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Pegylation approach applied to erlotinib–carbonic anhydrase inhibitors hybrids towards anticancer agents†

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Herein we report a first study on single molecular entities bearing both epidermal growth factor receptor (EGFR) and carbonic anhydrase (CA) inhibiting moieties as new tools for the management of hypoxic cancers. Specifically, we designed and synthesized a library of erlotinib (ERL)-based compounds bearing both the primary sulfonamide/coumarin moieties with the intent to selectively interfere with EGFR and CA targets respectively. The compounds obtained were investigated *in silico* and *in vitro* for their ability to interact with the appropriate targets followed by the assessment on selected compounds for the anti-proliferative activity using human (h) TNBC cell line MDA-MB-231. We are confident that the data provided in this study are fundamental for paving the way toward the development of multi-targeting molecular structures useful for the management of chronic diseases such as hypoxic tumors.

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1. Introduction

Over the years, the definition of cancer has evolved to reflect an increasing understanding of its overall complexity and heterogeneity,^{1,2} although data on its morbidity and mortality consistently underscore the ongoing Public Health emergency.^{3,4} The current gold standard in cancer treatment, which combines surgery, radiotherapy, and chemotherapy, often faces significant challenges, including limited effectiveness and the associated high failure rates.⁵ Recently, the development of immunotherapy and gene-based

therapeutics has emerged in response to the urgent need for new effective treatment options, aligning with the rising trend towards personalized medicine.⁶ Conversely, the more traditional approach of small molecules in cancer treatment still focuses on modulating enzymatic activities that are crucial for the disease progression and the novelty of the mode of action, in the attempt to avoid cross-resistance phenomena.^{7–9} In particular, a valuable strategy consists of anticancer agents endowed with multi-targeted mode of action^{10–12} and relies on hybrid compounds^{13–17} that combine multiple pharmacophoric features, enabling them to target various pathways and mechanisms simultaneously,^{18,19} often spaced by simple flexible linkers as methylene units.²⁰ In this context, several known anticancer drugs undergo this chemical modification^{21–24} and, among them, one of the most employed is erlotinib (ERL, Fig. 1).^{25–32} This small molecule selectively inhibits the epidermal growth factor receptor (EGFR) tyrosine kinase, which is a key player in cell proliferation and disease progression in various cancers, such as non-small cell lung cancer (NSCLC) and pancreatic cancer.^{33,34} Mechanistically, ERL binds reversibly the tyrosine kinase domain of EGFR, specifically to the ATP-binding site in a competitive manner,³⁵ thereby, blocking the phosphorylation of EGFR and, consequently, the signaling pathways of PI3K–AKT and RAS–RAF–MEK–ERK,^{36–38} mainly involved in tumor growth, survival, and metastasis, along with increased aggressiveness and poor prognosis.^{39–41} Interestingly, although the binding to the targeted receptor is

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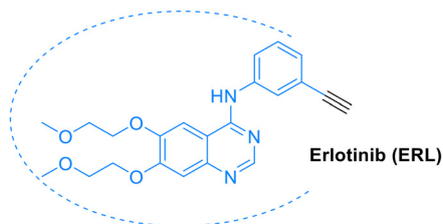
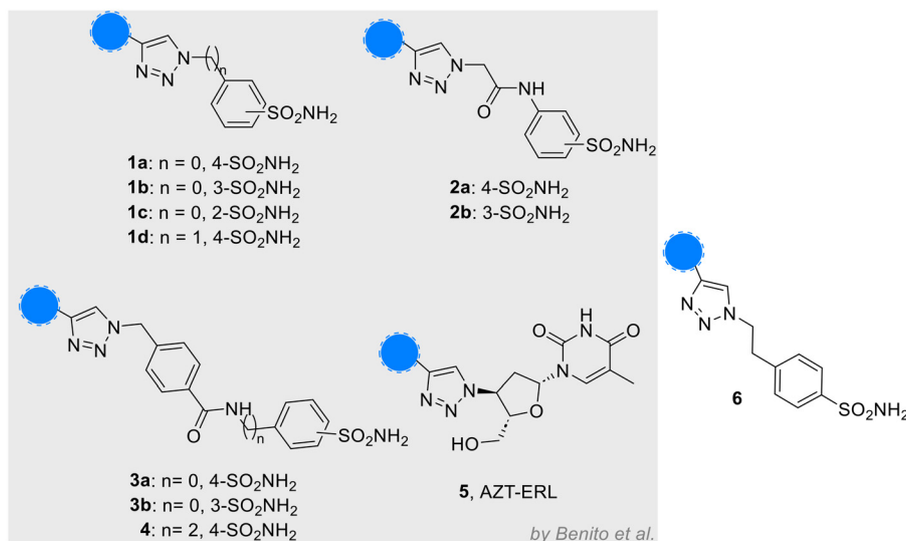
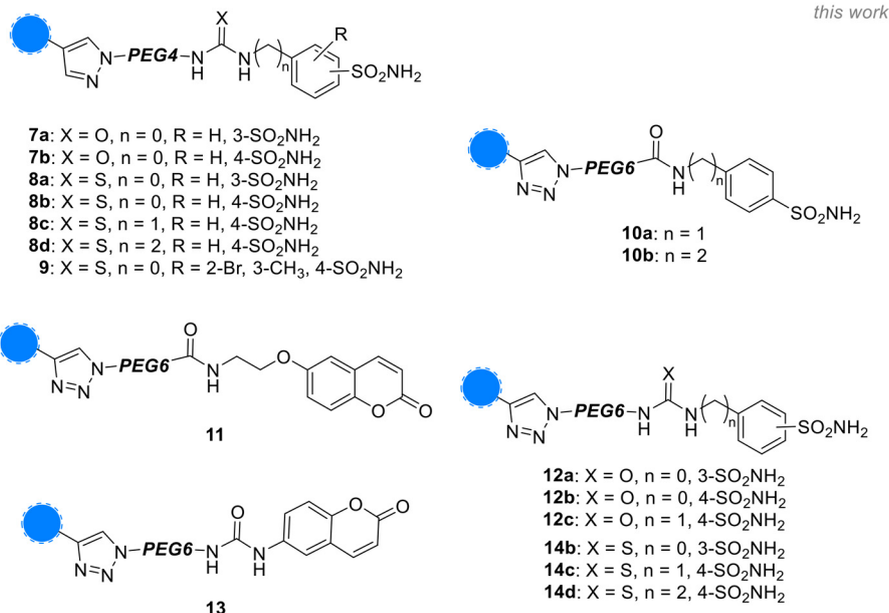
1st series of ERL-clicked derivatives (Hybrids)2nd series of ERL-clicked derivatives (Pegylated Hybrids)

Fig. 1 Structures of ERL and clicked derivatives from the 1st (1–6) and 2nd (pegylated 7–14) series.

governed by relevant networks of interactions also including hydrophobic contacts between the triad of residues (Thr766, leu764, and Lys721) and the acetylene terminal function,^{42,43} the latter has often been modified to generate various libraries of compounds, being often employed as starting point for derivatizations^{29,30,44–46} with only slight decrease in

the inhibitory potency towards EGFR.⁴⁴ Specifically, it served for the copper-catalyzed azide-alkyne cycloaddition (CuAAC), a cornerstone of click chemistry, to form the characteristic 1,4-disubstituted 1,2,3-triazole. The introduction of the triazole ring, a known surrogate for amides, esters, and carboxylic acids,^{45,47,48} is reported to often increase the



affinity for the target, allowing the establishment of additional interactions, such as hydrogen bonds and dipole interactions.⁴⁹

On the other hand, recent findings highlighted the high potential of another family of enzymes in the frame of anticancer treatment development: carbonic anhydrases (CAs, EC: 4.2.1.1), being widely reported during the last century due to their ubiquitary nature and their pivotal role in cell life.⁵⁰ These metalloenzymes catalyze the interconversion of carbon dioxide (CO₂) to bicarbonate (HCO₃⁻) and proton (H⁺), thereby regulating pH and CO₂ homeostasis, electrolyte secretion in various tissues and organs, and other crucial biological processes.^{51–53} In particular, the α -CA subfamily is the only one among the identified eight genetically distinct ones (α -, β -, γ - δ -, ζ -, η -, θ -, and ι -CAs) found in humans and is composed of 16 members (I–IV, VA, VB, VI–XV).^{50,54} Both the isoforms I and II are physiologically abundant in red blood cells, playing a role in the CO₂ transport, whereas hCA II levels are high also in kidneys, assisting in HCO₃⁻-resorption and diuresis.^{55,56} Conversely, hCAs overexpression is often associated with different diseases,^{10,52,57} such as glaucoma for hCA II⁵⁸ and hypoxic cancer for isoforms IX⁵⁹ and XII,⁶⁰ to name but a few. In particular, the latter isozymes were highly exploited and several libraries of CA inhibitors (CAIs), especially bearing the sulfonamide or the coumarin CA inhibiting chemotypes, have been described till now.^{61–63}

2. Material and methods

2.1. Chemistry

2.1.1. General chemistry. Anhydrous solvents and all reagents were purchased from Merck srl, TCI, and Fluorochem. All reactions involving air- or moisture-sensitive compounds were performed under a nitrogen atmosphere using dried glassware and syringes techniques to transfer solutions. Nuclear magnetic resonance (¹H- and ¹³C-NMR) spectra were recorded using a Bruker Advance III 400 MHz spectrometer in DMSO-*d*₆. 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, quartet; m, multiplet; dd, doublet of doublets; bs, broad singlet; ap s, apparent singlet; ap d, apparent doublet; ap t, apparent triplet; ap q, apparent quartet. The assignment of exchangeable protons (OH and NH) was confirmed by the addition of D₂O. Analytical thin-layer chromatography (TLC) was carried out on Merck silica gel F-254 plates. Flash chromatography purifications were performed on Merck silica gel 60 (230–400 mesh ASTM) as the stationary phase and methanol/dichloromethane (MeOH/DCM) or ethyl acetate/hexane (EtOAc/Hex) was used as eluents. The solvents used in Mass Spectrometry (MS) measures were acetone, acetonitrile (Chromasolv grade), and 56 mQ water 18 M Ω , obtained from Millipore's simplicity system (Milan-Italy). The mass spectra were obtained using a Varian 1200 L triple

quadrupole system (Palo Alto, USA) equipped with Electrospray Source (ESI) operating in both positive and negative modes. Stock solutions of analytes were prepared in acetone at 1.0 mg mL⁻¹ and stored at 4 °C. Working solutions of each analyte were freshly prepared by diluting stock solutions in a mixture of mQ H₂O/ACN 1/1(v/v) up to a concentration of 1.0 μ g mL⁻¹. The MS spectra of each analyte were acquired by introducing, *via* a syringe pump at 10 L min⁻¹, the working solution. RawQdata were collected and processed by Varian Workstation Vers. 6.8.

2.1.2. General procedure for the synthesis of compounds 19a and 19b. The appropriate sulfonamide **16a** (ref. 64) or **16b** (ref. 65) (1 equiv.) and the commercial 2-(2-(2-(2-azidoethoxy)ethoxy)ethoxy)ethan-1-amine **15** (1.2 equiv.) were dissolved in ACN (10 mL) and the reaction mixture was stirred at reflux temperature overnight. Then, slush was added to quench the reaction and extracted with EtOAc thrice. The combined organic layers were dried over anhydrous Na₂SO₄, filtered, and evaporated at reduced pressure to yield compounds **19a** or **19b**.

3-(3-(2-(2-(2-(2-Azidoethoxy)ethoxy)ethoxy)ethyl)ureido)benzenesulfonamide (19a). Yellow oil. Yield: 65%. ¹H NMR (400 MHz, DMSO-*d*₆) δ (ppm): 8.00 (1H, s, NH, exchange with D₂O), 7.54 (1H, d, *J* = 7.50 Hz), 7.42 (t, 1H, *J* = 7.55 Hz, NH, exchange with D₂O), 7.19 (2H, t, *J* = 7.76 Hz), 6.79 (2H, bs, SO₂NH₂, exchange with D₂O), 6.28 (1H, t, *J* = 5.37 Hz), 3.62 (2H, t, *J* = 3.77 Hz), 3.58 (8H, m), 3.50 (2H, t, *J* = 5.37 Hz), 3.41 (2H, m), 3.30 (2H, m). ¹³C NMR (100 MHz, DMSO-*d*₆) δ (ppm): 154.3, 139.9, 136.2, 127.0, 124.8, 122.9, 119.8, 70.4, 70.3, 70.2, 70.1, 70.0, 69.7, 50.0, 41.5; MS (ESI positive) *m/z*: 417.15 [M + H]⁺. Elemental analysis: calculated C, 43.26; H, 5.81; N, 20.18; found C, 43.25; H, 5.82; N, 20.16.

4-(3-(2-(2-(2-(2-Azidoethoxy)ethoxy)ethoxy)ethyl)ureido)benzenesulfonamide (19b). Yellow oil. Yield: 60%. ¹H NMR (400 MHz, DMSO-*d*₆) δ (ppm): 8.02 (1H, s, NH, exchange with D₂O), 7.72 (2H, d, *J* = 7.89 Hz), 7.57 (2H, d, *J* = 7.89 Hz), 6.80 (2H, bs, SO₂NH₂, exchange with D₂O), 6.36 (1H, t, *J* = 5.56 Hz, NH, exchange with D₂O), 3.63 (3H, t, *J* = 4.79 Hz), 3.59 (8H, s), 3.42 (3H, m), 3.30 (2H, q, *J* = 6.12 Hz). ¹³C NMR (100 MHz, DMSO-*d*₆) δ (ppm): 154.3, 142.6, 136.5, 129.5, 129.4, 118.1, 118.0, 70.4, 70.3, 70.2, 70.1, 70.0, 69.7, 50.0, 41.5; MS (ESI positive) *m/z*: 417.15 [M + H]⁺. Elemental analysis: calculated C, 43.26; H, 5.81; N, 20.18; found C, 43.25; H, 5.82; N, 20.19.

2.1.3. General procedure for the synthesis of compounds 20a–d and 21. The appropriate sulfonamide **17a–d** (ref. 66–68) or **18** (ref. 69) (1 equiv.) and the commercial 2-(2-(2-(2-azidoethoxy)ethoxy)ethoxy)ethan-1-amine **15** (1.2 equiv.) were dissolved in ACN (10 mL) and the reaction mixture was stirred at room temperature overnight. Then, slush added to quench the reaction and extracted with EtOAc thrice. The combined organic layers were dried over anhydrous Na₂SO₄, filtered, and evaporated at reduced pressure to yield compounds **20a–d** and **21**.

3-(3-(2-(2-(2-(2-Azidoethoxy)ethoxy)ethoxy)ethyl)thioureido)benzenesulfonamide (20a). Yellow oil. Yield: 95%. ¹H NMR (400 MHz, DMSO-*d*₆) δ (ppm): 9.94 (1H, s, NH, exchange with



D₂O), 7.99 (2H, d, *J* = 18.6 Hz), 7.76 (1H, d, *J* = 7.11 Hz), 7.55 (2H, bs, SO₂NH₂, exchange with D₂O), 7.40 (1H, s, NH, exchange with D₂O), 4.06 (2H, d, *J* = 7.22 Hz), 3.60 (14H, m). ¹³C NMR (100 MHz, DMSO-*d*₆) δ (ppm): 179.5, 140.0, 137.4, 129.7, 127.1, 124.7, 123.3, 70.6, 70.5, 70.4, 70.2, 70.1, 70.0, 50.0, 45.3; MS (ESI positive) *m/z*: 433.13 [M + H]⁺. Elemental analysis: calculated C, 41.66; H, 5.59; N, 19.43; found C, 41.64; H, 5.58; N, 19.45.

4-(3-(2-(2-(2-(2-Azidoethoxy)ethoxy)ethoxy)ethyl)thioureido)benzenesulfonamide (**20b**). Yellow oil. Yield: 95%. ¹H NMR (400 MHz, DMSO-*d*₆) δ (ppm): 9.97 (1H, s, NH, exchange with D₂O), 8.01 (1H, s, NH, exchange with D₂O), 7.74 (4H, q, *J* = 8.07 Hz), 7.31 (2H, bs, SO₂NH₂, exchange with D₂O), 3.61 (16H, m). ¹³C NMR (100 MHz, DMSO-*d*₆) δ (ppm): 179.5, 141.7, 136.9, 129.6, 129.5, 122.9, 122.8, 70.5, 70.4, 70.3, 70.2, 70.1, 70.0, 50.0, 45.3; MS (ESI positive) *m/z*: 433.13 [M + H]⁺. Elemental analysis: calculated C, 41.66; H, 5.59; N, 19.43; found C, 41.65; H, 5.58; N, 19.44.

4-(15-Azido-3-thioxo-7,10,13-trioxa-2,4-diazapentadecyl)benzenesulfonamide (**20c**). Yellow oil. Yield: 97%. ¹H NMR (400 MHz, DMSO-*d*₆) δ (ppm): 8.10 (1H, s, NH, exchange with D₂O), 7.80 (2H, d, *J* = 7.59 Hz), 7.67 (1H, s, NH, exchange with D₂O), 7.47 (2H, d, *J* = 7.59 Hz), 7.35 (2H, bs, SO₂NH₂, exchange with D₂O), 4.77 (2H, s), 4.07 (2H, d, *J* = 6.51 Hz), 3.58 (14H, m). ¹³C NMR (100 MHz, DMSO-*d*₆) δ (ppm): 182.3, 141.7, 141.1, 128.3, 128.2, 127.3, 127.2, 70.6, 70.5, 70.4, 70.2, 70.1, 70.0, 50.8, 50.0, 45.3; MS (ESI positive) *m/z*: 447.14 [M + H]⁺. Elemental analysis: calculated C, 43.04; H, 5.87; N, 18.82; found C, 43.05; H, 5.86; N, 18.83.

4-(16-Azido-4-thioxo-8,11,14-trioxa-3,5-diazahexadecyl)benzenesulfonamide (**20d**). Yellow oil. Yield: 97%. ¹H NMR (400 MHz, DMSO-*d*₆) δ (ppm): 7.79 (2H, d, *J* = 7.59 Hz), 7.60 (1H, s, NH, exchange with D₂O), 7.54 (1H, s, NH, exchange with D₂O), 7.45 (2H, d, *J* = 7.59 Hz), 7.34 (2H, bs, SO₂NH₂, exchange with D₂O), 3.64 (5H, m), 3.57 (14H, m), 2.92 (2H, m). ¹³C NMR (100 MHz, DMSO-*d*₆) δ (ppm): 181.8, 142.6, 140.9, 128.4, 128.3, 128.1, 128.0, 70.5, 70.4, 70.3, 70.2, 70.1, 70.0, 50, 47.8, 45.3, 35.3; MS (ESI positive) *m/z*: 461.16 [M + H]⁺. Elemental analysis: calculated C, 44.33; H, 6.13; N, 18.25; found C, 44.34; H, 6.12; N, 18.26.

4-(3-(2-(2-(2-(2-azidoethoxy)ethoxy)ethoxy)ethyl)thioureido)-3-bromo-2-methylbenzenesulfonamide (**21**). Yellow oil. Yield: 75%. ¹H NMR (400 MHz, DMSO-*d*₆) δ (ppm): 9.40 (1H, s, NH, exchange with D₂O), 8.44 (1H, s), 8.05 (1H, s), 7.51 (2H, bs, SO₂NH₂, exchange with D₂O), 7.28 (1H, s, NH, exchange with D₂O), 4.14 (2H, q, *J* = 5.19 Hz), 3.61 (8H, m), 3.21 (5H, m). ¹³C NMR (100 MHz, DMSO-*d*₆) δ (ppm): 179.5, 140.3, 138.3, 137.1, 126.5, 126.0, 120.9, 70.5, 70.4, 70.1, 70.4, 70.0, 50.0, 45.3, 13.6; MS (ESI positive) *m/z*: 525.05 [M + H]⁺. Elemental analysis: calculated C, 36.57; H, 4.80; N, 15.99; found C, 36.58; H, 4.81; N, 15.97.

2.1.4. General procedure for the synthesis of compounds 7a-d, 8a-d, and 9. Erlotinib hydrochloride (ERL) (1 equiv.), copper nanosized (0.5 equiv.), tetramethylammonium chloride (4.5 equiv.) and the appropriate azide derivative **19-21** (1.2 equiv.) were dissolved in DMF (2.5 mL) and

stirred overnight at 60 °C. Then, the reaction mixture was filtered through a cake of Celite 521© and the filtrate was treated with slush. The crude was and extracted with DCM thrice. The combined organic layers were dried over anhydrous Na₂SO₄, filtered, and evaporated at reduced pressure. The crude material was purified by flash column chromatography (MeOH/DCM: 5:95), to yield compounds **7a-b**, **8a-d**, and **9**.

3-(3-(2-(2-(2-(2-(4-(3-((6,7-Bis(2-methoxyethoxy)quinazolin-4-yl)amino)phenyl)-1H-1,2,3-triazol-1-yl)ethoxy)ethoxy)ethoxy)ethyl)ureido)benzenesulfonamide (**7a**). Yellow oil. Yield: 5%. ¹H NMR (400 MHz, DMSO-*d*₆) δ (ppm): 9.58 (1H, s, NH, exchange with D₂O), 8.90 (1H, s), 8.54 (1H, s), 8.48 (1H, s), 8.28 (1H, s, NH, exchange with D₂O), 7.95 (2H, d, *J* = 10.52 Hz), 7.88 (1H, d, *J* = 7.94 Hz), 7.55 (1H, d, *J* = 7.57 Hz), 7.48 (2H, t, *J* = 8.31 Hz), 7.38 (1H, t, *J* = 7.79 Hz), 7.33 (1H, d, *J* = 7.74 Hz), 7.29 (2H, bs, SO₂NH₂, exchange with D₂O), 7.23 (1H, s), 6.24 (1H, t, *J* = 5.23 Hz), 4.58 (2H, t, *J* = 4.63 Hz), 4.31 (4H, m), 3.88 (2H, t, *J* = 4.72 Hz), 3.79 (2H, m), 3.75 (2H, m), 3.45 (16H, m), 3.21 (2H, m). ¹³C NMR (100 MHz, DMSO-*d*₆) δ (ppm): 156.4, 154.9, 153.6, 152.9, 148.1, 146.9, 146.2, 144.5, 140.9, 140.0, 131.0, 129.2, 129.0, 121.8, 121.7, 120.3, 118.7, 118.0, 114.4, 108.9, 108.1, 103.2, 101.8, 70.1, 70.0, 69.7, 69.6, 68.6, 68.3, 68.0, 58.4, 58.3, 49.6, 28.9; MS (ESI positive) *m/z*: 810.32 [M + H]⁺. Elemental analysis: calculated C, 54.87; H, 5.85; N, 15.57; found C, 54.88; H, 5.84; N, 15.55.

4-(3-(2-(2-(2-(2-(4-(3-((6,7-Bis(2-methoxyethoxy)quinazolin-4-yl)amino)phenyl)-1H-1,2,3-triazol-1-yl)ethoxy)ethoxy)ethoxy)ethyl)ureido)benzenesulfonamide (**7b**). Yellow oil. Yield: 19%. ¹H NMR (400 MHz, DMSO-*d*₆) δ (ppm): 9.59 (1H, s, NH, exchange with D₂O), 8.96 (1H, s), 8.55 (1H, s), 8.48 (1H, s), 8.27 (1H, s), 7.93 (1H, s, NH, exchange with D₂O), 7.88 (1H, d, *J* = 7.65 Hz), 7.65 (2H, d, *J* = 8.29 Hz), 7.55 (1H, d, *J* = 7.8 Hz), 7.51 (2H, d, *J* = 8.76 Hz), 7.46 (1H, t, *J* = 7.87 Hz), 7.23 (1H, s), 7.17 (2H, bs, SO₂NH₂, exchange with D₂O), 6.31 (1H, t, *J* = 5.4 Hz), 4.58 (2H, t, *J* = 4.94 Hz), 4.30 (4H, m), 3.87 (2H, t, *J* = 5.0 Hz), 3.79 (2H, m), 3.75 (2H, m), 3.52 (8H, m), 3.36 (6H, m), 3.21 (2H, m), 3.16 (2H, d, *J* = 5.17 Hz). ¹³C NMR (100 MHz, DMSO-*d*₆) δ (ppm): 156.4, 154.7, 153.6, 152.9, 148.1, 146.9, 146.2, 143.6, 140.0, 136.0, 131.0, 129.0, 126.8, 121.8, 121.7, 120.3, 118.7, 116.7, 108.9, 108.1, 103.2, 70.1, 70.0, 69.7, 69.6, 68.6, 68.3, 68.0, 58.4, 58.3, 49.6, 48.6; MS (ESI positive) *m/z*: 810.32 [M + H]⁺. Elemental analysis: calculated C, 54.87; H, 5.85; N, 15.57; found C, 54.89; H, 5.86; N, 15.56.

3-(3-(2-(2-(2-(2-(4-(3-((6,7-Bis(2-methoxyethoxy)quinazolin-4-yl)amino)phenyl)-1H-1,2,3-triazol-1-yl)ethoxy)ethoxy)ethoxy)ethyl)thioureido)benzenesulfonamide (**8a**). Yellow oil. Yield: 27%. ¹H NMR (400 MHz, DMSO-*d*₆) δ (ppm): 9.86 (1H, m, NH, exchange with D₂O), 9.57 (1H, s), 8.54 (1H, s), 8.48 (1H, m), 8.28 (1H, s, NH, exchange with D₂O), 7.96 (1H, m), 7.93 (1H, s), 7.88 (1H, d, *J* = 7.18 Hz), 7.70 (1H, d, *J* = 7.33 Hz), 7.53 (2H, t, *J* = 7.98 Hz), 7.46 (2H, m), 7.36 (2H, bs, SO₂NH₂, exchange with D₂O), 7.23 (1H, m), 4.59 (2H, t, *J* = 4.93 Hz), 4.30 (4H, m), 3.88 (2H, t, *J* = 5.19 Hz), 3.79 (3H, t, *J* = 4.94 Hz), 3.75 (3H, m), 3.50 (12H, m), 3.37 (4H, m). ¹³C NMR (100 MHz, DMSO-*d*₆) δ (ppm): 181.4, 180.6, 165.4, 156.4, 153.6,



153.0, 148.1, 147.0, 146.8, 146.2, 144.3, 140.0, 131.1, 134.7, 129.0, 125.7, 121.9, 121.8, 120.3, 118.8, 108.1, 103.2, 70.1, 70.0, 69.7, 69.6, 68.7, 68.4, 68.0, 58.4, 58.3, 49.6, 48.6, 43.6; MS (ESI positive) m/z : 826.30 $[M + H]^+$. Elemental analysis: calculated C, 53.81; H, 5.74; N, 15.26; found C, 53.82; H, 5.73; N, 15.24.

4-(3-(2-(2-(2-(4-(3-((6,7-Bis(2-methoxyethoxy)quinazolin-4-yl)amino)phenyl)-1H-1,2,3-triazol-1-yl)ethoxy)ethoxy)ethoxy)ethyl)thioureido)benzenesulfonamide (8b). Yellow oil. Yield: 35%. 1H NMR (400 MHz, DMSO- d_6) δ (ppm): 9.64 (1H, s, NH, exchange with D₂O), 8.55 (1H, s), 8.50 (1H, s), 8.28 (1H, s), 8.09 (1H, s, NH, exchange with D₂O), 7.96 (1H, s), 7.88 (1H, d, $J = 7.34$ Hz), 7.76 (2H, d, $J = 9.15$ Hz), 7.66 (1H, m), 7.55 (1H, m), 7.47 (1H, d, $J = 7.92$ Hz), 7.41 (1H, d, $J = 7.92$ Hz), 7.32 (2H, bs, SO₂NH₂, exchange with D₂O), 7.24 (1H, s), 4.72 (2H, m), 4.59 (2H, m), 4.30 (4H, m), 4.13 (2H, m), 3.88 (2H, t, $J = 5.09$ Hz), 3.79 (2H, m), 3.75 (2H, m), 3.55 (3H, m), 3.52 (10H, m), 3.16 (3H, d, $J = 4.85$ Hz). ^{13}C NMR (100 MHz, DMSO- d_6) δ (ppm): 185.4, 156.3, 154.8, 146.2, 140.0, 136.0, 131.0, 129.4, 129.0, 127.0, 126.7, 121.8, 120.7, 118.7, 116.7, 103.3, 90.2, 70.1, 69.9, 69.7, 69.6, 69.1, 68.6, 68.0, 65.3, 61.9, 60.2, 58.4, 49.7, 49.6, 29.0; MS (ESI positive) m/z : 826.30 $[M + H]^+$. Elemental analysis: calculated C, 53.81; H, 5.74; N, 15.26; found C, 53.79; H, 5.75; N, 15.28.

4-(15-(4-(3-((6,7-Bis(2-methoxyethoxy)quinazolin-4-yl)amino)phenyl)-1H-1,2,3-triazol-1-yl)-3-thioxo-7,10,13-trioxa-2,4-diazapentadecyl)benzenesulfonamide (8c). Yellow oil. Yield: 42%. 1H NMR (400 MHz, DMSO- d_6) δ (ppm): 9.64 (1H, m, NH, exchange with D₂O), 8.59 (1H, s), 8.32 (1H, s), 8.00 (1H, s, NH, exchange with D₂O), 7.90 (1H, d, $J = 7.38$ Hz), 7.80 (2H, d, $J = 7.72$ Hz), 7.59 (1H, d, $J = 7.72$ Hz), 7.51 (1H, d, $J = 8.13$ Hz), 7.46 (2H, d, $J = 7.72$ Hz), 7.35 (2H, bs, SO₂NH₂, exchange with D₂O), 7.33 (1H, m), 4.62 (2H, m), 4.34 (5H, m), 3.92 (2H, m), 3.81 (5H, m), 3.54 (14H, m), 3.41 (8H, m). ^{13}C NMR (100 MHz, DMSO- d_6) δ (ppm): 163.2, 156.4, 155.8, 153.6, 152.8, 148.1, 146.8, 146.2, 142.5, 140.0, 137.6, 131.1, 129.0, 127.8, 127.4, 125.6, 121.8, 120.3, 118.8, 108.1, 103.3, 70.1, 70.0, 69.7, 69.6, 68.8, 68.6, 68.4, 68.0, 58.4, 58.4, 49.6, 48.6; MS (ESI positive) m/z : 840.31 $[M + H]^+$. Elemental analysis: calculated C, 53.98; H, 5.78; N, 15.18; found C, 53.99; H, 5.79; N, 15.16.

4-(16-(4-(3-((6,7-Bis(2-methoxyethoxy)quinazolin-4-yl)amino)phenyl)-1H-1,2,3-triazol-1-yl)-4-thioxo-8,11,14-trioxa-3,5-diazahexadecyl)benzenesulfonamide (8d). Yellow oil. Yield: 72%. 1H NMR (400 MHz, DMSO- d_6) δ (ppm): 9.69 (1H, m, NH, exchange with D₂O), 8.55 (2H, m), 8.28 (1H, s), 7.94 (1H, s, NH, exchange with D₂O), 7.84 (1H, m), 7.75 (3H, m), 7.57 (1H, d, $J = 7.51$ Hz), 7.48 (1H, m), 7.42 (3H, m), 7.32 (2H, bs, SO₂NH₂, exchange with D₂O), 7.32 (1H, s), 4.58 (2H, m), 4.30 (4H, m), 3.87 (2H, m), 3.79 (2H, m), 3.75 (2H, m), 3.49 (16H, m), 3.35 (4H, m), 2.88 (2H, m). ^{13}C NMR (100 MHz, DMSO- d_6) δ (ppm): 190.0, 158.9, 158.3, 153.7, 148.3, 148.2, 146.5, 146.2, 143.4, 142.2, 131.1, 129.7, 129.3, 129.2, 126.3, 126.0, 125.7, 122.5, 121.9, 103.2, 80.2, 70.1, 70.0, 69.7, 69.6, 69.3, 68.9, 68.7, 68.4, 68.1, 58.4, 58.3, 50.0, 49.6; MS (ESI positive) m/z : 854.33 $[M + H]^+$. Elemental

analysis: calculated C, 54.85; H, 6.02; N, 14.76; found C, 54.84; H, 6.01; N, 14.78.

4-(3-(2-(2-(2-(2-(4-(3-((6,7-Bis(2-methoxyethoxy)quinazolin-4-yl)amino)phenyl)-1H-1,2,3-triazol-1-yl)ethoxy)ethoxy)ethoxy)ethyl)thioureido)-3-bromo-2-methylbenzenesulfonamide (9). Yellow oil. Yield: 46%. 1H NMR (400 MHz, DMSO- d_6) δ (ppm): 9.57 (1H, s, NH, exchange with D₂O), 8.53 (1H, s), 8.46 (2H, m), 8.28 (1H, s), 8.11 (1H, m), 7.93 (1H, s, NH, exchange with D₂O), 7.88 (1H, d, $J = 8.26$ Hz), 7.65 (1H, d, $J = 8.68$ Hz), 7.54 (1H, d, $J = 7.44$ Hz), 7.46 (2H, m), 7.23 (1H, s), 7.20 (2H, bs, SO₂NH₂, exchange with D₂O), 4.58 (2H, m), 4.31 (4H, m), 3.87 (2H, t, $J = 5.37$ Hz), 3.79 (2H, t, $J = 4.22$ Hz), 3.75 (2H, t, $J = 4.21$ Hz), 3.54 (8H, m), 3.45–3.40 (9H, m). ^{13}C NMR (100 MHz, DMSO- d_6) δ (ppm): 182.2, 180.4, 175.7, 173.9, 172.0, 168.6, 152.9, 147.3, 144.8, 141.6, 140.1, 138.7, 134.5, 130.3, 121.8, 120.2, 119.0, 117.3, 116.2, 112.4, 107.8, 105.5, 103.3, 75.3, 70.0, 69.6, 68.5, 68.3, 68.0, 67.3, 63.4, 58.3, 55.3, 46.8, 31.9; MS (ESI positive) m/z : 918.22 $[M + H]^+$. Elemental analysis: calculated C, 52.17; H, 5.63; N, 14.22; found C, 52.18; H, 5.64; N, 14.21.

2.1.5. General procedure for the synthesis of compounds 25a–b and 26. The commercially available sulfonamide **23a**, **b** or coumarin **24** (ref. 70) (1 equiv.), commercially available 2,5-dioxopyrrolidin-1-yl 1-azido-3,6,9,12,15-pentaoxaoctadecan-18-oate **22** (1.1 equiv., azido-PEG5-NHS ester) and DIPEA (1.5 equiv.) were dissolved in DMF (5 mL) and the reaction mixture was stirred at room temperature overnight. Then, the reaction mixture was treated with slush and aq. NH₄Cl, extracted with EtOAc thrice. The combined organic layers were dried over anhydrous Na₂SO₄, filtered, and evaporated at reduced pressure. The crude material was purified by flash column chromatography (MeOH/DCM: 5:95), to yield compounds **25a–b** and **26**.

1-Azido-N-(4-sulfamoylbenzyl)-3,6,9,12,15-pentaoxaoctadecan-18-amide (25a). Yellow oil. Yield: 61%. 1H NMR (400 MHz, DMSO- d_6) δ (ppm): 8.45 (1H, t, $J = 5.32$ Hz, NH, exchange with D₂O), 7.75 (2H, d, $J = 8.20$ Hz), 7.41 (2H, d, $J = 7.91$ Hz), 7.30 (2H, bs, SO₂NH₂, exchange with D₂O), 4.33 (2H, d, $J = 5.73$ Hz), 3.63 (2H, t, $J = 7.09$ Hz), 3.60 (2H, t, $J = 4.84$ Hz), 3.50 (16H, m), 3.38 (2H, t, $J = 4.91$ Hz), 2.39 (2H, t, $J = 6.09$ Hz). ^{13}C NMR (100 MHz, DMSO- d_6) δ (ppm): 174.3, 142.3, 138.2, 130.3, 129.2, 70.6, 70.4, 69.5, 68.4, 49.1, 45.8, 25.9; MS (ESI positive) m/z : 504.2 $[M + H]^+$. Elemental analysis: calculated C, 47.70; H, 6.61; N, 13.91; found C, 47.71; H, 6.62; N, 13.90.

1-Azido-N-(4-sulfamoylphenethyl)-3,6,9,12,15-pentaoxaoctadecan-18-amide (25b). Yellow oil. Yield: 49%. 1H NMR (400 MHz, DMSO- d_6) δ (ppm): 7.93 (1H, t, $J = 5.70$ Hz, NH, exchange with D₂O), 7.73 (2H, d, $J = 8.15$ Hz), 7.38 (2H, d, $J = 8.15$ Hz), 7.28 (2H, bs, SO₂NH₂, exchange with D₂O), 3.58 (4H, m), 3.53 (4H, m), 3.49 (8H, m), 3.47 (4H, m), 3.38 (2H, t, $J = 5.14$ Hz), 3.27 (2H, t, $J = 6.14$ Hz), 2.77 (2H, t, $J = 7.14$ Hz), 2.28 (2H, t, $J = 6.42$ Hz). ^{13}C NMR (100 MHz, DMSO- d_6) δ (ppm): 179.3, 142.5, 133.8, 130.9, 128.5, 71.4, 70.6, 70.4, 69.1, 67.3, 45.4, 42.7, 37.4, 30.4; MS (ESI positive) m/z : 517.22 $[M + H]^+$.



Elemental analysis: calculated C, 48.73; H, 6.82; N, 13.53; found C, 48.72; H, 6.83; N, 13.54.

1-Azido-N-(2-((2-oxo-2H-chromen-6-yl)oxy)ethyl)-3,6,9,12,15-pentaoxaoctadecan-18-amide (**26**). Yellow oil. Yield: 70%. ¹H NMR (400 MHz, DMSO-*d*₆) δ (ppm): 8.12 (1H, t, *J* = 5.82 Hz, NH, exchange with D₂O), 8.00 (1H, d, *J* = 9.41 Hz), 7.34 (1H, d, *J* = 9.09 Hz), 7.30 (1H, d, *J* = 2.76 Hz), 7.20 (1H, dd, *J* = 8.08, 9.02 Hz), 6.49 (1H, d, *J* = 9.61 Hz), 4.02 (2H, t, *J* = 5.64 Hz), 3.58 (4H, m), 3.52 (4H, m), 3.49 (4H, m), 3.45 (10H, m), 3.38 (2H, t, *J* = 4.73 Hz), 2.34 (2H, t, *J* = 6.20 Hz). ¹³C NMR (100 MHz, DMSO-*d*₆) δ (ppm): 171.9, 165.2, 158.1, 154.7, 142.0, 119.3, 115.2, 113.4, 111.5, 71.3, 70.8, 70.6, 70.3, 69.8, 65.5, 61.4, 48.1, 39.3, 29.2; MS (ESI positive) *m/z*: 523.2 [M + H]⁺. Elemental analysis: calculated C, 55.16; H, 6.56; N, 10.72; found C, 55.17; H, 6.55; N, 10.71.

2.1.6. General procedure for the synthesis of compounds 29a–c. The appropriate sulfonamide **16a–c** (ref. 66) (1 equiv.) and 17-azido-3,6,9,12,15-pentaoxaheptadecan-1-amine **27** (1.1 equiv.) were dissolved in ACN (10 mL) and the reaction mixture was stirred at reflux temperature overnight. Then, the reaction mixture was treated with slush and extracted with EtOAc thrice. The combined organic layers were dried over anhydrous Na₂SO₄, filtered, and evaporated at reduced pressure. The crude material was purified by flash column chromatography (MeOH/DCM: 5:95), to yield compounds **29a–c**.

3-(3-(17-Azido-3,6,9,12,15-pentaoxaheptadecyl)ureido)benzenesulfonamide (**29a**). Yellow oil. Yield: 67%. ¹H NMR (400 MHz, DMSO-*d*₆) δ (ppm): 8.88 (1H, s, NH, exchange with D₂O), 7.96 (1H, s, NH, exchange with D₂O), 7.50 (1H, d, *J* = 7.51 Hz), 7.39 (1H, t, *J* = 7.93 Hz), 7.33 (1H, d, *J* = 7.51 Hz), 7.28 (2H, bs, SO₂NH₂, exchange with D₂O), 6.24 (1H, t, *J* = 5.18 Hz), 3.59 (2H, t, *J* = 4.47 Hz), 3.52 (16H, m), 3.46 (2H, t, *J* = 5.35 Hz), 3.38 (2H, t, *J* = 4.07 Hz), 3.26 (2H, m). ¹³C NMR (100 MHz, DMSO-*d*₆) δ (ppm): 155.6, 145.1, 141.4, 130.0, 121.2, 118.9, 115.1, 70.4, 70.3, 70.2, 69.9, 50.7, 31.3; MS (ESI positive) *m/z*: 505.1 [M + H]⁺. Elemental analysis: calculated C, 45.23; H, 6.39; N, 16.66; found C, 45.22; H, 6.38; N, 16.67.

4-(3-(17-Azido-3,6,9,12,15-pentaoxaheptadecyl)ureido)benzenesulfonamide (**29b**). Yellow oil. Yield: 32%. ¹H NMR (400 MHz, DMSO-*d*₆) δ (ppm): 8.95 (1H, s, NH, exchange with D₂O), 7.66 (2H, d, *J* = 8.27 Hz), 7.52 (2H, d, *J* = 9.05 Hz), 7.14 (2H, bs, SO₂NH₂, exchange with D₂O), 6.32 (1H, t, *J* = 5.51 Hz, NH, exchange with D₂O), 3.58 (2H, t, *J* = 5.11 Hz), 3.52 (16H, m), 3.46 (2H, t, *J* = 5.38 Hz), 3.38 (2H, t, *J* = 4.69 Hz), 3.25 (2H, dt, *J* = 10.71, 11.34 Hz). ¹³C NMR (100 MHz, DMSO-*d*₆) δ (ppm): 159.8, 143.7, 137.4, 130.6, 128.1, 70.6, 70.4, 70.5, 69.4, 50.2, 46.3; MS (ESI positive) *m/z*: 527.2 [M + H]⁺. Elemental analysis: calculated C, 45.23; H, 6.39; N, 16.66; found C, 45.21; H, 6.40; N, 16.68.

4-(21-Azido-3-oxo-7,10,13,16,19-pentaoxa-2,4-diazahenicosyl)benzenesulfonamide (**29c**). Yellow oil. Yield: 52%. ¹H NMR (400 MHz, DMSO-*d*₆) δ (ppm): 7.75 (2H, d, *J* = 8.27 Hz), 7.39 (2H, d, *J* = 8.27 Hz), 7.26 (2H, bs, SO₂NH₂, exchange with D₂O), 6.51 (1H, t, *J* = 6.39 Hz, NH, exchange with D₂O), 6.02 (1H, t, *J* = 4.88 Hz, NH, exchange with D₂O), 4.25 (2H, d, *J* =

5.26 Hz), 3.59 (2H, t, *J* = 5.13 Hz), 3.52 (16H, m), 3.38 (4H, m), 3.17 (2H, t, *J* = 5.47 Hz). ¹³C NMR (100 MHz, DMSO-*d*₆) δ (ppm): 163.7, 142.3, 137.4, 129.4, 129.1, 128.5, 71.7, 70.8, 70.6, 70.4, 67.3, 65.9, 50.2, 45.6, 44.9; MS (ESI positive) *m/z*: 519.2 [M + H]⁺. Elemental analysis: calculated C, 45.60; H, 6.47; N, 16.50; found C, 45.61; H, 6.48; N, 16.49.

2.1.7. General procedure for the synthesis of compounds 32b–d. The appropriate sulfonamide **17b–d** (1 equiv.) and the commercial 17-azido-3,6,9,12,15-pentaoxaheptadecan-1-amine **27** (1.1 equiv.) were dissolved in ACN (5 mL). K₂CO₃ (1.5 equiv.) was then added, and the reaction mixture was stirred at room temperature overnight. Then, the reaction mixture was treated with slush and aq. NH₄Cl, extracted with EtOAc thrice. The combined organic layers were dried over anhydrous Na₂SO₄, filtered, and evaporated at reduced pressure. The crude material was purified by flash column chromatography (MeOH/DCM: 5:95), to yield compounds **18b–d**.

4-(3-(17-Azido-3,6,9,12,15-pentaoxaheptadecyl)thioureido)benzenesulfonamide (**32b**). Yellow oil. Yield: 67%. ¹H NMR (400 MHz, DMSO-*d*₆) δ (ppm): 9.92 (1H, s, NH, exchange with D₂O), 8.02 (1H, s, NH, exchange with D₂O), 7.72 (2H, d, *J* = 8.69 Hz), 7.67 (2H, d, *J* = 8.99 Hz), 7.26 (2H, bs, SO₂NH₂, exchange with D₂O), 3.64 (2H, m), 3.59 (5H, m), 3.54 (15H, m), 3.38 (2H, t, *J* = 4.71 Hz). ¹³C NMR (100 MHz, DMSO-*d*₆) δ (ppm): 179.2, 140.1, 138.7, 129.1, 119.6, 119.5, 71.1, 70.8, 70.6, 69.8, 65.8, 50.2, 44.7; MS (ESI positive) *m/z*: 521.2 [M + H]⁺. Elemental analysis: calculated C, 43.83; H, 6.20; N, 16.14; found C, 43.82; H, 6.21; N, 16.12.

4-(21-Azido-3-thioxo-7,10,13,16,19-pentaoxa-2,4-diazahenicosyl)benzenesulfonamide (**32c**). Yellow oil. Yield: 63%. ¹H NMR (400 MHz, DMSO-*d*₆) δ (ppm): 8.03 (1H, s, NH, exchange with D₂O), 7.75 (2H, d, *J* = 8.39 Hz), 7.63 (1H, s, NH, exchange with D₂O), 7.42 (2H, d, *J* = 8.44 Hz), 7.31 (2H, bs, SO₂NH₂, exchange with D₂O), 4.73 (2H, m), 3.54 (22H, m), 3.38 (2H, t, *J* = 5.26 Hz). ¹³C NMR (100 MHz, DMSO-*d*₆) δ (ppm): 168.9, 146.7, 136.2, 127.4, 124.7, 70.9, 70.7, 70.6, 69.3, 63.5, 47.5, 46.3, 44.4; MS (ESI positive) *m/z*: 535.1 [M + H]⁺. Elemental analysis: calculated C, 44.39; H, 6.30; N, 15.93; found C, 44.38; H, 6.31; N, 15.94.

4-(22-Azido-4-thioxo-8,11,14,17,20-pentaoxa-3,5-diazadocosyl)benzenesulfonamide (**32d**). Yellow oil. Yield: 60%. ¹H NMR (400 MHz, DMSO-*d*₆) δ (ppm): 7.74 (2H, d, *J* = 8.20 Hz), 7.53 (1H, s, NH, exchange with D₂O), 7.47 (1H, s, NH, exchange with D₂O), 7.41 (2H, d, *J* = 8.11 Hz), 7.29 (2H, bs, SO₂NH₂, exchange with D₂O), 3.59 (3H, t, *J* = 4.22 Hz), 3.51 (21H, m), 3.38 (2H, t, *J* = 4.87 Hz), 2.87 (2H, t, *J* = 7.06 Hz). ¹³C NMR (100 MHz, DMSO-*d*₆) δ (ppm): 144.4, 142.5, 129.8, 128.7, 126.4, 70.4, 70.3, 69.9, 69.6, 50.7, 35.2; MS (ESI positive) *m/z*: 549.3 [M + H]⁺. Elemental analysis: calculated C, 45.97; H, 6.61; N, 15.32; found C, 45.98; H, 6.62; N, 15.33.

2.1.8. General procedure for the synthesis of compounds 10a–b, 11, 12a–c, 13, and 14b–d. Erlotinib hydrochloride (ERL) (1 equiv.), sodium ascorbate (1.6 equiv.), CuSO₄·5H₂O (0.8 equiv.) tetramethylammonium chloride (4.5 equiv.) and the appropriate azide derivative **25a–b, 26, 28, 29a–c**, or



32b-d (1.1 equiv.) were dissolved in H₂O/*tert*-butanol (1:1) and stirred overnight at 60 °C. Then, the reaction mixture was filtered through a cake of Celite 521© and the filtrate was treated with slush. The crude was extracted with EtOAc thrice. The combined organic layers were dried over anhydrous Na₂SO₄, filtered, and evaporated at reduced pressure. The crude material was purified by flash column chromatography (MeOH/DCM: 5:95), to yield compounds **10a-b**, **11**, **12a-c**, **13**, and **14b-d**.

1-(4-(3-((6,7-Bis(2-methoxyethoxy)quinazolin-4-yl)amino)phenyl)-1H-1,2,3-triazol-1-yl)-N-(4-sulfamoylbenzyl)-3,6,9,12,15-pentaoxaoctadecan-18-amide (10a). Yellow powder. Yield: 47%. ¹H NMR (400 MHz, DMSO-*d*₆) δ (ppm): 9.55 (1H, s, NH, exchange with D₂O), 8.53 (1H, s), 8.48 (1H, s), 8.42 (1H, t, *J* = 5.90 Hz), 8.27 (1H, s), 7.93 (1H, s), 7.89 (1H, d, *J* = 9.02 Hz), 7.75 (2H, d, *J* = 7.98 Hz), 7.54 (1H, d, *J* = 7.63 Hz), 7.47 (1H, d, *J* = 8.06 Hz), 7.40 (2H, d, *J* = 8.29 Hz), 7.29 (2H, bs, SO₂NH₂, exchange with D₂O), 7.23 (1H, s), 4.58 (2H, t, *J* = 5.12 Hz), 4.30 (6H, m), 3.88 (2H, t, *J* = 5.12 Hz), 3.79 (2H, t, *J* = 4.44 Hz), 3.75 (2H, t, *J* = 4.44 Hz), 3.61 (2H, t, *J* = 6.15 Hz), 3.56 (2H, m), 3.50 (2H, m), 3.45 (14H, m), 3.38 (3H, s), 3.36 (3H, s), 2.38 (2H, t, *J* = 6.01 Hz). ¹³C NMR (100 MHz, DMSO-*d*₆) δ (ppm): 170.8, 156.6, 153.9, 153.1, 148.4, 147.0, 146.4, 143.8, 142.5, 140.0, 131.2, 129.3, 127.5, 125.8, 122.0, 121.9, 120.6, 119.0, 109.1, 108.3, 103.5, 70.3, 70.2, 69.9, 69.8, 69.7, 68.8, 68.6, 68.3, 67.0, 58.6, 58.5, 49.8, 41.9, 41.8, 36.3; MS (ESI positive) *m/z*: 897.4 [M + H]⁺. Elemental analysis: calculated C, 52.34; H, 6.41; N, 13.56; found C, 52.33; H, 6.42; N, 13.55.

1-(4-(3-((6,7-Bis(2-methoxyethoxy)quinazolin-4-yl)amino)phenyl)-1H-1,2,3-triazol-1-yl)-N-(4-sulfamoylphenethyl)-3,6,9,12,15-pentaoxonadecan-19-amide (10b). Green powder. Yield: 44%. ¹H NMR (400 MHz, DMSO-*d*₆) δ (ppm): 9.58 (1H, s, NH, exchange with D₂O), 8.56 (1H, s), 8.51 (1H, s), 8.45 (1H, t, *J* = 5.90 Hz), 8.30 (1H, s), 7.96 (1H, s), 7.92 (1H, d, *J* = 9.02 Hz), 7.78 (2H, d, *J* = 7.98 Hz), 7.57 (1H, d, *J* = 7.63 Hz), 7.50 (1H, d, *J* = 8.06 Hz), 7.43 (2H, d, *J* = 8.29 Hz), 7.32 (2H, bs, SO₂NH₂, exchange with D₂O), 7.26 (1H, s), 4.61 (2H, t, *J* = 5.12 Hz), 4.33 (6H, m), 3.91 (2H, t, *J* = 5.12 Hz), 3.82 (2H, t, *J* = 4.44 Hz), 3.78 (2H, t, *J* = 4.44 Hz), 3.64 (2H, t, *J* = 6.15 Hz), 3.59 (2H, m), 3.53 (2H, m), 3.48 (16H, m), 3.41 (3H, s), 3.39 (3H, s), 2.41 (2H, t, *J* = 6.01 Hz). ¹³C NMR (100 MHz, DMSO-*d*₆) δ (ppm): 175.9, 172.1, 157.4, 154.6, 153.7, 149.0, 147.0, 144.7, 142.0, 140.3, 131.5, 130.1, 130.0, 126.4, 123.0, 122.8, 121.6, 120.0, 109.6, 108.3, 103.8, 70.8, 70.7, 70.3, 70.2, 70.1, 69.2, 69.2, 68.9, 67.3, 62.7, 59.1, 56.1, 50.5, 49.3, 40.4, 36.7, 35.2; MS (ESI positive) *m/z*: 911.4 [M + H]⁺. Elemental analysis: calculated C, 54.21; H, 6.46; N, 13.00; found C, 54.20; H, 6.47; N, 13.01.

1-(4-(3-((6,7-Bis(2-methoxyethoxy)quinazolin-4-yl)amino)phenyl)-1H-1,2,3-triazol-1-yl)-N-(2-((2-oxo-2H-chromen-6-yl)oxy)ethyl)-3,6,9,12,15-pentaoxaoctadecan-18-amide (11). Green oil. Yield: 45%. ¹H NMR (400 MHz, DMSO-*d*₆) δ (ppm): 9.68 (1H, s, NH, exchange with D₂O), 8.54 (1H, s), 8.51 (1H, s), 8.26 (1H, s), 8.11 (1H, t, *J* = 4.93 Hz), 7.98 (1H, d, *J* = 9.42 Hz), 7.94 (1H, s), 7.87 (1H, d, *J* = 8.35 Hz), 7.56 (1H,

d, *J* = 7.59 Hz), 7.46 (1H, t, *J* = 7.84 Hz), 7.32 (1H, d, *J* = 8.85 Hz), 7.27 (1H, d, *J* = 2.85 Hz), 7.23 (1H, s), 7.19 (1H, dd, *J* = 8.87 Hz, 9.27 Hz), 6.48 (1H, d, *J* = 9.50 Hz), 4.58 (2H, t, *J* = 5.51 Hz), 4.30 (4H, m), 4.00 (2H, t, *J* = 5.55 Hz), 3.88 (2H, t, *J* = 5.10 Hz), 3.79 (2H, t, *J* = 4.80 Hz), 3.75 (2H, t, *J* = 4.35 Hz), 3.57 (4H, m), 3.49 (2H, m), 3.42 (12H, m), 3.37 (8H, m), 2.32 (2H, t, *J* = 6.37 Hz). ¹³C NMR (100 MHz, DMSO-*d*₆) δ (ppm): 179.7, 170.4, 160.0, 156.5, 154.7, 153.8, 152.5, 148.2, 147.9, 146.1, 144.0, 139.7, 131.1, 129.0, 121.9, 121.8, 120.5, 119.8, 119.1, 118.9, 117.3, 116.5, 111.5, 108.9, 107.5, 103.4, 70.1, 70.0, 69.7, 69.6, 69.4, 68.6, 68.4, 68.1, 67.0, 66.7, 58.4, 58.3, 49.6, 47.2, 38.1, 36.0, 30.6; MS (ESI positive) *m/z*: 916.9 [M + H]⁺. Elemental analysis: calculated C, 55.91; H, 6.41; N, 12.36; found C, 55.92; H, 6.42; N, 12.34.

3-(3-(17-(4-(3-((6,7-Bis(2-methoxyethoxy)quinazolin-4-yl)amino)phenyl)-1H-1,2,3-triazol-1-yl)-3,6,9,12,15-pentaoxaheptadecyl)ureido)benzenesulfonamide (12a). Yellow powder. Yield: 40%. ¹H NMR (400 MHz, DMSO-*d*₆) δ (ppm): 9.56 (1H, s, NH, exchange with D₂O), 8.87 (1H, s, NH, exchange with D₂O), 8.53 (1H, s), 8.48 (1H, s), 8.27 (1H, s), 7.96 (1H, m), 7.93 (1H, s), 7.89 (1H, d, *J* = 8.71 Hz), 7.54 (1H, d, *J* = 7.87 Hz), 7.48 (2H, m), 7.34 (2H, m), 7.28 (2H, bs, SO₂NH₂, exchange with D₂O), 7.22 (1H, s), 6.23 (1H, t, *J* = 5.58 Hz), 4.58 (2H, t, *J* = 5.27 Hz), 4.30 (4H, m), 3.88 (2H, t, *J* = 4.96 Hz), 3.79 (2H, t, *J* = 4.34 Hz), 3.75 (2H, t, *J* = 4.23 Hz), 3.55 (2H, m), 3.46 (16H, m), 3.38 (3H, s), 3.36 (3H, s), 3.24 (2H, m). ¹³C NMR (100 MHz, DMSO-*d*₆) δ (ppm): 156.4, 154.9, 153.6, 152.9, 148.1, 146.9, 146.2, 144.5, 140.9, 140.0, 131.0, 129.2, 128.9, 121.8, 121.7, 120.3, 120.2, 118.7, 118.0, 114.4, 109.0, 108.2, 103.3, 76.4, 72.9, 70.6, 70.1, 70.0, 69.7, 69.6, 68.6, 68.4, 68.0, 59.5, 58.4, 58.3, 56.6, 49.6, 47.2, 30.6; MS (ESI positive) *m/z*: 898.4 [M + H]⁺. Elemental analysis: calculated C, 54.84; H, 6.17; N, 14.04; found C, 54.86; H, 6.15; N, 14.03.

4-(3-(17-(4-(3-((6,7-Bis(2-methoxyethoxy)quinazolin-4-yl)amino)phenyl)-1H-1,2,3-triazol-1-yl)-3,6,9,12,15-pentaoxaheptadecyl)ureido)benzenesulfonamide (12b). Reddish oil. Yield: 25%. ¹H NMR (400 MHz, DMSO-*d*₆) δ (ppm): 9.59 (1H, s, NH, exchange with D₂O), 8.56 (1H, s, NH, exchange with D₂O), 8.50 (1H, s), 8.29 (1H, s), 7.95 (1H, s), 7.91 (1H, d, *J* = 8.23 Hz), 7.77 (2H, d, *J* = 7.84 Hz), 7.56 (1H, d, *J* = 8.07 Hz), 7.48 (1H, t, *J* = 7.81 Hz), 7.41 (2H, d, *J* = 8.22 Hz), 7.30 (2H, bs, SO₂NH₂, exchange with D₂O), 7.25 (1H, s), 6.54 (1H, t, *J* = 4.86 Hz), 6.05 (1H, t, *J* = 5.43 Hz), 4.61 (2H, t, *J* = 4.94 Hz), 4.32 (4H, m), 4.27 (2H, d, *J* = 5.87 Hz), 3.90 (2H, t, *J* = 5.10 Hz), 3.81 (2H, t, *J* = 4.34 Hz), 3.77 (2H, t, *J* = 3.06 Hz), 3.57 (2H, m), 3.47 (14H, m), 3.39 (8H, m). ¹³C NMR (100 MHz, DMSO-*d*₆) δ (ppm): 158.3, 155.7, 154.9, 149.5, 148.6, 145.6, 143.7, 137.4, 135.9, 135.2, 131.4, 129.3, 126.7, 124.2, 123.1, 122.0, 121.3, 116.6, 107.2, 105.0, 100.6, 72.6, 69.9, 69.7, 69.6, 69.0, 68.8, 68.6, 58.4, 57.3, 51.3, 49.7, 44.0, 36.3, 35.4, 29.0; MS (ESI positive) *m/z*: 898.4 [M + H]⁺. Elemental analysis: calculated C, 54.84; H, 6.17; N, 14.04; found C, 54.86; H, 6.16; N, 14.02.



4-(21-(4-(3-((6,7-Bis(2-methoxyethoxy)quinazoline-4-yl)amino)phenyl)-1H-1,2,3-triazol-1-yl)-3-oxo-7,10,13,16,19-pentaoxa-2,4-diazahenicosyl)benzenesulfonamide (**12c**). Yellow powder. Yield: 52%. ¹H NMR (400 MHz, DMSO-*d*₆) δ (ppm): 9.57 (1H, s, NH, exchange with D₂O), 8.54 (1H, s, NH, exchange with D₂O), 8.48 (1H, s), 8.27 (1H, s), 7.93 (1H, s), 7.89 (1H, d, *J* = 8.23 Hz), 7.75 (2H, d, *J* = 7.84 Hz), 7.54 (1H, d, *J* = 8.07 Hz), 7.46 (1H, t, *J* = 7.81 Hz), 7.39 (2H, d, *J* = 8.22 Hz), 7.28 (2H, bs, SO₂NH₂, exchange with D₂O), 7.23 (1H, s), 6.52 (1H, t, *J* = 4.86 Hz), 6.03 (1H, t, *J* = 5.43 Hz), 4.59 (2H, t, *J* = 4.94 Hz), 4.30 (4H, m), 4.25 (2H, d, *J* = 5.87 Hz), 3.88 (2H, t, *J* = 5.10 Hz), 3.79 (2H, t, *J* = 4.34 Hz), 3.75 (2H, t, *J* = 3.06 Hz), 3.55 (2H, m), 3.45 (14H, m), 3.37 (8H, m), 3.15 (2H, m). ¹³C NMR (100 MHz, DMSO-*d*₆) δ (ppm): 162.6, 160.9, 158.3, 154.9, 152.8, 149.5, 146.9, 144.8, 140.1, 138.2, 135.2, 133.1, 129.3, 126.7, 124.2, 123.2, 122.1, 121.3, 116.6, 112.1, 108.7, 102.3, 98.2, 69.9, 69.7, 69.6, 69.0, 68.8, 68.6, 58.4, 57.3, 49.7, 44.0, 29.0; MS (ESI positive) *m/z*: 912.6 [M + H]⁺. Elemental analysis: calculated C, 55.31; H, 6.30; N, 13.82; found C, 55.30; H, 6.31; N, 13.83.

1-(17-(4-(3-((6,7-Bis(2-methoxyethoxy)quinazolin-4-yl)amino)phenyl)-1H-1,2,3-triazol-1-yl)-3,6,9,12,15-pentaoxa heptadecyl)-3-(2-oxo-2H-chromen-6-yl)urea (**13**). White powder. Yield: 54%. ¹H NMR (400 MHz, DMSO-*d*₆) δ (ppm): 9.56 (1H, s, NH, exchange with D₂O), 8.81 (1H, s, NH, exchange with D₂O), 8.53 (1H, s), 8.48 (1H, s), 8.27 (1H, s), 8.01 (1H, d, *J* = 9.6 Hz), 7.93 (1H, s), 7.89 (1H, d, *J* = 7.8 Hz), 7.79 (1H, d, *J* = 2.5 Hz), 7.54 (1H, d, *J* = 7.6 Hz), 7.49 (1H, dd, *J* = 9.2 Hz, *J* = 2.7 Hz), 7.45 (1H, t, *J* = 7.9 Hz), 7.27 (1H, d, *J* = 8.9 Hz), 7.22 (1H, s), 6.43 (1H, d, *J* = 9.5 Hz), 6.28 (1H, t, *J* = 5.4 Hz), 4.58 (2H, t, *J* = 4.9 Hz), 4.30 (4H, m), 3.88 (2H, t, *J* = 5.0 Hz), 3.79 (2H, t, *J* = 4.7 Hz), 3.75 (2H, t, *J* = 4.5 Hz), 3.55 (2H, m), 3.46 (16H, m), 3.37 (3H, s), 3.36 (3H, s), 3.23 (2H, m). ¹³C NMR (100 MHz, DMSO-*d*₆) δ (ppm): 179.7, 170.4, 160.0, 156.5, 154.7, 153.8, 152.5, 148.2, 147.9, 146.1, 144.0, 139.7, 131.1, 129.0, 121.9, 121.8, 120.5, 119.8, 119.1, 118.9, 117.3, 116.5, 111.5, 108.9, 107.5, 103.4, 70.1, 70.0, 69.7, 69.6, 69.4, 68.6, 68.4, 68.1, 67.0, 66.7, 58.4, 58.3, 49.6, 38.1, 36.0; MS (ESI positive) *m/z*: 887.6 [M + H]⁺. Elemental analysis: calculated C, 59.58; H, 6.14; N, 12.63; found C, 59.56; H, 6.15; N, 12.64.

4-(3-(17-(4-(3-((6,7-Bis(2-methoxyethoxy)quinazolin-4-yl)amino)phenyl)-1H-1,2,3-triazol-1-yl)-3,6,9,12,15-pentaoxaheptadecyl)thioureido)benzenesulfonamide (**14b**). Yellow powder. Yield: 36%. ¹H NMR (400 MHz, DMSO-*d*₆) δ (ppm): 9.95 (1H, s, NH, exchange with D₂O), 8.55 (1H, s, NH, exchange with D₂O), 8.49 (1H, s), 8.28 (1H, s), 7.94 (1H, s), 7.90 (1H, d, *J* = 8.23 Hz), 7.76 (2H, d, *J* = 7.84 Hz), 7.55 (1H, d, *J* = 8.07 Hz), 7.47 (1H, t, *J* = 7.81 Hz), 7.40 (2H, d, *J* = 8.22 Hz), 7.29 (2H, bs, SO₂NH₂, exchange with D₂O), 7.24 (1H, s), 6.53 (1H, t, *J* = 4.86 Hz), 6.04 (1H, t, *J* = 5.43 Hz), 4.60 (2H, t, *J* = 4.94 Hz), 4.31 (4H, m), 4.26 (2H, d, *J* = 5.87 Hz), 3.89 (2H, t, *J* = 5.10 Hz), 3.80 (2H, t, *J* = 4.34 Hz), 3.76 (2H, t, *J* = 3.06 Hz), 3.56 (2H, m), 3.47 (14H, m), 3.39 (8H, m). ¹³C NMR (100 MHz, DMSO-*d*₆) δ (ppm): 162.9, 148.8, 147.3, 146.5, 144.3, 141.9, 140.0, 138.9, 131.3, 129.5, 129.4, 128.4, 126.0, 125.9, 125.7, 122.1, 120.8, 119.2, 117.6, 112.1, 104.5, 70.3, 70.2, 69.9,

68.8, 68.7, 68.5, 60.3, 58.6, 56.5, 55.4, 49.9, 48.7, 36.0, 29.1, 28.8, 22.2; MS (ESI positive) *m/z*: 914.4 [M + H]⁺. Elemental analysis: calculated C, 53.87; H, 6.07; N, 13.79; found C, 53.85; H, 6.06; N, 13.78.

4-(21-(4-(3-((6,7-Bis(2-methoxyethoxy)quinazoline-4-yl)amino)phenyl)-1H-1,2,3-triazol-1-yl)-3-thioxo-7,10,13,16,19-pentaoxa-2,4-diazahenicosyl)benzenesulfonamide (**14c**). Yellow powder. Yield: 41%. ¹H NMR (400 MHz, DMSO-*d*₆) δ (ppm): 10.15 (1H, s, NH, exchange with D₂O), 8.59 (1H, s, NH, exchange with D₂O), 8.53 (1H, s), 8.32 (1H, s), 7.98 (1H, s), 7.94 (1H, d, *J* = 8.23 Hz), 7.80 (2H, d, *J* = 7.84 Hz), 7.59 (1H, d, *J* = 8.07 Hz), 7.51 (1H, t, *J* = 7.81 Hz), 7.44 (2H, d, *J* = 8.22 Hz), 7.33 (2H, bs, SO₂NH₂, exchange with D₂O), 7.28 (1H, s), 6.57 (1H, t, *J* = 4.86 Hz), 6.08 (1H, t, *J* = 5.43 Hz), 4.64 (2H, t, *J* = 4.94 Hz), 4.35 (4H, m), 4.30 (2H, d, *J* = 5.87 Hz), 3.93 (2H, t, *J* = 5.10 Hz), 3.84 (2H, t, *J* = 4.34 Hz), 3.80 (2H, t, *J* = 3.06 Hz), 3.60 (2H, m), 3.50 (14H, m), 3.42 (8H, m), 3.20 (2H, m). ¹³C NMR (100 MHz, DMSO-*d*₆) δ (ppm): 155.4, 148.8, 146.5, 145.1, 144.3, 141.8, 140.0, 133.0, 131.3, 129.6, 129.3, 128.5, 126.0, 125.9, 125.7, 122.1, 120.8, 119.1, 115.8, 112.0, 104.5, 70.4, 70.3, 69.9, 68.8, 68.7, 68.4, 66.4, 61.9, 60.3, 58.6, 49.9, 48.7, 36.0, 29.1, 28.8, 22.3; MS (ESI positive) *m/z*: 928.4 [M + H]⁺. Elemental analysis: calculated C, 54.35; H, 6.19; N, 13.58; found C, 54.32; H, 6.21; N, 13.56.

4-(22-(4-(3-((6,7-Bis(2-methoxyethoxy)quinazolin-4-yl)amino)phenyl)-1H-1,2,3-triazol-1-yl)-4-thioxo-8,11,14,17,20-pentaoxa-3,5-diazadocosyl)benzenesulfonamide (**14d**). Yellow powder. Yield: 42%. ¹H NMR (400 MHz, DMSO-*d*₆) δ (ppm): 9.84 (1H, s, NH, exchange with D₂O), 8.52 (1H, s, NH, exchange with D₂O), 8.46 (1H, s), 8.25 (1H, s), 7.91 (1H, s), 7.87 (1H, d, *J* = 8.23 Hz), 7.73 (2H, d, *J* = 7.84 Hz), 7.52 (1H, d, *J* = 8.07 Hz), 7.44 (1H, t, *J* = 7.81 Hz), 7.37 (2H, d, *J* = 8.22 Hz), 7.26 (2H, bs, SO₂NH₂, exchange with D₂O), 7.21 (1H, s), 6.50 (1H, t, *J* = 4.86 Hz), 6.01 (1H, t, *J* = 5.43 Hz), 4.57 (2H, t, *J* = 4.94 Hz), 4.28 (4H, m), 4.23 (2H, d, *J* = 5.87 Hz), 3.86 (2H, t, *J* = 5.10 Hz), 3.77 (2H, t, *J* = 4.34 Hz), 3.73 (2H, t, *J* = 3.06 Hz), 3.54 (2H, m), 3.45 (14H, m), 3.35 (8H, m), 3.13 (2H, m), 2.97 (2H, m). ¹³C NMR (100 MHz, DMSO-*d*₆) δ (ppm): 153.3, 148.8, 145.2, 144.3, 141.9, 140.0, 138.3, 131.3, 129.7, 129.5, 129.4, 128.4, 125.9, 125.7, 122.1, 120.8, 119.2, 119.2, 112.1, 111.0, 104.3, 72.8, 70.3, 70.2, 70.1, 69.9, 68.8, 68.7, 68.5, 60.3, 58.6, 52.0, 49.9, 48.7, 36.0, 29.1, 28.8, 22.2; MS (ESI positive) *m/z*: 942.2 [M + H]⁺. Elemental analysis: calculated C, 54.82; H, 6.31; N, 13.38; found C, 54.81; H, 6.33; N, 13.36.

2.1.9. Procedure for the synthesis of compound 1-(17-azido-3,6,9,12,15-pentaoxaheptadecyl)-3-(2-oxo-2H-chromen-6-yl)urea (25). The appropriate coumarin **28** (ref. 71) (1 equiv.) and the commercial 17-azido-3,6,9,12,15-pentaoxaheptadecan-1-amine **27** (1.1 equiv.) were dissolved in ACN (10 mL) and the reaction mixture was stirred at reflux temperature overnight. Then, the reaction mixture was treated with slush and extracted with EtOAc thrice. The combined organic layers were dried over anhydrous Na₂SO₄, filtered, and evaporated at reduced pressure. The crude material was purified by flash column chromatography (MeOH/DCM: 2:98), to yield compound **25**. Brownish oil, Yield: 47%. ¹H NMR (400 MHz, DMSO-*d*₆) δ



(ppm): 8.81 (1H, s, NH, exchange with D₂O), 8.07 (1H, d, *J* = 9.9 Hz), 7.85 (1H, s, NH, exchange with D₂O), 7.54 (1H, d, *J* = 8.7 Hz), 7.33 (1H, d, *J* = 9.0 Hz), 6.49 (1H, d, *J* = 8.9 Hz), 6.29 (1H, t, *J* = 4.9 Hz), 3.57 (2H, m), 3.30 (3H, m). MS (ESI positive) *m/z*: 494.2 [M + H]⁺. Elemental analysis: calculated C, 53.54; H, 6.33; N, 14.19; found C, 53.52; H, 6.32; N, 14.18.

2.2. Pan assays interference compounds

The behaviour of final compounds as pan-assay interference compounds (PAINS)⁷² was evaluated through two different web tools SwissADME (<http://www.swissadme.ch>, accessed 2024-01-07)⁷³ and False Positive Remover (<https://www.cbiligand.org/PAINS/login.php>, accessed 2024-01-07).⁷⁴ JChem for Office (21.15.704, 2023) by ChemAxon (<http://www.chemaxon.com>) was used for structure management, SMILES generation, and file conversion. All the analyzed compounds were “accepted” by both the web services.

2.3. *In vitro* carbonic anhydrase inhibition assay

The CA-catalyzed CO₂ hydration activity measurement was performed on an Applied Photophysics stopped-flow instrument using phenol red, at a concentration of 0.2 mM, as a pH indicator with 20 mM HEPES (pH 7.5) as the buffer, 20 mM Na₂SO₄, and following the initial rates of the CA-catalyzed CO₂ hydration reaction for a period of 10–100 s and working at the maximum absorbance of 557 nm. The CO₂ concentrations ranged from 1.7 to 17 mM.⁷⁵ For each inhibitor, six traces of the initial 5–10% of the reaction have been used to determine the initial velocity. The uncatalyzed reaction rates were determined in the same manner and subtracted from the total observed rates. Stock solutions of inhibitor (0.1 mM) were prepared in distilled water, and dilutions up to 0.01 nM were prepared. Solutions containing inhibitor and enzyme were preincubated for 15 min at room temperature before assay to allow the formation of the E-I complex. The inhibition constants were obtained as non-linear least-squares protocols using PRISM 3 and are the mean from at least three different measurements. All CAs were recombinant ones and were obtained in-house.⁷⁶

2.4. Molecular docking

The two-dimensional structures of the compounds were generated using the Maestro module of the Schrödinger Life-Sciences Suite 2024-1 software.⁷⁷ The LigPrep utility was then employed to prepare the compounds to obtain the three-dimensional geometry and identify all possible tautomers and protonation states at pH 7.0 ± 0.4, as determined by Epik.⁷⁸ The sulfonamide group was considered as deprotonated to evaluate the interaction with zinc ion, but after observing no interaction (even considering the zinc constraint and the core constraint), the sulfonamide group was used as unprotonated. The crystal structures of hCA I, hCA II, hCA IX, hCA XI, and EGFR were retrieved from the Protein Data Bank (PDB ID: 6I0J, 3K34, 8Q1A, 5LL5, and 3W2S, respectively).^{79–83}

Protein Preparation Workflow was used to correct, optimize and minimize the crystal structures. Molecular docking analyses were performed using the Glide software. Each enclosing box grid was generated using the centroid of the proper crystallographic ligand and the generated grid file was used for molecular docking using the SP-peptide module of Maestro.⁸⁴ In all CA isoforms, a water molecule was positioned in front of the zinc ion after no interaction between the zinc ion and the sulfonamide group was found. SP-peptide docking protocol was used by setting 5000 poses per ligand for the initial phase and 400 poses per ligand for energy minimization with the OPLS4 forcefield. The docking results were analyzed for the best-docked pose based on the Glide SP score.

2.5. *In vitro* cell proliferation assays

Human triple-negative breast cancer MDA-MB-231 cells (ATCC HTB-262) were cultured in Dulbecco's modified Eagle's medium (DMEM) with 10% fetal bovine serum (FBS), 1% penicillin/streptomycin, and 1% glutamine. Cells were kept at the low passage, returning to their original frozen stocks every 3–4 months. For hypoxic culture conditions cells were grown in 1% O₂ and 5% CO₂. Cell proliferation assays were performed seeding 10,000 cells per well in 48-well plates, and treatments were performed in 1% FBS with increasing concentrations of compounds **8b–d** and **7a**, **SLC-0111**, or **ERL**. After 72 h of incubation at 37 °C, cells detached using trypsin/2 mM EDTA, and cell counting was performed with a MACSQuant® Analyzer (Miltenyi Biotec).

3. Results and discussion

3.1. Synthetic approach

We recently reported the click chemistry approach applied to the hybridization of **ERL**, as an innovative strategy to tackle infections promoted by *Helicobacter pylori*.⁸⁵ Specifically, the hybrids obtained accounted for the **ERL** linked to the prototypic CA-inhibiting chemotype⁸⁶ (compounds **1–4**, Fig. 1, Table 1) and for the pyrimidine-2,4(1*H*,3*H*)-dione moiety as in the antiviral azidothymidine (**AZT**) drug (namely **5**, Fig. 1, Table 1) which is also endowed with antimicrobial features.^{87,88}

The main structural features contained in **1–5** were thought to inhibit the pathogenic expressed CA (HpαCA) which is fundamental for the microorganism survival at the harsh stomach pH values,^{89,90} and to induce overexpression of EGFR in the host cells^{91,92} along with its inhibition by the **ERL** portion.

The synthesized compounds allowed us to define the tolerance of introduced modifications, thus including a preliminary exploration on spacers connecting the two active portions. The enzymatic results on a panel of bacterial CAs were highly promising, highlighting some compounds to potently and selectively inhibit HpαCA (Table 1) over other bacterial isoenzymes and exhibit a moderate anti-*H. pylori* activity, as shown by minimal bacterial concentration (MIC)



Table 1 K_1 values of ERL-clicked derivatives 1–5 and reference drug AAZ, AZT, and ERL on a panel of HpaCA and hCAs by means of stopped-flow hydration assay¹⁰⁹

Cpd	Structure with R =	K_1 (nM)						
		HpaCA*	hCA I*	hCA II*	hCA VA	hCA VI	hCA IX	hCA XII
1a		63.3	70.0	2.0	13.2	360	18.5	n.a.
1b		70.1	85.4	22.0	11.6	174	18.1	n.a.
1c		252	3503	5.0	n.a.	60.1	18.3	28.0
1d		19.3	9.3	4.5	13.0	51.3	20.0	6.0
2a		56.3	3.3	10.4	8.3	39.7	443	7.0
2b		71.6	382	7.4	12.4	99.0	393	n.a.
3a		48.0	35.2	5.4	98.0	2601	40.0	n.a.
3b		43.6	31.2	3.0	107	192	561	n.a.
4		805	2.6	5.0	6.2	28.1	425	n.a.
5		19.3	7.1	3.0	145	103	1445	n.a.
AZT		n.a.	n.a.	684	n.a.	505	n.a.	590
ERL		n.a.	n.a.	3976	n.a.	453	n.a.	n.a.
AAZ		21.0	250	12.1	63.0	16.0	25.7	5.7

K_1 values are reported as means of three independent experiments. Errors are in the range of ± 5 –10% of the reported values. **AAZ** was used as a reference control in these assays. n.a.: not active at the highest concentration tested (100 μ M). Data on HpaCA and hCAs I and II (indicated with an asterisk, *) are already reported by Benito *et al.*⁸⁵



values.⁸⁵ Interestingly, the emerged compound **2b** was also investigated *in silico* and the modification of the ERL structure into the triazole-containing tail was proved not to severely impair the binding to EGFR.⁸⁵

Herein, as a follow-up of our research, this first series of ERL-clicked derivatives were tested on a selected panel of human (h) CAs such as the I, II, VA, VI, IX, and XII to assess their potency and selectivity profiles in comparison to ERL and AZT (Table 1).

Based on data in Table 1 relevant structure–activity relationships (SARs) can be drawn. The presence of the 4-benzenesulfonamide group (**1a**) seems to be responsible for a selective inhibitory profile for hCA II ($K_I = 2.0$ nM), whereas a low nanomolar inhibition was also detected towards hCA VA and hCA IX (K_I values of 13.2 and 18.5 nM, respectively) (selectivity index (SI) values for the tumor-associated hCAs are reported in Table S1 in the ESI† file). Interestingly, no activity was found against the XII isoform. By moving the sulfonamide function in the *meta*-position of the phenyl ring, as in compound **1b**, the potency towards isoforms VA and IX was maintained (K_I values of 11.6 and 18.1 nM, respectively), with no significant difference, such as the lack of activity against hCA XII. Similar activities were also found on Hp α CA and hCA I, whereas an 11-fold decrease in potency was noticed *versus* hCA II (K_I values of 22.0 nM) and a 2-fold lower K_I on hCA VI (= 360 and 174 nM for **1a** and **1b**, respectively). On the other hand, the *ortho*-isomer **1c** showed a very different inhibitory profile. In fact, the compound proved to be highly selective for hCA II, with a low nanomolar inhibition ($K_I = 5.0$ nM), and still maintained the potency against hCA IX ($K_I = 18.3$ nM). However, **1c** reverted the trend against the isoforms VA and XII; indeed, the complete loss of activity towards the former corresponded to a relevant gain in potency on hCA XII ($K_I = 28.0$ nM). Thereby, data for **1a–c** highlighted that the position of the zinc-binding group is not relevant in the inhibition of hCA IX, but fundamental to discriminate potency towards hCAs VA and XII.

A dramatic change was observed by introducing a methylene unit between the triazole ring and the *para*-benzenesulfonamide ring, as in compound **1d**: a net increase in the inhibitory potency is highlighted and low nanomolar K_I values were found for Hp α CA, hCA I, hCA VI and hCA XII (= 19.3, 9.3, 51.3, and 6.0 nM), although the potency against the isoforms II, VA, and IX were not relevantly affected with respect to **1a**. Introducing an amido moiety between the methylene unit and the benzenesulfonamide group in **2a** the inhibition of hCA XII was maintained ($K_I = 7.0$ nM), while resulting in a 2.9- and 22-fold weaker inhibitor of hCAs II and IX ($K_I = 56.3$ and 443 nM), respectively. Shifting the sulfonamide function in *meta*-position leading to compound **2b** resulted in a complete loss of activity towards hCA XII along with a general decrease in potency with the maximum example in the 115-fold reduced inhibition of hCA I ($K_I = 382$ nM), whereas the chemical modification was better tolerated by hCA II and hCA IX. The

affinity towards the hCA XII active site was clearly reset by an additional structural complication, *i.e.*, the insertion of a benzyl group in the tail, resulting in distancing the ERL core from the CA inhibiting function, as in **3a** and **3b**. Also, inhibition of hCA II was quite unaffected (K_I values of 5.4 and 3.0 nM), whereas higher discrepancy was highlighted by enzymatic data on hCA VI ($K_I = 2601$ and 192 nM for **3a** and **3b**, respectively) and hCA IX ($K_I = 40.0$ and 561 nM for **3a** and **3b**, respectively). Surprisingly, despite the isomerism, the compounds shared a similar inhibitory profile against Hp α CA and hCAs I, II, and VA. Additional tail elongation as in compound **4** severely worsened the K_I value till the submicromolar range (= 805 nM) against Hp α CA, while improving the potency against hCAs I, II, VA, and VI. As regards the tumor-associated hCAs IX and XII, while for the former a K_I of 425 nM was detected, inactivity emerged for the latter. In the end, when the hybrid **5** was tested on hCAs IX and XII, again no worthy-of-note inhibition can be highlighted. Moreover, CA inhibition profiling of AZT and ERL did not provide satisfactory results.

The dataset for this series of compounds highlighted that the shorter the linker the higher the inhibitory potency towards the tumor-associated hCAs, with **1a–d** being the most active on hCA IX. Thus, the introduction of additional moieties (amido groups as in **2a** and **2b**) and/or hydrophobic portions (phenyl rings as in **2a**, **2b**, and **4**) in the spacer seemed to cause a loss in activity. However, we tried to slightly increase the linker without implementing the chemical complexity, thereby designing compound **6**, being a homologue compound that bears only a methylene unit more (Table 2, synthesis in Scheme S1 in the ESI† file).

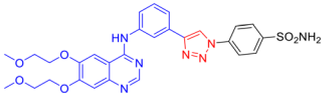
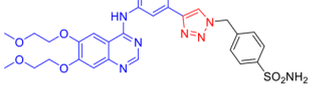
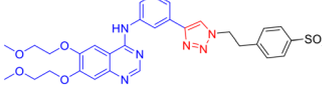
Data in Table 2 showed compound **6** being a weak inhibitor of hCA I, with a micromolar inhibitory potency *versus* this isoenzyme ($K_I = 1031$ nM), and possesses a stronger affinity towards hCAs II and IX, showing K_I values of 18.4 and 9.3 nM, respectively. Interestingly, the compound exhibited a higher preference for hCA IX with respect to its homologues **1a** and **1d**, as highlighted by the SI values. However, the activity towards hCA XII decreased reaching the medium nanomolar range ($K_I = 353$ nM), also impacting the SI.

However, representative compounds of the ERL-clicked series (**1a–c**) were selected along with the reference hCAs IX and XII inhibitor SLC-0111, and ERL to evaluate their anti-proliferative effect *in vitro* on non-small cell lung carcinoma A549 and pancreatic epithelioid carcinoma PANC-1 cell lines, both characterized by overexpression of the tumor-associated hCA IX^{93,94} and EGFR^{95,96} (Fig. S1 in the ESI† file). The reference compound SLC-0111 was found to be ineffective in reducing the proliferation of both the cell lines and only ERL and compound **1c** displayed an antiproliferative effect, with the latter being the most effective and possessing IC₅₀ values of 16.3 and 57.1 μ M on A549 and PANC-1 cells, respectively.

The overall promising results encouraged further investigation and the design of a second series of hybrids



Table 2 K_i values of ERL-clicked derivative **6** and homologues **1a** and **1d** on hCAs I, II, IX, and XII and related SIs by means of stopped-flow hydration assay

Cpd	Structure	K_i (nM)				SI			
		hCA I	hCA II	hCA IX	hCA XII	I/IX	II/IX	I/XII	II/XII
1a		63.3*	70.0*	18.5	n.a.	3.42	3.78	—	—
1d		19.3*	9.3*	20.0	6.0	0.96	0.47	3.21	1.55
6		1031	18.4	9.3	353	111	1.98	2.92	0.05

K_i values are reported as means of three independent experiments. Errors are in the range of ± 5 –10% of the reported values. **AAZ** was used as a reference control in these assays. n.a.: not active at the highest concentration tested (100 μ M). Data indicated with an asterisk (*) were already reported by Benito *et al.*⁸⁵ SI values are calculated as the ratio between the K_i values of the physiologically relevant CA isoform I or II (as indicated) and the CA isoform of interest (hCA IX or hCA XII, as indicated). The higher the SI value, the higher the isoform preference.

aimed at enhancing the compounds solubility. Thereby, in place of the traditional methylene chains,^{20,97} we introduced non-cleavable monodispersed poly(ethylene glycol) (PEG) spacers between the pharmacophores interacting with hCAs and EGFR. Thus, we selected 4- and 6-unit PEG spacers to hold a precise length and good flexibility, which should ensure the two active portions could better fit into their targeted binding sites. In fact, due to their non-toxicity and amphiphilic nature, PEG polymers have garnered attention in pharmaceutical technologies, being included in formulations as excipients or vehicles/carriers of small molecules, biopharmaceuticals (*e.g.*, peptides, antibody-drug conjugates), and drug delivery systems (*e.g.*, liposomes and nanoparticles) or covalently bound to the drug of interest.⁹⁸ In this case, PEGylated biopharmaceuticals⁹⁹ and small molecule drugs¹⁰⁰ resulted in decreased immunogenicity and improved half-life in blood and showed increased metabolic stability, solubility, and overall enhanced pharmacokinetic properties. Known examples are the FDA-approved PEGylated naloxegol,¹⁰¹ PEG-docetaxel,¹⁰² PEG-camptothecin,¹⁰³ and many others.¹⁰⁴ However, few small molecules-based hybrid compounds have been reported to possess PEG spacer¹⁰⁵ with respect to the widely explored PROTACs¹⁰⁶ and antibody-drug conjugates.¹⁰⁷

Thus, we generated seventeen new compounds as PEGylated **ERL**–CAI hybrids (**7**–**14**, Fig. 1) through the Huisgen click chemistry reaction. After synthesis, the compounds were then tested on hCAs I and II along with the tumor-associated isoforms IX and XII, and docking simulations helped the rationalization of the enzymatic results, and the affinity towards EGFR was investigated *in silico*. In the end, to assess the compounds antiproliferative effect, we moved towards the triple-negative breast cancer (TNBC) cell line MDA-MB-231, characterized by the upregulation of both EGFR and the tumor-associated hCAs.¹⁰⁸

3.2. Synthesis of the ERL-based library

The synthetic pattern aimed at introducing PEG (= 4 or 6 oxyethylene units) spacers between the CA-interacting functionalities, such as benzenesulfonamide and coumarin, and **ERL**. The first series of derivatives (**7**–**9**) was successfully obtained by a two-step procedure involving the insertion of the PEG-4 linker by means of the nucleophilic attack of the amino head of **15** to the electrophilic carbamates **16a–b** or isothiocyanates **17a–d** and **18** on the benzenesulfonamide tails. The obtained intermediates **19a–b**, **20a–d**, and **21** were then reacted with **ERL** via a CuAAC by using nanosized copper in presence of tetramethylammonium chloride (TMACl) in DMF, affording the final compounds **7**–**9** (Scheme 1).

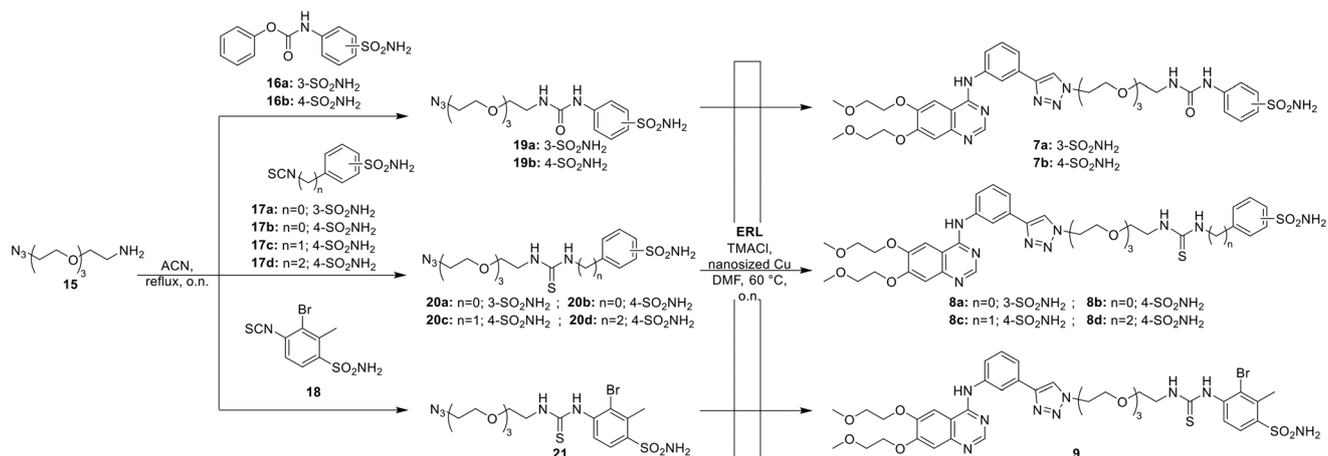
The second series of derivatives (**10**–**14**) was synthesized starting from the *N*-hydroxysuccinimide ester-functionalized PEG-6 (**22**) and suitable amines **23a–b** and **24**, gaining intermediates **25a–b** and **26**, tailored with benzenesulfonamide and coumarin cores, respectively (Scheme 2). Similarly, the amino group of the PEG-6 chemical **27** reacted with carbamates **16a–c** and **28** and thiocarbamates **17b–d** to obtain intermediates **29**–**32**. In the end, all the intermediates were clicked on **ERL** by using copper sulfate and sodium ascorbate in presence of TMACl, yielding the final compounds **10**–**14** (Scheme 2).

3.3. *In vitro* enzymatic inhibition on CAs

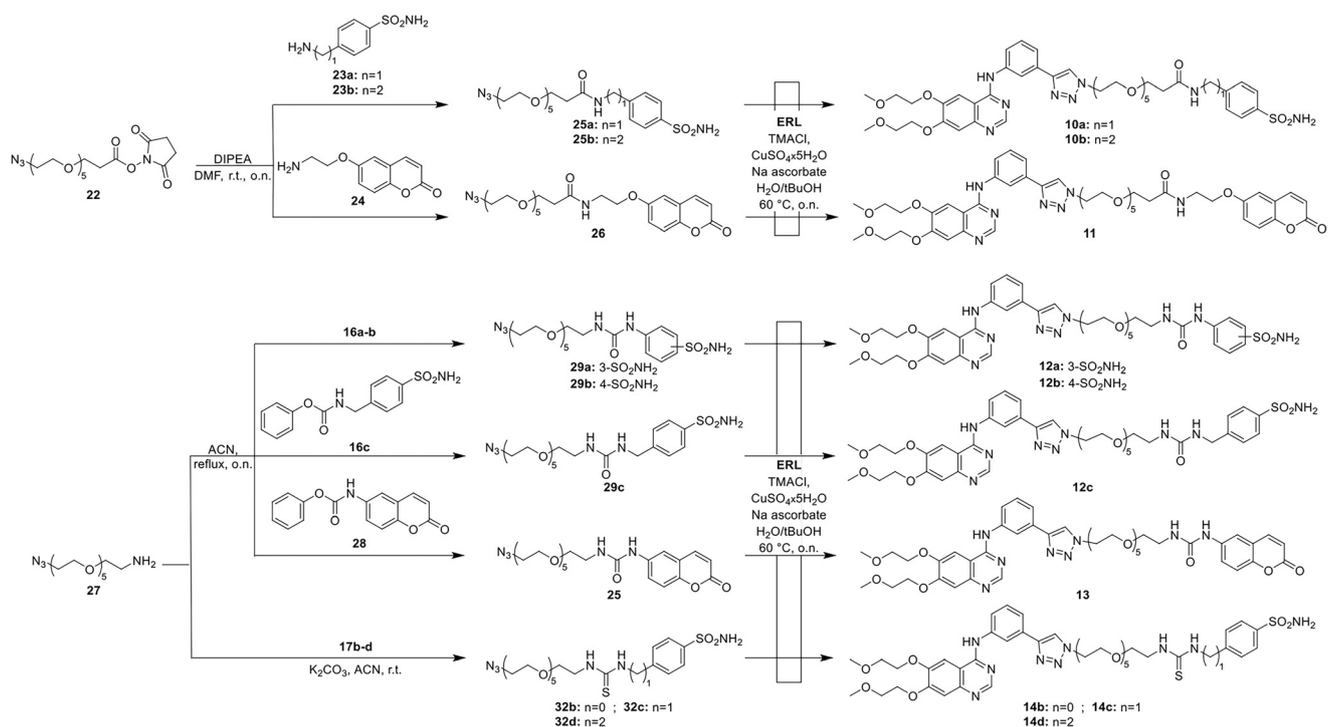
This library of **ERL**-derived compounds was tested for the capability to exert selective inhibition of a panel of four human CAs using the stopped-flow CO₂ hydration assay.¹⁰⁹ Data obtained were also compared with those of the reference **AAZ** and reported in Table 3 as K_i values.

As reported in Table 3, interesting SARs for each CA isoform can be deduced.





Scheme 1 Synthesis of hybrids 7–9.



Scheme 2 Synthesis of hybrids 10–14.

i) As for the hCA I, all the derivatives are not strongly effective inhibitors, being the associated K_i values in the micromolar range (except for nanomolar inhibitors **8d**, **10a**, **10b**, **14d**, and above all **12c**). Compounds showed less potency when the PEG units are 6 (**10–14**), while the inhibitory activity is improved with PEG-4 (**7–9**). The CA inhibiting warhead must be the benzenesulfonamide with respect to the coumarin ring (**11** and **13**). In the presence of PEG-4, thiourea is preferred over urea especially if not close to the ArSO₂NH₂ ($n = 2 > 1 > 0$). 3-SO₂NH₂ and 4-SO₂NH₂ contributed equally to the inhibitory activity. If the linker is PEG-6, thiourea is like urea especially if not

close to the ArSO₂NH₂ ($n = 2 > 1 > 0$). The amide was a good option to substitute the (thio)urea moiety but only if $n = 2 > 1$.

ii) Overall, the kinetic trend for the hCA II is better for almost all the compounds (**23c** is even in the picomolar range $K_i = 0.8$ nM). Again, PEG-6 is preferred over PEG-4 and the CA inhibiting warhead must be the benzenesulfonamide with respect to the coumarin ring (**11** and **13**). If the linker is PEG-4, thiourea is more favored than urea, especially if $n = 2 > 0 > 1$. Conversely, if the linker is PEG-6, urea is better ($n = 1 > 0 > 2$) than thiourea, whereas the amide moiety can be a valid alternative.



Table 3 K_1 and SI values of ERL-clicked derivatives 7–14 and reference drugs ERL and AAZ versus a panel of hCAs and by means of stopped-flow hydration assay¹⁰⁹

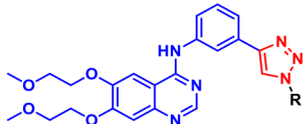
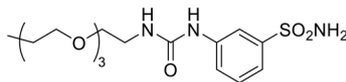
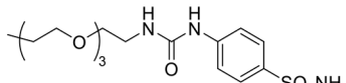
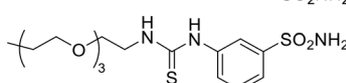
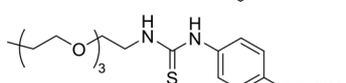
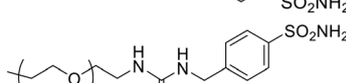
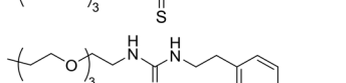
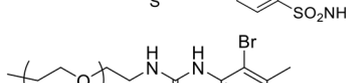
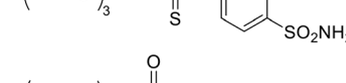
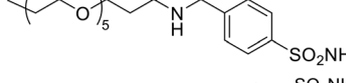
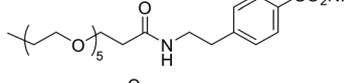
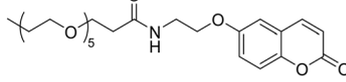
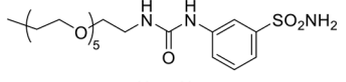
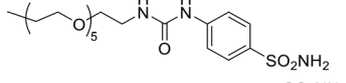
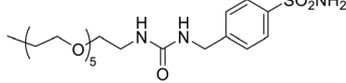
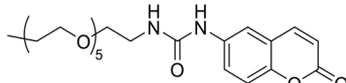
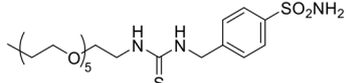
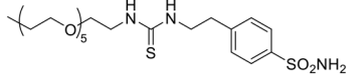
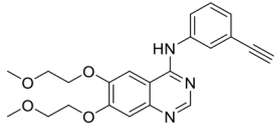
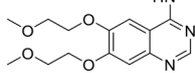
Cpd	Structure with R = 	K_1 (nM)				SI			
		hCA I	hCA II	hCA IX	hCA XII	I/IX	II/IX	I/XII	II/XII
7a		46 546	723	285	58.9	164	2.54	790	12.3
7b		36 052	950	1124	57.4	32.1	0.85	628	16.6
8a		45 796	3169	1924	72	23.8	1.65	636	44.0
8b		3144	135	112	5.33	28	1.2	590	25.4
8c		3277	1150	26	69.7	127	44.57	47.0	16.5
8d		923	106	28	6.49	32.6	3.73	142	16.3
9		4909	531.0	1345	69.8	3.65	0.39	70.3	7.61
10a		903	253	4158	3305	0.22	0.06	0.27	0.08
10b		632.4	33.5	3673	1363	0.17	0.01	0.46	0.02
11		70 140	33 700	4677	n.a.	15.0	7.21	—	—
12a		70 090	1023	4473	6403	15.7	0.23	11.0	0.16
12b		24 790	90.8	5620	75.3	4.41	0.02	329	1.21
12c		160	0.80	387	940	0.41	0.002	0.17	0.0008
13		28 301	11 364	242	8059	117	46.9	3.51	1.41
14b		43 340	9712	4928	1523	8.79	1.97	28.4	6.38



Table 3 (continued)

Cpd	Structure	K_i (nM)				SI			
		hCA I	hCA II	hCA IX	hCA XII	I/IX	II/IX	I/XII	II/XII
									
14c		4976	1011	3727	665	1.33	0.27	7.48	1.52
14d		957	32 703	1877	95.2	0.51	17.4	10.0	344
ERL		n.a.*	3976*	n.a.	n.a.	—	—	—	—
AZT		250	12.1	25.7	5.7	9.73	0.47	43.86	2.12

K_i values are reported as means of three independent experiments. Errors are in the range of ± 5 –10% of the reported values. **AZT** was used as a reference control in these assays. n.a.: not active at the highest concentration tested (100 μ M). Data indicated with an asterisk (*) were already reported by Benito *et al.*⁸⁵ SI values are calculated as the ratio between the K_i values of the physiologically relevant CA isoform I or II (as indicated) and the CA isoform of interest (hCA IX or hCA XII, as indicated). The higher the SI value, the higher the isoform preference.

iii) For hCA IX, the general situation is preferred for PEG-4. In this series, 4-SO₂NH₂ impacts positively the inhibitory values with respect to 3-SO₂NH₂, and thiourea and urea contributed equally ($n = 1 \geq 2 > 0$). In the subset PEG-6, compounds are almost in the same micromolar range (except for **12** and **13**).

iv) The most interesting data were registered for hCA XII: if the linker is PEG-4 (better than 6), 4-NH₂SO₂Ar improves the inhibitory values with respect to 3-NH₂SO₂Ar and, again, thiourea is preferred to urea with $n = 0-1 > 2$. In the subset PEG-6, coumarins are still not important for the activity (high micromolar activity for compound **13** and no activity for compound **11**) and amide is more tolerated than (thio)urea especially if the 4-SO₂NH₂ moiety is present and with $n = 2 > 0 > 1$ in the thiourea series and $n = 0 > 1$ for ureas.

Isoform selectivity could be easily assessed by analyzing SI values reported in Table 3. Taking into consideration the first subset of PEG-4 derivatives (**7**–**9**), the higher affinity towards hCA XII was demonstrated by a large number of compounds with respect to isoforms I and II, with SI (I/XII) values ranging from 47.0 up to 790. Conversely, hCA IX inhibitory data almost overlapped with the values determined against hCA II in terms of potency range, not showing remarkable selectivity. However, the PEG-6 derivatives series (**10**–**14**) required a more accurate analysis of SI values since each compound seem to show a different isoform preference.

3.4. *In silico* studies on targeted enzymes

To rationalize the enzyme inhibition data, structure-based computational studies were conducted on all studied CA

isoforms. The crystallographic structures of hCAs I, II, IX, and XII were obtained from the Protein Data Bank (PDB ID: 6I0J, 3K34, 8Q1A, and 5LL5, respectively). The selection of the PDB structures, validated in our previous works as well, was made taking into account the good crystallographic resolution and the absence of structural defects in the chosen structures. The investigation focused on compounds demonstrating the highest activity toward hCAs IX and XII while exhibiting selectivity over hCAs I and II. Thereby, the *in silico* study included compounds **8b**, **8c**, and **8d** (Table 3).

Given their chemical structure, dimension, and flexibility, the Glide SP-peptide docking method was employed. In the initial trial, the sulfonamide group was assumed to be deprotonated in order to ascertain the potential for interaction with the zinc ion in the active site. This binding mode has not yet been observed in any X-ray crystal structure of CA-sulfonamide adducts⁵⁴ and should be checked experimentally in order to be validated. Unexpectedly, the docking poses obtained without any constraint do not exhibit the sulfonamide functionality of the compounds coordinated with the zinc ion. In some cases, the **ERL**-derived portion of the three compounds interacts with some residues in the active site. The protocol was then modified by introducing a constraint on the zinc ion, yet no poses in which this ion coordinates the sulfonamide were obtained. Subsequently, the core constraint approach was employed to force the sulfonamide group to coordinate the metal center, but the results were again negative. Because of the evidence that there is no direct interaction between the zinc ion and the sulfonamide group of compounds in this theoretical *in silico* approach, a coordination water molecule was introduced to



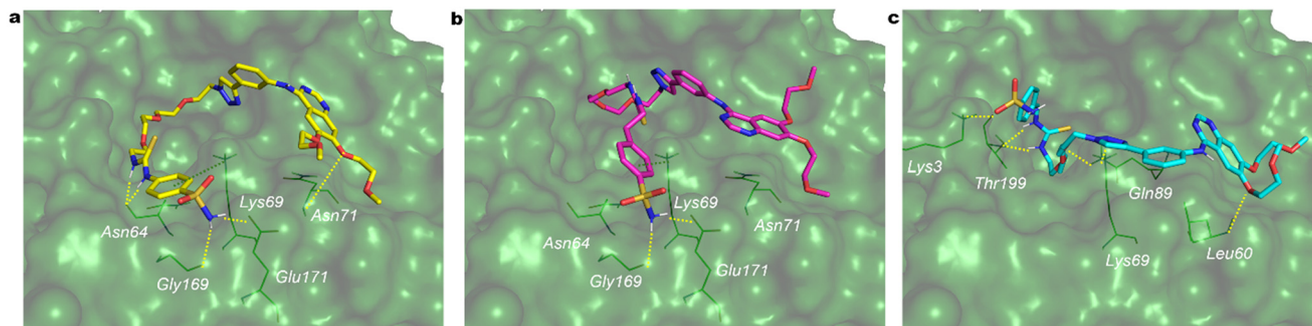


Fig. 2 Predicted 3D binding mode within hCA XII (green surface) of compounds (a) **8b** (yellow sticks), (b) **8d** (magenta sticks), and (c) **8c** (cyan sticks). H-bond interacting residues are shown as green lines while H-bonding and cation- π interactions as yellow and green dotted lines, respectively.

the zinc ion in the hCA isoforms, with the sulfonamide group assumed to be in its undeprotonated form at physiological pH. The docking poses obtained with the docking SP-peptide were clustered, and the representative geometry of the most abundant cluster for each compound and each enzyme was considered. In hCA XII, compounds **8b** and **8d** (Fig. 2a and b) are positioned outside the active site and form a cation- π interaction between Lys69 and the phenyl ring of the **ERL** fragment. Additionally, the sulfonamide NH_2 establishes two H-bonds with the carbonyl of the Gly169 backbone and the carbonyl oxygen of Glu171. In the case of **8d**, in addition to the H-bond between the amide nitrogen of Asn71 and the proximal oxygen of the 2-methoxyethoxy tail of **ERL**, the two nitrogen atoms of the thiourea directly linked to benzenesulfonamide mediate two H-bonds with Asn64.

The arrangement of compound **8c** (Fig. 2c), which possesses 10 times less activity, differs from that of the other compounds. It enters the active site with the thiourea group, whose two nitrogen atoms mediate two H-bonds with the oxygen of Thr199. Additionally, the first oxygen atom of the ethoxy linker forms an H-bond with the nitrogen of Gln89. At the edge of the active site, the sulfonamide forms an H-bond with the amine group of Lys3, while H-bonds between the nitrogen of the triazole and Lys69 and between Leu60 and the oxygen of the **ERL** tail are observed outside the active site.

Compounds **8c** and **8d** (Fig. 3b and c), exhibiting a potent inhibition of hCA IX, gain access to the active site of hCA IX, despite their differing geometries. The sulfonamide moiety of **8b** or **8d** (Fig. 3a and b) forms an H-bond with the zinc-coordinated water molecule, while the aromatic ring of **8d** engages in a π - π interaction with the zinc-coordinated His94. The nitrogen of the thiourea moiety of the three compounds forms an H-bond with His68.

In the region external to the active site, the quinazoline moiety of the **ERL** portion of **8c** (Fig. 3c) engages in a cation- π interaction with Arg62, whereas the terminal methoxy of the 6-(2-methoxyethoxy) substituent of **8d** forms an H-bond with Arg89. Furthermore, two oxygen atoms of the PEG linker of **8d** form H-bonds with Arg64. As regards compound **8b**, no strong interactions are observed outside the active site.

All three compounds are positioned on the surface of hCA I without entering the active site, justifying the lack of activity toward this isoform. However, they display distinct binding modes within the hCA II isoform. Interestingly, **8c** remains outside the active site, exhibiting no evidence of bonding that could justify its observed activity. On the other hand, **8d** (Fig. 4b) enters the active site with the **ERL** moiety, and the two methoxy groups form H-bonds with Asn62 and Gln92. The triazole moiety engages in a π - π interaction with Phe131, the nitrogen atom of the thiourea moiety forms an H-bond

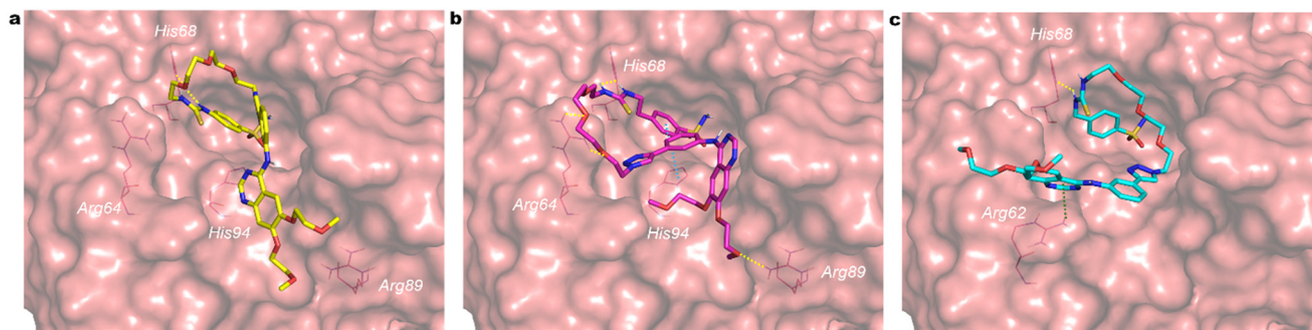


Fig. 3 Predicted 3D binding mode within hCA IX (pink surface) of compounds (a) **8b** (yellow sticks), (b) **8d** (magenta sticks), and (c) **8c** (cyan sticks). H-bond interacting residues are shown as magenta lines, while H-bonding, π - π interactions, and cation- π interactions as yellow, cyan, and green dotted lines, respectively.



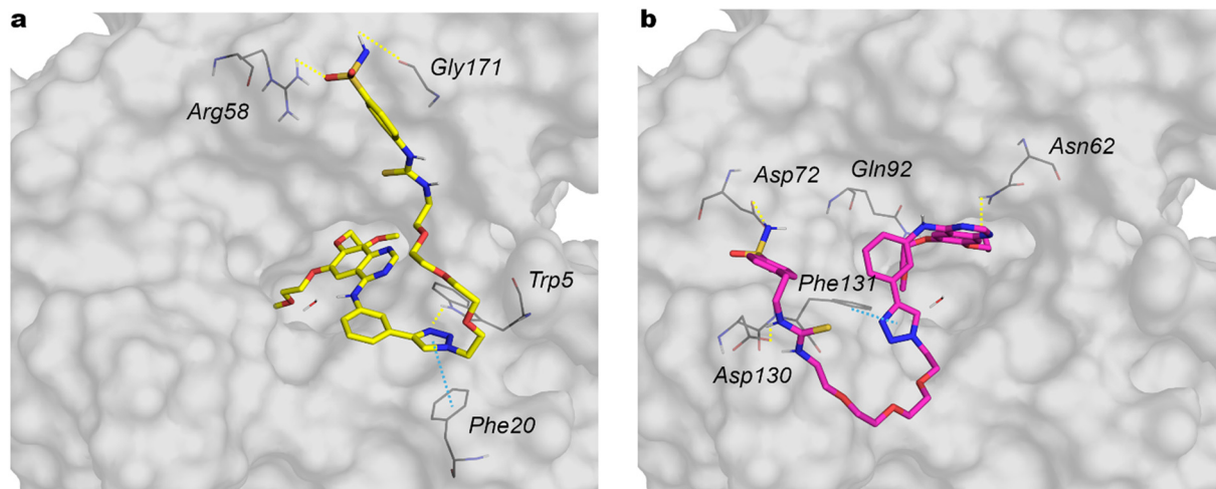


Fig. 4 Predicted 3D binding mode within hCA II (grey surface) of compounds (a) **8b** (yellow sticks) and (b) **8d** (magenta sticks). H-bond interacting residues are shown as green lines while H-bonding and π - π interactions as yellow and cyan dotted lines, respectively.

with Asp130, whereas the nitrogen and oxygen atoms of the sulfonamide moiety form H-bonds with Asp72 and the backbone of Phe131, respectively. Conversely, **8b** (Fig. 4a) enters the active site with the **ERL** portion, and the only notable interactions outside the active site are H-bonds between the triazole nitrogen and Trp5, and between the oxygen and nitrogen atoms belonging to the sulfonamide moiety and Arg58 and Gly171, respectively. Furthermore, a π - π interaction between the triazole and Phe20 can be observed.

In summary, compounds **8b**, **8c**, and **8d** were proven to establish favorable interactions within and outside the active site of the hCAs IX and XII isoforms, thereby justifying their observed activity and selectivity.

Although the activity toward EGFR kinases has yet to be evaluated, we conducted a docking study on the EGFR kinase domain with the same compounds to evaluate a potential dual activity. Among the available protein in the Protein Data Bank, the PDB ID 3W2S was selected on the basis of the dimensions of the crystal ligand, which is comparable to those of compounds under investigation. The three

compounds were observed to dock within the ATP binding site, forming hydrophobic interactions. Furthermore, compounds **8c** and **8d** (Fig. 5a and b) establish H-bonds between the sulfonamide and Lys875 (**8c** with Val876), while **8b** interacts with Ala859. Lys745 forms an H-bond with the oxygen of the oxyethylene linker of **8d** and with two oxygens of **8c**. The aromatic ring of the benzenesulfonamide of **8c** forms a cation- π interaction with Arg841, while the nitrogen atoms of the thiourea of **8b** and **8d** engage in H bond interactions with Gly857 and Asp855, respectively. These observations are consistent with the docking score and conserved key interactions, which indicate that **ERL** derivatives retain the structural features necessary for binding to the EGFR kinase domain.

3.5. Anti-proliferative effect on TNBC cells

TNBC is one of the most reported causes of death in the female population worldwide¹¹⁰ and one of its distinguishing features compared to other breast cancers is the overexpression of EGFR.¹¹¹⁻¹¹⁴ Interestingly, this receptor was

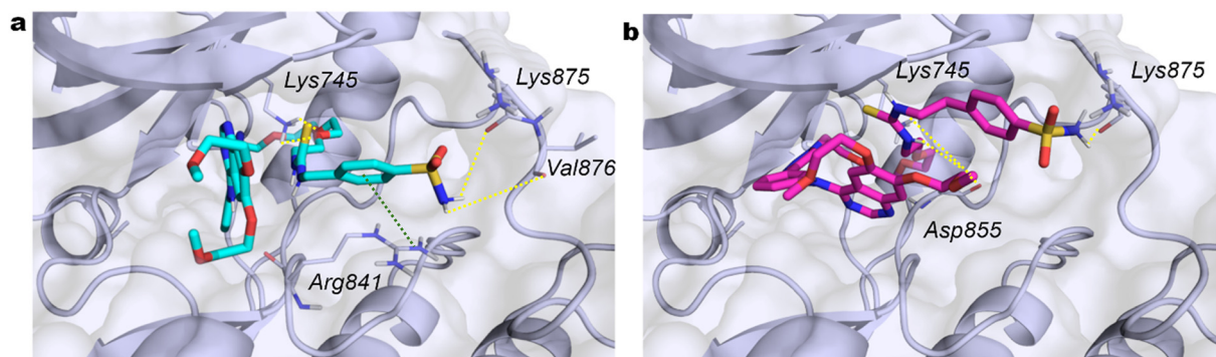


Fig. 5 Predicted 3D binding mode within EGFR kinase domain (blue-grey surface and cartoon) of compounds (a) **8c** (cyan sticks) and (b) **8d** (magenta sticks). H-bond interacting residues are shown as grey lines while H-bonding and cation- π interactions as yellow and green dotted lines, respectively.



proven to be activated in several TNBC cell lines¹¹⁵ and its inhibition was demonstrated to arrest tumor growth and metastasis in TNBC xenograft models^{116,117} by controlling the initial progression of the disease.¹¹⁸ However, drug resistance to **ERL** has been recorded in some patients^{119–121} with an unclear mechanism.¹⁰⁸ Also, **ERL** seems to inhibit the non-cancer stem cells, responsible for stemness and drug-resistance, or bulk TNBC cells.¹²²

Thus, we selected representative compounds of the new series (**8b–d** and **7a**) to evaluate their anti-proliferative effect *in vitro* on the human TNBC cell line MDA-MB-231 that expresses both the tumor-associated hCA isoenzymes and EGFR.^{60,123} We also tested the known hCAs IX and XII inhibitor **SLC-0111**, and **ERL**, as reference compounds (Fig. 6). Cancer cells were treated with the compounds at increasing concentrations under hypoxic culture conditions, and the anti-proliferative readout was assessed after 72 hours by flow cytometry-based cell counting. As shown in Fig. 6, the reference compound **SLC-0111** was found to be effective in reducing the proliferation of MDA-MB-231, with an IC_{50} of 23.4 μ M. Interestingly, compounds **8b** and **8d** displayed the most promising anti-proliferative effect, with IC_{50} values ranging from 30 and 48.51 μ M, respectively. A significantly lower effect on tumor cells was observed for compound **8c** (IC_{50} ~190 μ M). Notably, treatment with **ERL** resulted only in a mild reduction of cell proliferation observed only at the higher 100 μ M concentration tested, in accordance to our previous studies underlining its lack of efficacy on this

cell line,¹²² thus supporting the increased effect obtained with the novel **ERL**-derived compounds.

4. Conclusions

In summary, a second series of seventeen new Pegylated **ERL**-CAI hybrids was synthesized through the Huisgen click chemistry reaction, bearing the benzenesulfonamide or the coumarin moiety as pharmacophores. Two different linker lengths were selected, *i.e.* 4- and 6-unit PEG spacers, to hold good flexibility and simultaneously explore the best length for coordination with the zinc ion within the CA active cleft, thus enhancing pharmacokinetic properties. Most of the derivatives were found to strongly inhibit hCAs IX and XII at low-to-high nanomolar concentration, showing isoform selectivity in a large number of compounds. Concerning hCA IX the general situation shows a remarkable preference for PEG-4 derivatives. Moreover, no differences between thiourea and urea emerged, while the sulfonamide moiety position showed a high impact in K_1 values ($4-SO_2NH_2 > 3-SO_2NH_2$). The same linker length preference is confirmed against hCA XII, in which the PEG-4 linker showed the best results. As well as previous isoform, thiourea is preferred to urea for hCA XII inhibition, with $n = 0-1 > 2$, and, similarly to hCA IX, $4-SO_2NH_2$ improved inhibitory features with respect to $3-SO_2NH_2$. Selectivity index (SI) values were then used to assess isoform selectivity: compounds 7–9 (PEG-4) showed a high preference for hCA XII (SI up to 790.25) compared to hCA I and II, while compounds 10–14 (PEG-6) had similar

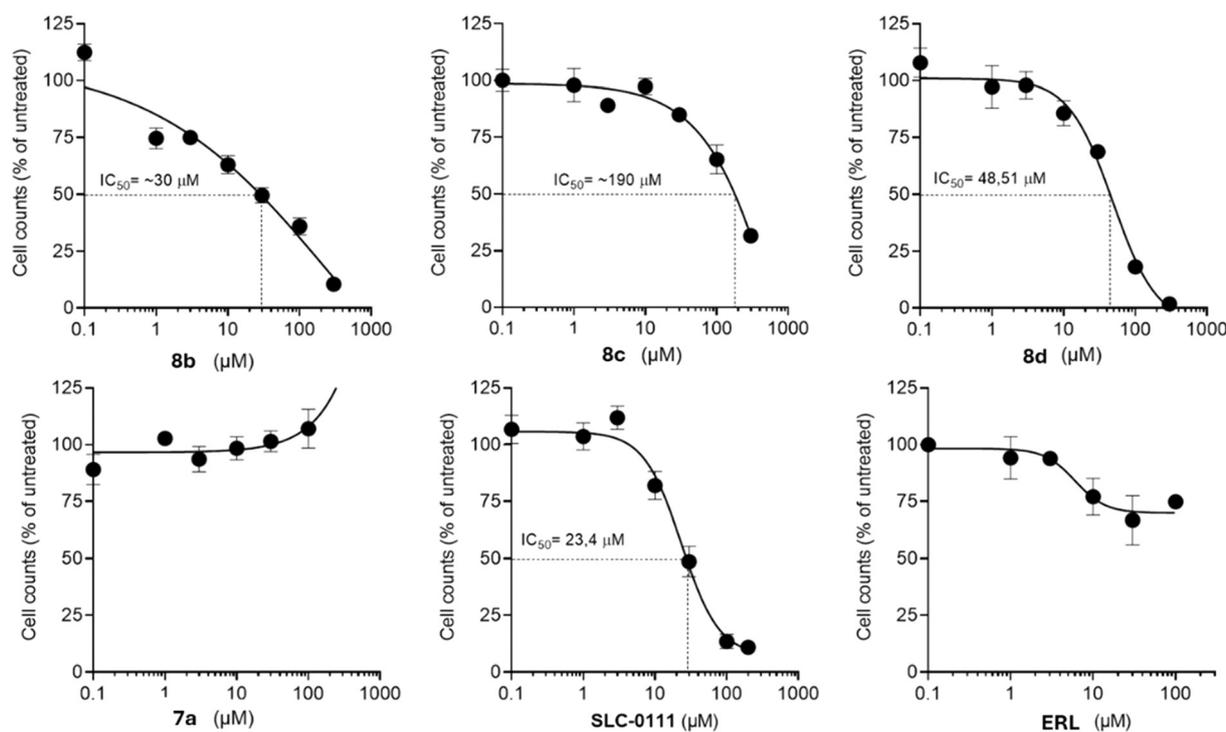


Fig. 6 Cell proliferation of TNBC MDA-MB-231 cells treated for 72 hours in hypoxic conditions with compounds **8b–d** and **7a**, **SLC-0111**, or **ERL**. Cell count is referred to the untreated/control considered as 100%, mean \pm SEM, and the IC_{50} are reported.



activity against hCA IX and II, with some selectivity over hCA I. The lower SI values (down to 345.52) for the latter group reflect their stronger inhibition of hCA IX and XII.

Computational molecular docking studies on hCA isoforms rationalize the inhibitory activity and selectivity of compounds **8b**, **8c**, and **8d**. Concerning hCA XII, **8b** and **8d** bind outside the active site, with key cation- π and hydrogen bonding interactions. Conversely, **8c** enters the active site differently. For hCA IX all three compounds showed binding within the active site, with the sulfonamide group of **8b** and **8d** forming an H-bond with a zinc-coordinated water molecule, and the aromatic ring of **8c** interacting with His94. Moreover, minimal interactions in the active sites of hCA I and II do support compounds' selectivity profile. Notably, docking study to the EGFR kinase domain indicated potential for dual activity. Thus, compounds **8b**, **8c**, and **8d** docked within the ATP binding site and retained key binding features, indicating the conservation of structural elements needed for binding to the EGFR domain, as well. We further evaluated new synthesized hybrids compounds for TNBC treatment, using the MDA-MB-231 cell line which expresses EGFR and hCAs. Compounds **8b** and **8d** demonstrated strong anti-proliferative activity (IC_{50} 30 and 48.51 μ M respectively), thus comparable to the single target hCA inhibitor **SLC-0111** (IC_{50} = 23.4 μ M) and in contrast to the low efficacy observed for the EGFR inhibitor **ERL**. These results highlight the potential of such compounds for further development within TNBC therapy and more.

Data availability

The data supporting this article have been included as part of the ESI†. The following data are made available in the ESI†: selectivity index (SI) values for the first ERL-clicked derivatives (Table S1†); synthesis of compound **6** (Scheme S1†); procedure for the synthesis of compound **6**); antiproliferative effect of the first series of ERL clicked derivatives (Fig. S1†); 1H and ^{13}C NMR of representative compounds. The ESI† file contains the NMR spectra of the compounds reported in this article.

Author contributions

Conceptualization, review and editing, S. C., I. D. A., A. A., F. C., and C. T. S.; methodology and formal analysis, G. R., S. F., M. F., G. B., M. L. M., R. S., C. C., R. R.; funding acquisition, F. C., R. R.; writing – original draft preparation, F. C., R. R., C. S., and I. D. A. All authors have read and agreed to the final version of the manuscript.

Conflicts of interest

There are no conflicts of interest to declare.

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