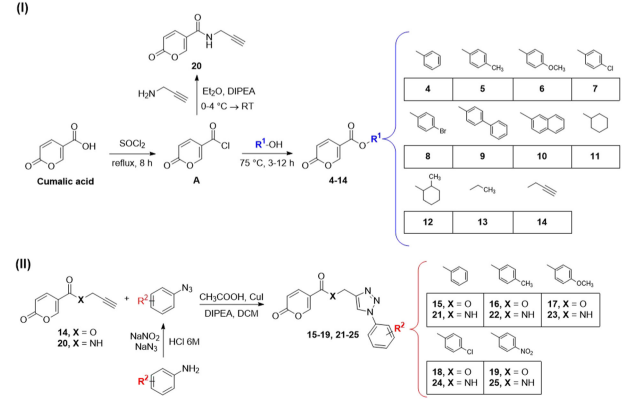
All the compounds have been evaluated against four hCA isoforms, the two cancer-related ones (hCA IX and XII) and the two off-targets hCA I and II. Moreover, to provide a structural elucidation of the detected $K_{\rm I}$ values for coumalic acid derivatives towards hCA IX and hCA XII, molecular docking simulations were performed. The further aim of this study is to understand the relationships between the coumalic acid isomers and their different inhibitory activities. The mechanism of action of coumalic acid derivatives in carbonic anhydrases inhibition is based on literature reports, showing that the coumalic acid ring is hydrolysed by the Zn(II)-coordinated hydroxide ion or water molecule within the hCAs active site, generating the corresponding E/Z cinnamic acid derivatives. ^{36,38,40}

2. Chemistry

Compounds 1-3, corresponding to the derivatives (molecules 1 and 2) and the coumalic acid analogue endowed with carboxylic acid moiety at position 6 (compound 3), have been commercially obtained. On the contrary, the derivatives 4-25 have been achieved by employing different synthetic procedures (Scheme 1I and II).

The synthesis of the compounds provided with ester linker (4-14) has been performed through a multi-step synthetic approach. In the first step, coumalic acid was converted into the corresponding acyl chloride A (Scheme 1I) by refluxing in thionyl chloride (SOCl2). The acyl chloride A was employed in the subsequent reaction with (un) substituted aromatic or cycloaliphatic alcohol, in absence of solvent, at a temperature of 75 °C for 3 h or 12 h depending on the reagents, to achieve the corresponding ester derivatives (Scheme 1I).41

Compound 20, the amidic analogue of derivative 14, has been synthesised by means of a slightly modified procedure, by reacting the acyl chloride A with propargyl amine in the presence of diisopropylethylamine (DIPEA) as base. The reaction was performed in diethyl ether (Et₂O) at reduced temperature (0-4 °C). Compounds 14 and 20 correspond to the alkyne intermediates employed in the "click" reactions exploited to obtain the triazole derivatives provided with the ester (15-19) or amidic (21-25) linker, respectively.³⁹ Particularly, these compounds (14 and 20) were reacted with azides (that in turn have been obtained from corresponding amines with NaN3 and NaNO2 in 6 M HCl), in the presence of CH3COOH, CuI, and DIPEA in



Scheme 1 Synthetic approaches employed for the synthesis of compounds 4-25.

dichloromethane through a copper(i)-catalysed click reaction.³⁹

Structural characterization of the final compounds 4–25 was obtained by means of 1 H-NMR and 13 C-NMR spectroscopic analysis. Elemental analyses for C, H, and N were recorded on a Perkin-Elmer 240 B microanalyzer and the analytical results were within $\pm 0.4\%$ of the theoretical values for all compounds (data not shown).

3. Inhibition assay

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Compounds 1-25 were evaluated for their ability to inhibit four different isoforms of hCA, the two off-targets hCA I and II and the two tumour-related isoforms hCA IX and XII. The inhibitory activity data (K_I) obtained for compounds 1-25 are reported in Table 1 along with the K_I values of coumalic acid and the related carboxamide analogues which exhibited the best outcomes in our previous work (8A, 9A and 11A, structures in blue in Table 1).36 The inhibitory activity data of these compounds (coumalic acid, 8A, 9A and 11A) have been added with the aim of performing an exhaustive structure-activity relationship (SAR) analysis. One of the first purposes of this study was to demonstrate the importance that the lactone moiety plays in the inhibition of hCA as suggested by our previously accomplished molecular modelling studies.³⁶ To confirm this aspect, the replacement of the lactone fragment with the lactam one has been performed (derivatives 1-2) showing the loss of inhibitory activity, these compounds being ineffective against all the evaluated isoforms ($K_{\rm I} > 100 \, \mu {\rm M}$). The other structural modification attempted concerns the change of position of the carboxylic acid group of the coumalic acid from position 5 to 6, to obtain the isomer 3. Surprisingly, compound 3 resulted to be devoid of inhibitory activity with $K_{\rm I} > 100~\mu{\rm M}$ against all the evaluated isoforms (see section 4, molecular modelling).

The compounds obtained by the amide to ester replacement of the linker moiety (4-14, Table 1) preserved the selectivity against the hCA IX and XII isoforms, these compounds being ineffective against the two off-targets ($K_{\rm I}$ hCA I/II $> 100 \mu M$). However, these derivatives showed a diminished affinity against the two cancer-related isoforms with $K_{\rm I}$ values falling in the micromolar range (Table 1). These results demonstrate that the NH/O isosteric replacement at the linker moiety had detrimental effects on potency. Besides this general observation, some SAR considerations can be performed. In particular, it is possible to note that the insertion of substituent groups (compounds 5-10) on the phenyl ring generally led to the increase of inhibitory activity, especially against hCA IX. As for hCA XII, the inhibition values were found to be highly dependent on the substituents considered. The best results were obtained for compound 8, endowed with the bromine atom at the para position of the phenyl ring. Indeed, concerning the unsubstituted parent compound 4, derivative 8 exhibited a 7-fold and a 71-fold increase of hCA IX and XII inhibition, respectively, resulting as the most potent hCA XII inhibitor of the entire series (KI hCA IX = 3.14 μ M; K_I hCA XII = 0.91 μ M, Table 1). As for compounds 7 (4-Cl), 9 (4-Ph) and 10 (1-naphthyl), comparable inhibition values against hCA IX were recorded when compared to the unsubstituted analogues 4. On the other hand, reduced (7-10) or absent (9) affinity against hCA XII isoform was observed. Considering the (cyclo)aliphatic derivatives (11-14), only compound 11 exhibited a single-digit micromolar inhibition of hCA XII $(K_I \text{ hCA XII} = 3.12 \text{ } \mu\text{M})$ the others being endowed with $K_{\rm I}$ s values greater than 10 μ M against both the isoforms. Interestingly, the carboxamide analogue of compound 11 (Table 1, 11A) was one of the most active compounds of the previously reported series, thus further confirming the unfavourable effects of the N/O isosteric substitution at the linker moiety. Keeping into account the beneficial effects that the insertion of triazole linker exerts on the coumarin core, 37-39 we proposed similar modifications on this scaffold, by combining the triazole ring with the carboxamide or ester linker group (compounds 15-19 and 21-25). Although maintaining the selectivity against the two tumour-related isoforms, being these compounds ineffective against hCA I and II ($K_{\rm I} > 100 \, \mu {\rm M}$), the inhibitory activity data still evidenced a detrimental effect of this molecular elongation with most of the compounds exhibiting poor inhibition of both the isoforms, regardless of linker type and/or substituent group. As a matter of fact, only two compounds endowed with the triazole ring (16 and 25), showed K_I values below 10 μ M against hCA IX; conversely, the affinity of most of the compounds was quite scarce with $K_{\rm I}$ values greater than 20 μ M.

4. Molecular modelling studies

Coumalic acid and compounds 3 and 11 were docked to the crystallographic structure of hCA IX (PDB-ID: 3IAI) and hCA XII (PDB-ID: 1JD0) by the GOLD program. Considering the inhibition mechanism of these derivatives through the hydrolysis of the lactone moiety, the resulting open enolic forms in both E and Z configurations of the double bond were also docked. The Zn(II)-bound water molecule was kept as a part of the receptor during molecular docking simulations.36,43 Docking results of the closed form of all the examined compounds showed that all of them are able to fit the hCA IX and hCA XII catalytic sites. Coumalic acid and 11 are able to anchor the Zn(II)-bound water molecule through their lactone moiety, while 3 is unable to form a coordination bond with it. Instead, 3 establishes only an H-bond interaction with the carboxyl moiety with the Zn(II)-bound water molecule, which likely fails to activate the lactone ringopening mechanism, thus explaining its lack of activity against both hCAs ($K_{\rm I}$ > 100). The binding mode of coumalic acid against hCA IX was reproduced such as in

Table 1 Inhibition data of hCA II, hCA II, hCA IX and hCA XII with compounds 1-25 and the standard sulfonamide inhibitor acetazolamide (AAZ) by a stopped flow CO_2 hydrase.⁴² The coumalic acid and the compounds 8A, 9A, and 11A (structures in blue) belong to the carboxamide analogues series previously published,³⁶ and have been added with the aim of performing a more concise SAR analysis

		$K_{ m I}{}^a\left(\mu{ m M} ight)$			
Compound	Structure	hCA I	hCA II	hCA IX	hCA XI
Coumalic acid ³⁶	О	>100	>100	0.073	0.083
1	ОРОН	>100	>100	>100	>100
2	O N O OH	>100	>100	>100	>100
3	ON CH ₃	>100	>100	>100	>100
4		>100	>100	21.4	65.7
5	O CH ₃	>100	>100	12.21	1.83
6	OCH ₃	>100	>100	10.17	1.65
7	CI	>100	>100	15.1	76.3
8	o Br	>100	>100	3.14	0.91
8A ³⁶	O O Br	>100	>100	0.082	0.089

Table 1 (continued)

			$K_{ m I}^{~a} \left(\mu { m M} ight)$			
Compound	Structure	hCA I	hCA II	hCA IX	hCA XII	
9		>100	>100	11.43	>100	
9A ³⁶		>100	>100	0.083	0.076	
10		>100	>100	14.8	68.9	
11		>100	>100	19.91	3.12	
11A ³⁶		>100	>100	0.098	0.086	
12		>100	>100	12.3	82.8	
13	O CH ₃	>100	>100	14.4	92.6	
14		>100	>100	17.9	76.3	
15		>100	>100	18.6	55.8	

Table 1 (continued)

hCA I >100 >100 >100	hCA II >100 >100 >100	hCA IX 9.8 11.4	62.6 54.8
>100	>100	11.4	54.8
>100	>100	21.9	61.3
>100	>100	20.0	62.8
>100	>100	22.8	68.1
>100	>100	21.3	62.0
	>100	>100 >100	>100 >100 22.8

Table	1 د	(continued

			$K_{ m I}{}^a\left(\mu{ m M} ight)$			
Compound	Structure	hCA I	hCA II	hCA IX	hCA XII	
22	O NH NN	>100	>100	14.5	57.8	
23	CH ₃	>100	>100	23.5	58.7	
24	H ₃ C'	>100	>100	18.2	61.2	
25	O N N N N N N N N N N N N N N N N N N N	>100	>100	7.5	63.4	
AAZ	NO ₂	0.25	0.012	0.026	0.006	

^a Mean from 3 different assays, by a stopped-flow technique (errors were in the range of ±5–10% of the reported values).

a previous study³⁶ (Fig. 2A), while its binding to hCA XII was simulated herein, showing that the molecule establishes H-bond interaction with Asn62, Gln92 and Thr199 residues (Fig. 2D). Compound 3 in hCA IX establishes H-bond interactions with Gln92, Tyr7 and a π - π stacking interaction with His94 (Fig. 2B), whereas in hCA XII establishes H-bond interactions with Thr200, Asn62 and π - π stacking interaction with His94 (Fig. 2E). Focusing on the derivative 11, it demonstrates a higher inhibitory activity against hCA XII ($K_I = 3.12 \mu M$) compared to hCA IX ($K_I = 19.91 \mu M$). Analysis of docking

results with the closed form of the molecule shows that in hCA IX, 11 establishes H-bond interaction with Gln92 and a π - π stacking interaction with His94, while the cyclohexyl moiety binds in proximity of a lipophilic Leu/ Val-rich cluster composed of Leu91, Val121, Leu123, Val131, Leu135, Leu141, Val134 (Fig. 2C). Similarly, in hCA XII, 11 establishes the same interaction network that is reinforced by an additional H-bond interaction with Tyr7 (Fig. 2F) which is not established in hCA IX.

Moreover, the analysis of the respective cinnamic acids in the open E and Z forms of 11 provides additional insight into

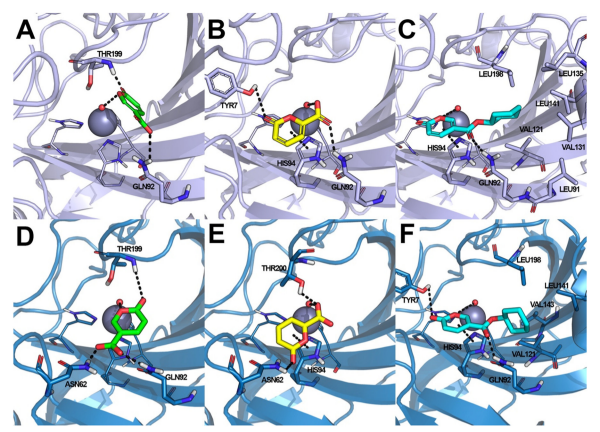


Fig. 2 Docking-based binding mode of coumalic acid (A and D), 3 (B and E) and 11 (C and F) derivatives within the catalytic site of hCA IX (A-C) and hCA XII (D-F). The structure of hCA IX is shown as violet cartoon, whereas the structure of hCA XII is shown as blue cartoon. Residues not involved in active interaction with the inhibitors are omitted for clarity. Coumalic acid is shown as green sticks, 3 as yellow sticks and 11 as cyan sticks. Interactions are highlighted by black dashed lines, while residues involved in the interactions are represented as sticks. The Zn(II) ion is shown as a grey sphere, while the coordinated water molecule is represented as a small red sphere.

the higher inhibitory activity of 11 against hCA XII compared to hCA IX. In fact, considering the Z conformation, which is reported to be more relevant than E in hCAs inhibition, 36 11 establishes with hCA XII an additional H-bond to Asn62. Furthermore, the cyclohexyl moiety occupies the hydrophobic cluster of hCA XII, unlike the pose obtained in hCA IX, where the same portion is docked on the opposite side (Fig. 3A). The E conformation of 11 binds to both hCA IX and hCA XII through the same interaction network: H-bond interactions with Gln92, Thr199, Thr200 and with the Zn(II)-bound water molecule (Fig. 3B-D).

Materials and methods

5.1 Chemistry

All the reagents, coumalic acid, 1-methyl-6-oxo-1,6dihydropyridine-3-carboxylic acid (compound 2) and other starting materials, employed in the synthetic procedures as well as HPLC analysis were obtained from commercial suppliers and were used without further purification. All melting points (m.p.) reported were uncorrected (temperatures are reported in °C) and measured on a Stuart® melting point apparatus SMP1. Nuclear magnetic resonance spectra (1H-NMR: 400 MHz and 13C-NMR: 101

MHz) were recorded in DMSO-d₆ or CDCl₃ using an Avance III 400 MHz spectrometer (Bruker, Milan, Italy). The processing and analyses of the NMR data were carried out with MestreNova software. Chemical shifts are reported in parts per million (ppm) and the coupling constants (J) are expressed in Hertz (Hz). Splitting patterns are designated as follows: s, singlet; d, doublet; t, triplet; q, quadruplet; m, multiplet; brs, broad singlet; dd, double of doublets.

Accurate mass measurements were performed with an Orbitrap Exploris 120 mass spectrometer (Thermo-Fisher Scientific, Waltham, MA, USA) coupled to an AP-MALDI (ng) ultra-high resolution (UHR) ion source (MassTech Inc., Columbia, MD, USA). The samples were prepared by adapting a spotting procedure reported elsewhere. 44 Briefly, 100 µl of acetonitrile/water (1:1 v/v) were added to ca. 1 mg of each synthesized compound and 1 µl of the suspensions thus obtained was dropped on the wells of the MALDI plate (Opti-TOF 184 spot sample insert, SCIEX, Italy). The analytes were then covered with an equal volume of a 5 mg ml⁻¹ solution of alpha-cyano-4-hydroxycinnamic acid (CHCA) dissolved in acetonitrile/ultrapure water/trifluoroacetic acid 50/50/0.1% (LaserBio Labs, Valbonne, France) used as a MALDI matrix.

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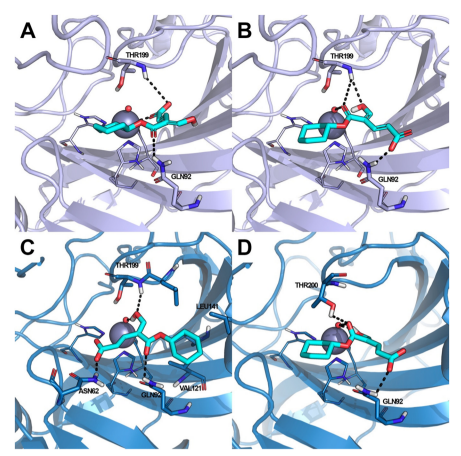


Fig. 3 Docking-based binding mode on hCA IX and hCA XII of the corresponding cinnamic acid in conformation E and Z of the double bond of 11. The structure of hCA IX is shown as violet cartoon, whereas the structure of hCA XII is shown as blue cartoon. Residues not involved in active interaction with the inhibitors are omitted for clarity. Compound 11 is shown as cyan sticks. Interactions are highlighted by black dashed lines, while residues involved in interaction are shown as sticks. The Zn(II) ion is shown as a grey sphere, while the coordinated water molecule is represented as a small red sphere.

After solvent evaporation, the samples were irradiated with an UV laser at a wavelength of 355 nm operating in spiral motion mode at a repetition rate of 1000 Hz with 1% laser energy. Other parameters, i.e. pos ionization voltage (V) and ion transfer tube temperature (°C), were in turn optimized to maximize the ionic intensity of each compound. Mass spectra were acquired in positive ion mode over a mass range of 150-500 Da with a mass resolution of $120\,000@m/z\ 200$ and a maximum injection time of 100 ms. The instrument was daily calibrated with Flexmix Calibration solution (Thermo-Fisher Scientific, Waltham, MA, USA) and fluorantrene was used as an internal lock mass by activating the EASY-IC source option allowing for a scan-to-scan mass scale recalibration in each analysis. Raw data were processed by using FreeStyle software v. 1.8 provided with the Xcalibur package. Δ values < 0.5 ppm between the theoretical and measured masses of the corresponding protonated species were obtained for all the compounds analysed. Due to the high mass accuracy achieved, measured masses were reported with five decimal places.

Thin layer chromatography (TLC) was performed on 0.2 mm thick silica gel-aluminium backed plates (60 F₂₅₄, Merck) was employed to monitor all the reactions. Preparative flash column chromatography was carried out on silica gel (230-400 mesh, G60 Merck). The yields shown are not optimised. Drying of organic solution was accomplished with anhydrous sodium sulphate. Evaporation of the solvent after reaction was carried out on a rotary evaporator. Elemental analyses for C, H, and N were recorded on a Perkin-Elmer 240 B microanalyzer (PerkinElmer, Waltham, MA, USA) and the analytical results are within $\pm 0.4\%$ of the theoretical values for all compounds (data not shown).

5.1.1 Synthetic procedure and characterization data of compounds 1-25

General procedure for the synthesis of 2-oxo-2H-pyran-5carbonyl chloride (A). To the powder of coumalic acid (1 eq.) placed in the round bottom flask, was dropwise added thionyl chloride (3.8 eq.). The synthesis was carried out under reflux conditions and stirred for 8 hours, after which a 1:4 mixture of dichloromethane and cyclohexane was added. The obtained suspension was filtered and the solid crystallised in cyclohexane to afford the title compound A as a brown solid (yield 55%, mp 72–75 °C). ¹H NMR (400 MHz, CDCl₃-d): δ 6.38 (dd, J = 9.9, 1.1 Hz, 1H, CH_{Pyr}), 7.76 (dd, J = 9.9, 2.8 Hz,

1H, CH_{Pyr}), 8.57 (dd, J = 2.8, 1.1 Hz, 1H, CH_{Pyr}). ¹³C NMR (101 MHz, CDCl₃-d): δ 115.3 (C_{Pyr}), 116.5 (CH_{Pyr}), 140.2 (CH_{Pyr}), 158.1 (CH_{Pyr}), 162.7 (CO), 162.8 (COCl).

General procedure B. A mixture of 2-oxo-2H-pyran-5carbonyl chloride (A) (1 eq.) and the proper alcohol (1 eq.) was heated at T = 75 °C under inert atmosphere (N₂) in absence of solvents and stirred for 3 to 12 h (TLC monitoring), depending on alcohol reactivity. The reaction quenched with slush and extracted dichloromethane (3 × 40 mL). The organics were reunited, washed with NaOH 2N and dried over sodium sulfate. The drying agent was removed by filtration and the organic phase was evaporated in vacuo. The obtained crude product was purified by silica gel column chromatography eluting with 33% ethyl acetate in petroleum ether to afford the intended product. When necessary, the solids obtained were further crystallised from petroleum ether to afford the desired compounds.

Phenyl 2-oxo-2H-pyran-5-carboxylate (4). Synthesised according to the **general procedure B** using phenol as starting material to afford the titled compound 4 as a white solid (51% yield). 1 H NMR (400 MHz, CDCl₃-d): δ 6.41 (dd, J = 9.8, 1.1 Hz, 1H, CH_{Pyr}), 7.16 (dd, J = 7.9, 1.7 Hz, 2H, Ar), 7.29 (t, J = 7.5 Hz, 1H, Ar), 7.43 (t, J = 7.9 Hz, 2H, Ar), 7.89 (dd, J = 9.8, 2.7 Hz, 1H, CH_{Pyr}), 8.51 (dd, J = 2.7, 1.1 Hz, 1H, CH_{Pyr}). 13 C NMR (101 MHz, CDCl₃-d): δ 111.6 (CH_{Pyr}), 115.5 (C_{Pyr}), 121.4 (2 × Ar), 126.4 (Ar), 129.6 (2 × Ar), 141.5 (Ar), 150.0 (CH_{Pyr}), 159.1 (CH_{Pyr}), 159.5 (CO), 161.6 (COOR). Measured monoisotopic mass [C₁₂H₈O₄ + H]⁺: m/z 217.04954 (Δ : 0.00 ppm) – theoretical monoisotopic mass: m/z 217.04954.

p-Tolyl 2-oxo-2*H-pyran-5-carboxylate* (5). Synthesised according to the **general procedure B** using *p*-cresol as starting material to afford the titled compound 5 as a white solid (25% yield). ¹H NMR (400 MHz, DMSO-d₆): δ 2.33 (s, 3H, CH₃), 6.50 (dd, J = 9.8, 1.1 Hz, 1H, CH_{Pyr}), 7.08–7.18 (m, 2H, Ar), 7.26 (d, J = 8.2 Hz, 2H, Ar), 7.93 (dd, J = 9.8, 2.7 Hz, 1H, CH_{Pyr}), 8.85 (d, J = 2.3 Hz, 1H, CH_{Pyr}). ¹³C NMR (101 MHz, DMSO-d₆): δ 20.8 (CH₃), 111.5 (C_{Pyr}), 115.3 (CH_{Pyr}), 121.9 (2 × Ar), 130.4 (2 × Ar), 135.9 (Ar), 142.5 (Ar), 148.3

(CH_{Pyr}), 160.0 (CH_{Pyr}), 160.8 (CO), 162.3 (COOR). Measured monoisotopic mass $[C_{13}H_{10}O_4 + H]^+$: m/z 231.06511 (Δ : -0.35 ppm) – theoretical monoisotopic mass: m/z 231.06519.

4-Methoxyphenyl 2-oxo-2H-pyran-5-carboxylate (6). Synthesized according to the **general procedure B** using 4-methoxyphenol as starting material to afford the titled compound 6 as a yellow solid (36% yield). ¹H NMR (400 MHz, DMSO-d₆): δ 3.78 (s, 3H, CH₃), 6.50 (d, J = 9.8 Hz, 1H, CH_{pyr}), 7.00 (d, J = 8.8 Hz, 2H, Ar), 7.18 (d, J = 8.7 Hz, 2H, Ar), 7.93 (dd, J = 9.9, 2.7 Hz, 1H, CH_{pyr}), 8.85 (d, J = 2.6 Hz, 1H, CH_{pyr}), 1³C NMR (101 MHz, DMSO-d₆): δ 55.9 (CH₃), 111.5 (C_{pyr}), 114.9 (2 × Ar), 115.3 (CH_{pyr}), 123.0 (2 × Ar), 142.6 (Ar), 143.8 (Ar), 157.6 (CH_{pyr}), 160.0 (CH_{pyr}), 160.8 (CO), 162.4 (COOR). Measured monoisotopic mass [C₁₃H₁₀O₅ + H]⁺: m/z 247.06009 (Δ: -0.04 ppm) – theoretical monoisotopic mass: m/z 247.06010.

4-Chlorophenyl 2-oxo-2H-pyran-5-carboxylate (7). Synthesized according to the **general procedure B** using 4-chlorophenol as starting material to afford the titled compound 7 as a white solid (61% yield). 1 H NMR (400 MHz, DMSO- d_6): δ 6.51 (dd, J=9.8, 1.1 Hz, 1H, CH_{Pyr}), 7.25–7.44 (m, 2H, Ar), 7.47–7.69 (m, 2H, Ar), 7.94 (dd, J=9.8, 2.7 Hz, 1H, CH_{Pyr}), 8.88 (dd, J=2.7, 1.1 Hz, 1H, CH_{Pyr}). 13 C NMR (101 MHz, DMSO- d_6): δ 111.3 (CH_{Pyr}), 115.3 (C_{Pyr}), 124.2 (2 × Ar), 130.0 (2 × Ar), 130.8 (Ar), 142.5 (Ar), 149.2 (CH_{Pyr}), 159.9 (CH_{Pyr}), 161.1 (CO), 162.0 (COOR). Measured monoisotopic mass [C₁₂H₇ClO₄ + H]⁺: m/z 251.01050 (Δ: -0.24 ppm) – theoretical monoisotopic mass: m/z 251.01056.

4-Bromophenyl 2-oxo-2H-pyran-5-carboxylate (8). Synthetized according to the **general procedure B** using

4-bromophenol as starting material to afford the titled compound **8** as a white solid (67% yield). ¹H NMR (400 MHz, DMSO- d_6): δ 6.51 (dd, J = 9.8, 1.1 Hz, 1H, CH_{Pyr}), 7.24–7.29 (m, 2H, Ar), 7.63–7.70 (m, 2H, Ar), 7.93 (dd, J = 9.8, 2.7 Hz, 1H, CH_{Pyr}), 8.87 (dd, J = 2.7, 1.1 Hz, 1H, CH_{Pyr}). ¹³C NMR (101 MHz, DMSO- d_6): δ 111.3 (CH_{Pyr}), 115.3 (C_{Pyr}), 119.0 (Ar), 124.6 (2 × Ar), 132.9 (2 × Ar), 142.5 (Ar), 149.7 (CH_{Pyr}), 159.9 (CH_{Pyr}), 161.1 (CO), 161.9 (COOR). Measured monoisotopic mass [C₁₂H₇BrO₄ + H]⁺: m/z 249.96006 (Δ : 0.03 ppm) – theoretical monoisotopic mass: m/z 294.96005.

[1,1'-Biphenyl]-4-yl 2-oxo-2H-pyran-5-carboxylate (9). Synthetized according to the **general procedure B** using [1,1'-biphenyl]-4-ol as starting material to afford the titled compound 9 as a white solid (60% yield). δ ¹H NMR (400 MHz, DMSO-d₆): δ 6.55–6.48 (m, 1H, CH_{Pyr}), 7.43–7.32 (m, 3H, Ar), 7.48 (t, J = 7.5 Hz, 2H, Ar), 7.72–7.66 (m, 2H, Ar), 7.79–7.72 (m, 2H, Ar), 7.96 (dd, J = 9.8, 2.7 Hz, 1H, CH_{Pyr}), 8.91–8.85 (m, 1H, CH_{Pyr}). ¹³C NMR (101 MHz, DMSO-d₆): δ 111.5 (C_{Pyr}), 115.3 (CH_{Pyr}), 122.7 (2 × Ar), 127.1 (2 × Ar), 128.1 (Ar), 128.3 (2 × Ar), 129.4 (2 × Ar), 138.7 (Ar), 139.7 (Ar) 142.6 (CH_{Pyr}), 150.0 (Ar), 160.1 (CH_{Pyr}), 161.0 (COOR), 162.2 (CO). Measured monoisotopic mass [C₁₈H₁₂O₄ + H]⁺: m/z 293.08081 (Δ : -0.10 ppm) – theoretical monoisotopic mass: m/z 293.08084.

Naphthalen-2-yl 2-oxo-2H-pyran-5-carboxylate (10). Synthetized according to the **general procedure B** using naphthalen-2-ol as starting material to afford the titled compound **10** as a white solid (60% yield). ¹H NMR (400 MHz, DMSO-d₆): δ 6.53 (dd, J = 9.8, 1.1 Hz, 1H, CH_{Pyr}), 7.45 (dd, J = 8.9, 2.4 Hz, 1H, CH_{Pyr}), 7.57 (tt, J = 7.0, 5.2 Hz, 3H, Ar), 7.83 (d, J = 2.3 Hz, 1H, Ar), 7.91–8.08 (m, 3H, Ar), 8.86–9.01 (m, 1H, CH_{Pyr}). ¹³C NMR (101 MHz, DMSO-d₆): δ 111.5 (C_{Pyr}), 115.3 (CH_{Pyr}), 119.1 (Ar), 121.8 (Ar), 126.5 (Ar), 127.3 (Ar), 128.0 (Ar), 128.2 (Ar), 129.9 (Ar), 131.6 (Ar), 133.7 (Ar), 142.5 (Ar), 148.1 (CH_{Pyr}), 160.0 (CH_{Pyr}), 161.0 (CO), 162.4 (COOR). Measured monoisotopic mass [C₁₆H₁₀O₄ + H]⁺: m/z

267.06517 (Δ : -0.07 ppm) – theoretical monoisotopic mass: m/z 267.06519.

Cyclohexyl 2-oxo-2H-pyran-5-carboxylate (11). Synthetized according to the **general procedure B** using cyclohexanol as starting material to afford the titled compound **11** as a white solid (58% yield). ¹H NMR (400 MHz, DMSO-d₆): δ 1.24–1.43 (m, 3H, CH_{2Chx}), 1.46–1.55 (m, 3H, CH_{2Chx}), 1.67–1.87 (m, 4H, CH_{2Chx}), 4.89 (tt, J = 8.3, 3.7 Hz, 1H, CH_{Chx}), 6.43 (dd, J = 9.8, 1.3 Hz, 1H, CH_{Pyr}), 7.82 (dd, J = 9.8, 2.6 Hz, 1H, CH_{Pyr}), 8.59 (dd, J = 2.7, 1.3 Hz, 1H, CH_{Pyr}). ¹³C NMR (101 MHz, DMSO-d₆): δ 23.3 (2 × CH_{2Chx}), 25.3 (CH_{2Chx}), 31.3 (2 × CH_{2Chx}), 73.5 (CH_{2Chx}), 112.2 (C_{Pyr}), 115.2 (CH_{Pyr}), 142.5 (CH_{Pyr}), 159.5 (CH_{Pyr}), 160.2 (CO), 162.6 (COOR). Measured monoisotopic mass [C₁₂H₁₄O₄ + H]⁺: m/z 233.09650 (Δ: 0.04 ppm) – theoretical monoisotopic mass: m/z 233.09649.

2-Methylcyclohexyl 2-oxo-2H-pyran-5-carboxylate (12). Synthetized according to the general procedure B using 2-methylcyclohexanol as starting material to afford the titled compound 12 as a white solid (22% yield). 1H NMR (400 MHz, DMSO- d_6): δ 0.87 (dd, J = 9.8, 6.6 Hz, 3H, CH₃), 1.04-1.16 (m, 1H, CH_{2Chx}), 1.22 (dt, J = 12.5, 3.5 Hz, 1H, CH_{2Chx}), 1.29-1.37 (m, 1H, CH_{2Chx}), 1.47 (dq, J = 8.7, 4.4 Hz, 1H, CH_{2Chx}), 1.57-1.68 (m, 2H, CH_{2Chx}), 1.74 (tt, J = 9.1, 4.1 Hz, 2H, CH_{2Chx}), 1.91–1.98 (m, 1H, CH_{Chx}), 4.52 (td, J = 9.9, 4.4 Hz, 1H, CH_{Chx}), 6.44 (dd, J = 9.8, 4.0 Hz, 1H, CH_{Pyr}), 7.82 (dd, J = 9.7, 2.7 Hz, 1H, CH_{Pvr}), 8.61 (dd, J = 11.1, 2.6 Hz, 1H, CH_{Pyr}). 13 C NMR (101 MHz, DMSO- d_6): δ 18.7 (CH₃), 24.5 (2 × CH_{2Chx}), 25.1 (2 × CH_{2Chx}), 31.5 (2 × CH_{2Chx}), 33.1 (2 × CH_{2Chx}), 37.0 (CH_{Chx}), 79.4 (CH_{Chx}), 112.0 (C_{Pvr}) , 115.2 (CH_{Pvr}) , 142.5 (CH_{Pvr}) , 159.5 (CH_{Pvr}) , 160.1 (CO), 162.9 (COOR). Measured monoisotopic mass $[C_{13}H_{16}O_4 + H]^+$: m/z 237.11216 (Δ : 0.08 ppm) theoretical monoisotopic mass: m/z 237.11214.

Ethyl 2-oxo-2H-pyran-5-carboxylate (13). Synthetized according to the general procedure B using ethanol as starting material to afford the titled compound 13 as a white solid (53% yield). 1 H NMR (400 MHz, DMSO-d₆): δ 1.27 (t, J = 7.1 Hz, 3H, CH₃), 4.26 (q, J = 7.1 Hz, 2H, CH₂), 6.42 (dd, J = 9.8, 1.1 Hz, 1H, CH_{Pvr}), 7.81 (dd, J = 9.8, 2.6 Hz, 1H, CH_{Pyr}), 8.57 (dd, J = 2.6, 1.2 Hz, 1H, CH_{Pyr}). ¹³C NMR (101 MHz, DMSO-d₆): δ 14.4 (CH₃), 61.5 (CH₂), 112.0 (C_{Pvr}) , 115.2 (CH_{Pvr}) , 142.5 (CH_{Pvr}) , 159.6 (CH_{Pvr}) , 160.1 (CO), 163.2 (COOR). Measured monoisotopic mass $[C_8H_8O_4 + H]^{\dagger}$: m/z 169.04955 (Δ : 0.06 ppm) – theoretical monoisotopic mass: m/z 169.04954.

Prop-2-yn-1-yl 2-oxo-2H-pyran-5-carboxylate (14). To 1 eq. of intermediate A, were added 2 eq. of propargyl alcohol under inert atmosphere (N_2) at T = 75 °C. The mixture was stirred for 12 h and quenched with slush. The mixture was extracted with dichloromethane (3 × 40 mL), the organics were reunited dried over sodium sulfate and filtered to remove the drying agent. The organic phase was evaporated in vacuo and the crude purified by silica gel column chromatography eluting with 33% ethyl acetate in n-hexane to afford the titled compound 14 as a white solid (yield 58%, mp 80-84 °C). ¹H NMR (400 MHz, DMSO-d₆): δ 3.64 (q, J = 2.2 Hz, 1H, CH), 4.91 (t, J = 1.8 Hz, 2H, CH₂), 6.44 (d, J = 9.8Hz, 1H, CH_{Pvr}), 7.82 (dt, J = 9.8, 1.9 Hz, 1H, CH_{Pvr}), 8.55–8.73 (m, 1H, CH_{Pvr}). ¹³C NMR (101 MHz, DMSO-d₆): δ 53.2 (CH₂), 78.4 (C≡CH), 78.8 (C≡CH), 111.3 (C_{Pyr}), 115.3 (CH_{Pyr}), 142.3 (CH_{Pyr}), 160.0 (CH_{Pyr}), 160.2 (CO), 162.6 (COOR). Measured monoisotopic mass $[C_9H_6O_4 + H]^+$: m/z 179.03392 (Δ : 0.17 ppm) – theoretical monoisotopic mass: m/z 179.03389.

General procedure C. To a mixture of CuI (0.01 eq.), DIPEA (0.1 eq.), and CH₃COOH (0.1 eq.) in dichloromethane was added 2-oxo-N-(prop-2-yn-1-yl)-2H-pyran-5-carboxamide (14, 1 eq.) or prop-2-yn-1-yl 2-oxo-2*H*-pyran-5-carboxylate (20, 1 eq.) dissolved in dichloromethane, at room temperature. The solution was left to stir for 10 minutes and then the proper azide (1.05 eq.), was synthesized as previously reported.³⁶ The resultant mixture was stirred for 12 h (TLC monitoring). Then depending on compound behavior, filtration or extraction

was performed. The derivatives producing precipitate were filtered and purified by silica gel column chromatography eluting with 10% ethyl acetate in n-hexane to afford the final compound. The derivatives that did not precipitate were extracted with dichloromethane (3 × 40 mL), the organics were reunited, dried over sodium sulfate and filtered to remove the drying agent. The organic phase was evaporated in vacuo and the crude was purified with chromatographic procedures on silica gel employing 10% ethyl acetate in *n*-hexane as mobile phase.

(1-Phenyl-1H-1,2,3-triazol-4-yl)methyl 2-oxo-2H-pyran-5carboxylate (15). Synthetized according to the general procedure D to afford the titled compound 15 as a bright brown solid (37% yield). 1 H NMR (400 MHz, CDCl₃-d): δ 5.51 (s, 2H, CH₂), 6.34 (dd, J = 9.8, 1.1 Hz, 1H, CH_{Pyr}), 7.45-7.50 (m, 1H, Ar), 7.51-7.58 (m, 2H, Ar), 7.72-7.76 (m, 2H, Ar), 7.79 (dd, J = 9.9, 2.6 Hz, 1H, CH_{Pvr}), 8.10 (s, 1H, CH_{Triaz}), 8.34 (dd, J = 2.6, 1.1 Hz, 1H, CH_{Pyr}). ¹³C NMR (101 MHz, CDCl₃-d): δ 58.3 (CH₂), 111.7 (C_{Pyr}), 115.4 (CH_{Pyr}), 120.7 (2 \times Ar), 122.5 (CH_{Triaz}), 129.2 (Ar), 129.9 (2 \times Ar), 136.8 (Ar), 141.5 (CH_{Pyr}), 142.7 (C_{Triaz}), 158.7 (CH_{Pyr}), 159.6 (CO_{Pyr}), 163.0 (COOR). Measured monoisotopic mass $[C_{15}H_{11}N_3O_4 + H]^+$: m/z 298.08228 (Δ : 0.17 ppm) theoretical monoisotopic mass: m/z 298.08223.

(1-(p-Tolyl)-1H-1,2,3-triazol-4-yl)methyl2-oxo-2H-pyran-5carboxylate (16). Synthetized according to the general procedure D to afford the titled compound 16 as a white solid (28% yield). ¹H NMR (400 MHz, CDCl₃-d): δ 2.42 (s, 3H, CH₃), 5.49 (s, 2H, CH₂), 6.33 (dd, J = 9.9, 1.1 Hz, 1H, CH_{Pyr}), 7.28-7.42 (m, 2H, Ar), 7.54–7.67 (m, 2H, Ar), 7.78 (dd, J = 9.9, 2.6 Hz, 1H, CH_{Pvr}), 8.06 (s, 1H, CH_{Triaz}), 8.33 (dd, J = 2.6, 1.1 Hz, 1H, CH_{Pyr}). 13 C NMR (101 MHz, CDCl₃-d): δ 21.1 (CH₃), 58.4 (CH_2) , 111.7 (C_{Pyr}) , 115.4 (CH_{Pyr}) , 120.6 $(2 \times Ar)$, 122.5

(CH_{Triaz}), 130.4 (2 × Ar), 134.5 (Ar), 139.3 (Ar), 141.5 (CH_{Pyr}), 142.6 (C_{Triaz}), 158.7 (CH_{Pyr}), 159.6 (CO_{Pyr}), 163.0 (COOR). Measured monoisotopic mass [$C_{16}H_{13}N_3O_4 + H$]⁺: m/z 312.09794 (Δ : 0.19 ppm) – theoretical monoisotopic mass: m/z 312.09788.

(1-(4-Methoxyphenyl)-1H-1,2,3-triazol-4-yl)methyl 2-oxo-2H-pyran-5-carboxylate (17). Synthetized according to the **general procedure D** to afford the titled compound 17 as an orange solid (26% yield). 1 H NMR (400 MHz, CDCl₃-d): δ 3.87 (s, 3H, CH₃), 5.49 (s, 2H, CH₂), 6.33 (dd, J = 9.9, 1.2 Hz, 1H, CH_{Pyr}), 6.97–7.09 (m, 2H, Ar), 7.58–7.67 (m, 2H, Ar), 7.78 (dd, J = 9.8, 2.7 Hz, 1H, CH_{Pyr}), 8.01 (s, 1H, CH_{Triaz}), 8.33 (dd, J = 2.6, 1.1 Hz, 1H, CH_{Pyr}). 13 C NMR (101 MHz, CDCl₃-d): δ 55.7 (CH₃), 58.4 (CH₂), 111.7 (C_{Pyr}), 114.9 (2 × Ar), 115.4 (CH_{Pyr}), 122.3 (2 × Ar), 122.6 (CH_{Triaz}), 127.7 (Ar), 130.2 (Ar), 141.5 (CH_{Pyr}), 142.5 (C_{Triaz}), 158.7 (CH_{Pyr}), 159.6 (CO_{Pyr}), 160.1 (Ar), 163.0 (COOR). Measured monoisotopic mass [C₁₆H₁₃N₃O₅ + H]⁺: m/z 328.09281 (Δ: 0.03 ppm) – theoretical monoisotopic mass: m/z 328.09280.

(1-(4-Chlorophenyl)-1H-1,2,3-triazol-4-yl)methyl 2-oxo-2H-pyran-5-carboxylate (18). Synthetized according to the **general procedure D** to afford the titled compound 18 as a white solid (24% yield). ¹H NMR (400 MHz, CDCl₃-d): δ 5.50 (s, 2H, CH₂), 6.34 (dd, J = 9.8, 1.1 Hz, 1H, CH_{Pyr}), 7.48–7.58 (m, 2H, Ar), 7.66–7.73 (m, 2H, Ar), 7.78 (dd, J = 9.8, 2.6 Hz, 1H, CH_{Pyr}), 8.08 (s, 1H, CH_{Triaz}), 8.34 (dd, J = 2.6, 1.1 Hz, 1H, CH_{Pyr}). ¹³C NMR (101 MHz, CDCl₃-d): δ 58.2 (CH₂), 111.6 (C_{Pyr}), 115.4 (CH_{Pyr}), 121.8 (2 × Ar), 122.4 (CH_{Triaz}), 130.1 (2 × Ar), 135.0 (Ar), 135.3 (Ar), 141.4

(CH_{Pyr}), 143.0 (C_{Triaz}), 158.7 (CH_{Pyr}), 159.6 (CO_{Pyr}), 163.0 (COOR). Measured monoisotopic mass [C₁₅H₁₀ClN₃O₄ + H]⁺: m/z 332.04328 (Δ : 0.06 ppm) – theoretical monoisotopic mass: m/z 332.04326.

(1-(4-Nitrophenyl)-1H-1,2,3-triazol-4-yl)methyl 2-oxo-2H-pyran-5-carboxylate (19). Synthetized according to the **general procedure D** to afford the titled compound 19 as a white solid (15% yield). 1 H NMR (400 MHz, CDCl₃-d): δ 5.52 (s, 2H, CH₂), 6.34 (dd, J = 9.9, 1.1 Hz, 1H, CH_{pyr}), 7.78 (dd, J = 9.8, 2.6 Hz, 1H, CH_{pyr}), 7.94–8.05 (m, 2H, Ar), 8.23 (s, 1H, CH_{Triaz}), 8.34 (dd, J = 2.6, 1.1 Hz, 1H, CH_{pyr}), 8.40–8.47 (m, 2H, Ar). 13 C NMR (101 MHz, CDCl₃-d): δ 58.0 (CH₂), 111.5 (C_{pyr}), 115.5 (CH_{pyr}), 120.7 (2 × Ar), 122.4 (CH_{Triaz}), 125.7 (2 × Ar), 140.9 (Ar), 141.3 (CH_{pyr}), 143.7 (C_{Triaz}), 147.5 (Ar), 158.8 (CH_{pyr}), 159.5 (CO_{pyr}), 163.0 (COOR). Measured monoisotopic mass [C₁₅H₁₀N₄O₆ + H]⁺: m/z 343.06733 (Δ: 0.06 ppm) – theoretical monoisotopic mass: m/z 343.06731.

2-Oxo-N-(prop-2-yn-1-yl)-2H-pyran-5-carboxamide (20). To a solution of 2-oxo-2H-pyran-5-carbonyl chloride (A, 1 eq.) in diethyl ether, N,N-diisopropylethylamine (DIPEA, 1.5 eq.) and the propargyl amine (1.1 eq.) were added. The mixture was stirred at T = 4 °C (ice-bath) for 12 h (TLC monitoring). Then the reaction was added of diethyl ether and the organic phase extracted with HCl 2N (3 × 20 mL). The organic phase was dried over sodium sulfate, filtered to remove the drying agent and evaporated in vacuo. The crude has been purified by silica gel column chromatography eluting with 50% ethyl acetate in n-hexane, affording the titled compound 20 as a yellow powder (yield 51%, mp 131-135 °C). H NMR (400 MHz, DMSO- d_6): δ 3.14 (t, J = 2.5 Hz, 1H, CH), 4.01 (dd, J =5.4, 2.6 Hz, 2H, CH_2), 6.41 (dd, J = 9.8, 1.2 Hz, 1H, CH_{Pvr}), 7.86 (dd, J = 9.8, 2.8 Hz, 1H, CH_{Pvr}), 8.38 (dd, J =2.7, 1.2 Hz, 1H, CH_{PVI}), 8.86 (t, J = 5.5 Hz, 1H, NHCO).

¹³C NMR (101 MHz, DMSO- d_6): δ 28.4 (CH₂), 73.3 (C≡CH), 80.7 (C≡CH), 114.4 (C_{Pyr}), 114.6 (CH_{Pyr}), 142.1 (CH_{Pyr}), 155.7 (CH_{Pyr}), 160.0 (CO), 161.9 (NHCO). Measured monoisotopic mass [C₉H₇NO₃ + H]⁺: m/z 178.04989 (Δ: 0.11 ppm) – theoretical monoisotopic mass: m/z 178.04987.

2-Oxo-N-((1-phenyl-1H-1,2,3-triazol-4-yl)methyl)-2H-pyran-5-carboxamide (21). Synthetized according to the **general procedure** C to afford the titled compound 21 as a white solid (32% yield). ¹H NMR (400 MHz, DMSO- d_6): δ 3.74 (d, J = 5.6 Hz, 2H, CH₂), 5.51–5.66 (m, 1H, CH_{Pyr}), 6.60–6.67 (m, 1H, Ar), 6.69–6.83 (m, 2H, Ar), 7.01–7.07 (m, 2H, Ar), 7.09 (dd, J = 9.8, 2.6 Hz, 1H, CH_{Pyr}), 7.59 (dd, J = 9.8, 2.6 Hz, 1H, CH_{Triaz}), 8.17 (t, J = 5.5 Hz, 1H, NHCO). ¹³C NMR (101 MHz, DMSO- d_6): δ 35.1 (CH₂), 114.9 (C_{Pyr}), 115.2 (CH_{Pyr}), 120.4 (2 × Ar), 121.5 (CH_{Triaz}), 128.9 (Ar), 130.2 (2 × Ar), 137.2 (Ar), 142.8 (CH_{Pyr}), 146.1 (C_{Triaz}), 155.9 (CH_{Pyr}), 160.5 (CO_{Pyr}), 162.6 (NHCO). Measured monoisotopic mass [C₁₅H₁₂N₄O₃ + H]⁺: m/z 297.09827 (Δ: 0.17 ppm) – theoretical monoisotopic mass: m/z 297.09822.

2-Oxo-N-((1-(p-tolyl)-1H-1,2,3-triazol-4-yl)methyl)-2H-pyran-5-carboxamide (22). Synthetized according to the **general procedure** C to afford the titled compound 22 as a white solid (26% yield). 1 H NMR (400 MHz, DMSO- d_6): δ 2.37 (s, 3H, CH₃), 4.55 (d, J = 5.6 Hz, 2H, CH₂), 6.36–6.46 (m, 1H, CH_{Pyr}), 7.38 (d, J = 8.1 Hz, 2H, Ar), 7.76 (d, J = 8.4 Hz, 2H, Ar), 7.91 (dd, J = 9.8, 2.7 Hz, 1H, CH_{Pyr}), 8.38–8.50 (m, 1H, CH_{Pyr}), 8.63 (s, 1H, CH_{Triaz}), 9.00 (t, J = 5.6 Hz, 1H, NHCO). 13 C NMR (101 MHz, DMSO- d_6): δ 21.0 (CH₃), 35.1 (CH₂), 115.0 (C_{Pyr}), 115.2 (CH_{Pyr}), 120.3 (2 × Ar), 121.6 (CH_{Triaz}), 130.7 (2 × Ar), 134.9 (Ar), 138.7 (Ar), 142.9 (CH_{Pyr}), 146.1 (C_{Triaz}), 156.0 (CH_{Pyr}), 160.6 (CO_{Pyr}), 162.7 (NHCO). Measured monoisotopic mass [C₁₆H₁₄N₄O₃ + H]⁺:

m/z 311.11389 (Δ : 0.06 ppm) – theoretical monoisotopic mass: m/z 311.11387.

N-((1-(4-Chlorophenyl)-1H-1,2,3-triazol-4-yl)methyl)-2-oxo-2H-pyran-5-carboxamide (24). Synthetized according to the general procedure C to afford the titled compound 24 as a white solid (34% yield). ¹H NMR (400 MHz, CDCl₃): δ 4.56 (d, J = 5.6 Hz, 2H, CH₂), 6.42 (dd, J = 9.8, 1.1 Hz, 1H, CH_{Pyr}), 7.63–7.68 (m, 2H, Ar), 7.91 (dd, J = 9.8, 2.7 Hz, 1H, CH_{Pyr}), 7.93–7.97 (m, 2H, Ar), 8.42 (dd, J = 2.7, 1.1 Hz, 1H, CH_{Pyr}), 8.72 (s, 1H, CH_{Triaz}), 9.02 (t, J = 5.6 Hz, 1H, NHCO). ¹³C NMR (101 MHz, CDCl₃): δ 35.1 (CH₂), 115.0 (C_{Pyr}), 115.2 (CH_{Pyr}), 121.8 (CH_{Triaz}), 122.1 (2 × Ar), 130.3 (2 × Ar), 133.4 (Ar), 135.9 (Ar), 142.9 (C_{Triaz}), 146.4 (CH_{Pyr}), 156.0 (CH_{Pyr}), 160.6 (CO_{Pyr}), 162.7 (NHCO). Measured monoisotopic mass [C₁₅H₁₁ClN₄O₃ + H]⁺: m/z 331.05931 (Δ : 0.21 ppm) – theoretical monoisotopic mass: m/z 331.05924.

N-((1-(4-Methoxyphenyl)-1H-1,2,3-triazol-4-yl)methyl)-2-oxo-2H-pyran-5-carboxamide (23). Synthetized according to the general procedure C to afford the titled compound 23 as an orange solid (16% yield). 1 H NMR (400 MHz, DMSO- d_6): δ 3.82 (s, 3H, CH₃), 4.54 (d, J = 5.6 Hz, 2H, CH₂), 6.40 (dd, J = 9.8, 1.1 Hz, 1H, CH_{Pyr}), 7.04–7.18 (m, 2H, Ar), 7.73–7.85 (m, 2H, Ar), 7.90 (dd, J = 9.8, 2.7 Hz, 1H, CH_{Pyr}), 8.41 (dd, J = 2.7, 1.1 Hz, 1H, CH_{Pyr}), 8.55 (s, 1H, CH_{Triaz}), 8.98 (t, J = 5.6 Hz, 1H, NHCO). 13 C NMR (101 MHz, DMSO- d_6): δ 35.1 (CH₂), 56.0 (CH₃), 114.9 (C_{Pyr}), 115.2 (CH_{Pyr}), 115.3 (2 × Ar), 121.7 (CH_{Triaz}), 122.0 (2 × Ar), 130.6 (Ar), 142.8 (CH_{Pyr}), 145.9 (C_{Triaz}), 156.0 (Ar), 159.7 (CH_{Pyr}), 160.5 (CO_{Pyr}), 162.6 (NHCO). Measured monoisotopic mass [C₁₆H₁₄N₄O₄ + H]⁺: m/z

327.10890 (Δ : 0.37 ppm) – theoretical monoisotopic mass: m/z 327.10878.

N-((1-(4-Nitrophenyl)-1H-1,2,3-triazol-4-yl)methyl)-2-oxo-2H-pyran-5-carboxamide (25). Synthetized according to the general procedure C to afford the titled compound 25 as a white solid (25% yield). 1 H NMR (400 MHz, DMSO- d_6): δ 4.58 (d, J = 5.6 Hz, 2H, CH₂), 6.41 (dd, J = 9.8, 1.1 Hz, 1H, CH_{Pyr}), 7.91 (dd, J = 9.8, 2.7 Hz, 1H, CH_{Pyr}), 8.23 (d, J = 9.1 Hz, 2H, Ar), 8.28–8.51 (m, 3H, Ar + H_{Triaz}), 8.89 (s, 1H, CH_{Pyr}), 9.05 (t, J = 5.6 Hz, 1H, NHCO). 13 C NMR (101 MHz, DMSO- d_6): δ 35.0 (CH₂), 115.0 (C_{Pyr}), 115.2 (CH_{Pyr}), 120.9 (2 × Ar), 122.2 (Ar), 126.0 (2 × Ar), 141.4 (C_{Triaz}), 142.8 (CH_{Triaz}), 147.0 (Ar), 147.1 (CH_{Pyr}), 156.0 (CH_{Pyr}), 160.5 (CO), 162.7 (NHCO). Measured monoisotopic mass [C₁₅H₁₁N₅O₅ + H]⁺: m/z 342.08330 (Δ : 0.03 ppm) – theoretical monoisotopic mass: m/z 342.08329.

5.2 Inhibition assay

Research Article

An Applied Photophysics stopped-flow instrument was used to assay the CA-catalyzed CO2 hydration activity. 42 Phenol red (at a concentration of 0.2 mM) was used as an indicator, working at the absorbance maximum of 557 nm, with 20 mM Hepes (pH 7.4) as a buffer and 20 mM Na₂SO₄ (to maintain constant ionic strength), following the initial rates of the CA-catalyzed CO2 hydration reaction for a period of 10-100 s. The CO₂ concentrations ranged from 1.7 to 17 mM for the determination of the kinetic parameters and inhibition constant. Enzyme concentrations ranged between 5-12 nM. For each inhibitor, at least six traces of the initial 5-10% of the reaction were used to determine the initial velocity. The uncatalyzed rates were determined in the same manner and subtracted from the total observed rates. Stock solutions of the inhibitor (0.1 mM) were prepared in distilled-deionized water and dilutions up to 0.01 nM were conducted thereafter with the assay buffer. Inhibitor and enzyme solutions were preincubated together for 6 h at room temperature prior to the assay, to allow for the formation of the E-I complex. The inhibition constants were obtained by non-linear least-squares methods using PRISM 3 and the Cheng-Prusoff equation as reported earlier and represent the mean from at least three different determinations.

5.3 Molecular modelling

The crystallographic structures of hCA IX at 2.2 Å resolution (PDB-ID: 3IAI, chain A)⁴⁵ and hCA XII at 1.50 Å resolution (PDB-ID: 1JD0, chain A),⁴⁶ both in complex with acetazolamide (AAZ), were used as rigid receptors in molecular docking simulations. In hCA IX, sequence alignment with human hCA IX deposited in the UniProt database (Q16790)⁴⁷ revealed the C14S single point mutation compared to the hCA IX crystallized sequence. However, this mutation is at a distance higher than 20 Å from the catalytic site, thus it was not further accounted for in molecular docking simulations as in previous studies.36 The PDB sequence of PDB-ID: 1JD0 matches the UniProt sequence O43570, so no mutations are present. Small molecules were sketched in 2D with the Picto software (version 4.5.4.1, OpenEye Cadence Molecular Sciences, Santa Fe, NM) and converted into a 3D structure with OMEGA (version 4.2.0.1, OpenEye Cadence Molecular Sciences, Santa Fe, NM). 48,49 Ligand protonation state was assigned by QUACPAC (version 2.2.0.1, OpenEye Cadence Molecular Sciences, Santa Fe, NM),⁵⁰ while energy minimization was carried out by SZYBKI (version 2.5.0.1, OpenEye Cadence Molecular Sciences, Santa FE, NM)⁵¹ using the MMFF94S force field.^{52,53} Molecular docking was carried out with GOLD (The Cambridge Crystallographic Data Centre, Cambridge, UK) version 2023.1.54,55 In both structures, re-docking of AAZ was performed to validate the docking protocol, and the obtained pose overlaps with the co-crystalized AAZ (RMSD < 2.0 Å) in both crystals with the same docking settings. The ChemPLP fitness function for docking and scoring was used. Based on structural and literature evidence, the binding site for docking simulations was centred on the catalytic Zn(II) ion, and it was assigned a radius of 10 Å. The Zn(II)-coordinated water molecule was modelled as described previously,56,57 and it was kept as fixed in molecular docking (toggle "on" option). The receptor was prepared with the GOLD docking wizard implemented in Hermes visualizer, which is an integral part of the GOLD interface. Ten runs for each ligand were stored and submitted to visual inspection.

6. Conclusion

In this study, a series of novel coumalic acid derivatives was synthesized and the compounds were evaluated for their inhibitory activity against human carbonic anhydrase isoforms, particularly the tumour-associated isoforms hCA IX and hCA XII. The results from this work contribute valuable insights into the structure-activity relationship (SAR) of coumalic acid-based inhibitors, with the goal of developing selective inhibitors for cancer therapy. The key findings from the SAR analysis highlight the critical role of the lactone moiety in the inhibition of hCA IX and XII. The replacement of the lactone with a lactam (compounds 1–2) resulted in a complete loss of activity, confirming the importance of the lactone ring and its open form (cinnamic acid) in enzyme inhibition. Furthermore, the position of the carboxylic acid

group within the coumalic acid core was found to significantly influence inhibitory potency. Indeed, the moving this group from the position 5 of the coumalic acid to position 6 (compound 3) provoked the dramatic loss of inhibitory activity against isoforms IX and XII, emphasizing the structural precision required for effective binding to the enzyme active site. The modifications of the linker moiety, particularly the isosteric replacement of amide with ester (compounds 4-14), demonstrated a clear reduction in potency against hCA IX and XII, although selectivity for these isoforms was maintained. Notably, the introduction of substituents on the phenyl ring (compounds 5-10) enhanced inhibitory activity, with compound 8 emerging as the most potent inhibitor of hCA XII, showcasing its potential as a lead compound for further development. The incorporation of a triazole ring as part of a "click chemistry" approach (compounds 15-25) aimed at molecular elongation did not yield compounds with improved potency, suggesting that excessive structural complexity may hinder the desired inhibitory effect. However, compounds 16 and 25 did show some promise, indicating that further optimization of the triazole-linked derivatives could still provide valuable insights. Molecular docking studies supported experimental results, showing that the coumalic acid derivatives bind effectively to the active sites of hCA IX and XII, with key interactions involving the Zn(II)-bound water molecule, critical for enzyme inhibition. In particular, compound 11 demonstrated a higher selectivity and potency for hCA XII, reinforcing the importance of specific structural features in achieving selective inhibition. In conclusion, the coumalic acid-based derivatives described in this study offer valuable starting points for the development of selective inhibitors targeting tumour-related carbonic anhydrase isoforms. These findings provide a deeper understanding of the molecular mechanisms underlying hCA inhibition and suggest potential avenues for further refinement of these compounds as therapeutic agents for cancer treatment. Future work should focus on optimizing the lead compounds identified in this study, exploring additional modifications to enhance their potency, selectivity, and pharmacological

Data availability

properties.

The data supporting this article have been included as part of the ESI.†

Author contributions

Conceptualization, review and editing: P. G., V. P., C. T. S., D. S., and P. C. Supervision, P. G., D. S., C. T. S. Investigation: V. P., F. A., M. C., E. D. F., A. B., M. M., M. S., L. C., A. A., E. B., C. S., A. T., F. P., A. D. N. and A. G. Writing - original draft preparation: P. G., V. P., M. M., and L. C. Writing - review & editing: S. C., C. T. S., D. S., P. C., and A. A. All authors have read and agreed to the published version of the manuscript.

Conflicts of interest

The authors declare no conflict of interest.

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