

### **BRIEF REPORT**

**3** OPEN ACCESS



# 5-(Sulfamoyl)thien-2-yl 1,3-oxazole inhibitors of carbonic anhydrase II with hydrophilic periphery

Stanislav Kalinin<sup>a,b</sup> (D), Alexander Kovalenko<sup>a</sup>, Annika Valtari<sup>b</sup>, Alessio Nocentini<sup>c</sup> (D), Maxim Gureev<sup>d</sup> (D), Arto Urtti<sup>a,b,e</sup> (D), Mikhail Korsakov<sup>f</sup>, Claudiu T. Supuran<sup>c</sup> (D) and Mikhail Krasavin<sup>a,g</sup> (D)

<sup>a</sup>Institute of Chemistry, Saint Petersburg State University, St. Petersburg, Russian Federation; <sup>b</sup>School of Pharmacy, University of Eastern Finland, Kuopio, Finland; <sup>c</sup>Department of Neurofarba, Universita degli Studi di Firenze, Florence, Italy; <sup>d</sup>Digital Biodesign and Personalized Healthcare Research Center, Sechenov First Moscow State Medical University, Moscow, Russian Federation; <sup>e</sup>Drug Research Program, Division of Pharmaceutical Biosciences, Faculty of Pharmacy, University of Helsinki, Helsinki, Finland; <sup>f</sup>Pharmaceutical Technology Transfer Center, Ushinsky Yaroslavl State Pedagogical University, Yaroslavl, Russian Federation; <sup>g</sup>Immanuel Kant Baltic Federal University, Kaliningrad, Russian Federation

### **ABSTRACT**

Hydrophilic derivatives of an earlier described series of carbonic anhydrase inhibitors have been designed, prepared and profiled against a panel of carbonic anhydrase isoforms, including the glaucoma-related hCA II. For all hydrophilic derivatives, computational prediction of intraocular permeability routes showed the predominance of conjunctival rather than corneal absorption. The potentially reactive primary or secondary amine periphery of these compounds makes them suitable candidates for bioconjugation to polymeric drug carriers. As was shown previously, the most active hCA II inhibitor is efficacious in alleviating intraocular pressure in normotensive rabbits with efficacy matching that of dorzolamide.

#### **GRAPHICAL ABSTRACT**

#### **ARTICLE HISTORY**

Received 17 February 2022 Revised 15 March 2022 Accepted 18 March 2022

#### **KEYWORDS**

Glaucoma; intraocular pressure; hydrophilicity; bioconjugation; intraocular delivery

# Introduction

Glaucoma-related high intraocular pressure can be alleviated by the use of eye drops of prostaglandin analogues, beta blocking agents and carbonic anhydrase inhibitors (CAIs)<sup>1</sup>. The recent approval of rho kinase inhibitors and NO donors significantly expands the range of treatment options<sup>2,3</sup>. The clinically used topical CAIs for glaucoma treatment include dorzolamide (1) and brinzolamide (2), compounds that are (a) relatively lipophilic and (b) non-selective as inhibitors of a particular carbonic anhydrase isoform<sup>4</sup>. Acetazolamide (3) and methazolamide (4) are also used as anti-glaucoma agents (Figure 1), but they are oral medications which frequently cause adverse drug reactions<sup>5</sup>. Potent and selective inhibition of carbonic anhydrase II isoform (hCA II) is an important mechanism of action due to the critical importance of this enzyme in reduction of glaucoma-related intraocular pressure<sup>6</sup>.

Topical ocular drugs are typically designed as rather lipophilic, because they absorb to the eye across the cornea<sup>7</sup>. Lipophilicity leads to decreased water solubility and, thus, lowers the achievable drug concentration in the tear fluid. On the contrary, higher concentration in the tear fluid can be achieved with hydrophilic compounds. Such compounds may absorb into their ocular targets via conjunctiva and sclera that allow permeation of relatively hydrophilic compounds<sup>8</sup>. Specifically designing hydrophilic compounds that can utilise this route will lower the loss of hydrophilic compounds to the blood stream across conjunctiva<sup>8</sup>. Anti-glaucoma CAIs exert their action in the ciliary body located next to sclera, thereby making non-corneal absorption of highly potent, hydrophilic derivatives an interesting approach. Moreover, in comparison to the cornea, the conjunctiva has wider inter-cellular space for permeation of hydrophilic compounds<sup>9</sup>.

Previously, we described a series of 5-(sulfamoyl)thien-2-yl 1,3-oxazoles **5a-c** which displayed a remarkably potent inhibition

CONTACT Claudiu T. Supuran claudiu.supuran@unifi.it Department of Neurofarba, Universita degli Studi di Firenze, Florence, Italy; Mikhail Krasavin mkrasavin@hotmail.com, krasavintm@gmail.com la Institute of Chemistry, Saint Petersburg State University, St. Petersburg, Russian Federation

**(b)** Supplemental data for this article can be accessed <u>here.</u>

Figure 1. Clinically used antiglaucoma carbonic anhydrase inhibitors.

$$K_{i}$$
 ( $hCA$  II),  $nM$ 
 $K_{i}$  ( $hCA$  II),  $nM$ 
 $R'$ 
 $NH$ 
 $NH$ 

Figure 2. Earlier reported potent hCAII inhibitors 5(6)a-c and their modified hydrophilic thiophene analogues 7 designed and investigated in this work.

profile towards human carbonic anhydrase (CA, EC 4.2.1.1) and, in particular, its hCA II isoform<sup>10</sup> which is the primary target for intraocular pressure-reducing antiglaucoma drugs.<sup>6</sup> Later on, a related – and similarly potent against hCA II – benzenesulfonamide series (6a-c) showed high efficacy in vivo lowering ocular hypertension in rabbits. Furthermore, the high potency and the pronounced selectivity towards the CA isoform of this series was rationalised by X-ray crystallographic structure of complex of 6c with the protein<sup>11</sup>. Considering that compounds **5a-c** contain the primary sulphonamide group linked to a thiophene moiety, it makes them structurally closer to the clinically used drugs 1-4 all of which have a five-membered heterocyclic core as a primary sulphonamide-bearing scaffold. Thus, we selected carboxamides **5b-c** as the prototype scaffold for the introduction of peripheral functional groups which would increase the resulting compounds' hydrophilicity and also a reactive 'handle' for subsequent chemical conjugation to polymer nanoparticles. These notions resulted in the design of series 7 (Figure 2).

Eye drop treatment for glaucoma is notoriously hampered by the poor patient compliance and the progression of the disease and loss of vision<sup>12</sup>. Longer-acting intraocular drug delivery with polymeric systems could potentially solve this issue<sup>13</sup>. New compounds 7 were designed with this downstream goal in mind, since their structure could allow conjugation to the polymeric carriers via amide and other potentially biodegradable linkages. On the other hand, the inherent hydrophilicity of these compounds was seen as potentially beneficial as hydrophilic compounds, even when liberated from a polymer carrier, display slower clearance from the intraocular spaces<sup>14</sup>. Thus, even with similar on-target potencies, more hydrophilic drugs, once delivered to the intraocular space, are expected to have a lower clearance and would require smaller dose per day to exert their actions. Even taken alone, more hydrophilic hCA II inhibitors will have potential as traditional eye drop medications if they could be potentially delivered across the conjunctiva-sclera route to the ciliary body.

As cautioned earlier<sup>15</sup>, 'decorating' a more lipophilic potent hCA II inhibitor with outright hydrophilic moieties (i.e. moving from **5** to **7**) carries a potential risk of losing the desired hCA II potency. As one must bear in mind, the active site of carbonic

anhydrase has a very characteristic topology where a hydrophobic half of the protein surface is clearly delineated from the hydrophilic one  $^{16}$ . Thus, replacing a relatively hydrophopic groups in **5a–c** with a large hydrophilic carboxamide groups could, in principle, deprive **7** of desired affinity to hCA II. Despite these potential risks we set off to synthesise a set of compounds **7** for investigation of their carbonic anhydrase inhibitory potency *in vitro* and subsequent efficacy study of the best inhibitor intraocular pressure-lowering agents *in vivo*. Herein, we report the results of these studies.

#### **Results and discussion**

The key building block – ethyl 5–(4-sulfamoylphenyl)oxazole-2-carboxylate (8) – was synthesised in several straightforward steps from  $\alpha$ -aminoacetophenone hydrochloride as described previously  $^{10,15}$ . The electron-withdrawing influence of the sulphonamide group on the electrophilicity of the ester functionality in 8 turned out to be of advantage in subsequent synthesis of the target compounds **7a–e**. Indeed, on reaction requiring no additional activation, with 2.5-fold excess of mono-Boc-protected dibasic amines **9a–e** at r.t. in MeOH, respective amides **10a–e** were obtained and deprotected with TFA in 1,4-dioxane at 60 °C and purified chromatographically to give the target compounds **7a–e** (Scheme 1).

The inhibitory profile obtained for sulphonamides **7a–e** in a stopped-flow kinetics assay against human CA I, II, IV and XII is shown in Table 1. In addition to hCA II, the other three isoforms were selected to preliminarily gauge the off-target profile of the compounds intended to inhibit the target isoform. Moreover, inhibition profile against hCA IV and XII was thought to be of significance as these isoforms are also involved in the secretion of the intraocular liquor<sup>17</sup>.

To our delight, all four inhibitors **6a–d** preserved the potent inhibition profile against the target *h*CA II isoform (although their *h*CA II potency deteriorated somewhat compared to the less hydrophilic initial leads **5a–c**) and a clearly better *h*CA II selectivity profile compared to acetazolamide (**4**) employed as a

Scheme 1. Synthesis of hydrophilic sulphonamides 7a-e investigated in this work.

Table 1. Inhibitory activity of compounds 7a-e against the target (hCA II) as well as selected off-target (hCA I, IV and XII) isoforms.

		K <sub>i</sub> (nM) <sup>a</sup>			
Compound	Structure	hCA I	<i>h</i> CA II	hCA IV	<i>h</i> CA XII
7a	H <sub>2</sub> N S S N NH	4.0	0.069	21.6	3.9
7b	O NH H <sub>2</sub> N S S	56.8	0.92	23.7	8.9
7c	$H_2N$ $O$ $N$ $N$ $N$ $N$ $N$	31.3	0.41	30.6	5.7
7d	$0 \longrightarrow N \longrightarrow NH_2$ $H_2N \longrightarrow 0$	72.9	3.9	5.2	9.3
7e	$0 \qquad N \qquad NH_2$ $H_2N \qquad 0 \qquad N$	58.3	3.1	4.6	8.8
3 <sup>b</sup>		250	12	75	5.7

<sup>&</sup>lt;sup>a</sup>Mean from three different assays by stopped flow technique (errors were in the range of  $\pm$  5–10% of the reported values).

reference inhibitor. Interestingly, the replacement of the morpholine oxygen atom in 5c with hydrogen bond donating/accepting piperazine NH in compound 7a (a rather drastic change from the standpoint of potential molecular interactions which resulted in the change of the binding mode, vide infra) led to only a

three-fold drop in hCA II potency. This clearly makes compound stand out as the hydrophilic (and potentially 'bioconjugatable') follow-on to compound 5c. Of course, the ultimate efficacy profile of this inhibitor reducing the glaucomarelated intraocular pressure (IOP) would depend on a multitude

<sup>&</sup>lt;sup>b</sup>Sulfonamide inhibitor acetazolamide (AAZ) used as a reference pan-CA inhibitor in stopped flow CO<sub>2</sub> hydrase assay.

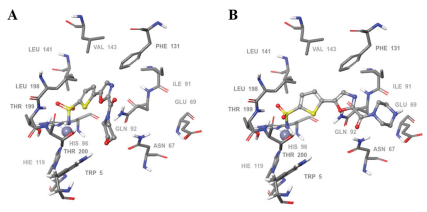


Figure 3. Binding poses of 5c (A) and 7a (B) in the hCAII active site.

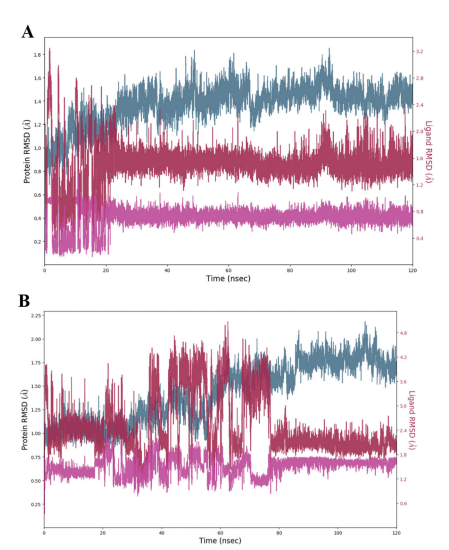


Figure 4. RMSD changes observed for the complexes 'acetazolamide – hCA II' (A) and 'compound 7a – hCA II' (B) during a 120 ns molecular dynamics simulation.

of factors among which permeability characteristics (intrinsically linked to a favourable set of molecular parameters) will be of significance.

In order to visualise the binding of the prototype compound  $\mathbf{5c}$  in comparison to the hydrophilic lead derivative  $\mathbf{7a}$  and to possibly understand the origins of the essentially preserved hCA II potency in case of the latter, we performed the docking of both ligands into the active site hCA II. In the case of both prototype molecule  $\mathbf{5c}$  and the advanced hydrophilic lead compound  $\mathbf{7a}$  the

thiophene sulphonamide moiety, predictably, acted as a zinc binding group displaying typical orientation which is well known from a wide range of crystallographic studies<sup>18</sup>. Specifically, the sulphonamide moiety interacted with the catalytic Zn<sup>2+</sup> ion as well as with Thr199. At the same time, the thiophene ring was oriented towards the hydrophobic pocket lined up with the residues Leu141, Val143, and Phe131. Furthermore, the 1,3-oxazole ring of the ligands was involved in interactions with Phe131 and formed a hydrogen bond with Gln92. Interestingly, we found the

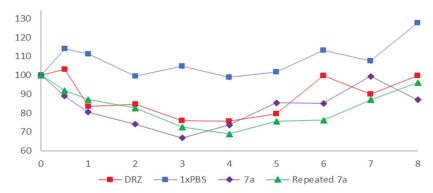


Figure 5. Percentage change in IOP (y axis) over time (x axis) after administration of compound 7a (two independent experiments), negative control phosphate buffered saline (PBS) and positive control dorzolamide (DRZ) in albino rabbits (n = 6).

Table 2. Chemical descriptors of carbonic anhydrase inhibitors 7a-e (calculated using ACDLabs 12.0).<sup>a</sup>

Compound	НВа	HBd	HBtot	LogP	MW	LogD <sub>8.0</sub>	PSA	logPSA
7a	3	11	-0.42	0.000861	342.39	0.38	155.15	2.191
7b	4	12	-1.46	-0.93	356.42	0.84	163.94	2.215
7c	4	12	-1.24	-0.68	356.42	0.42	169.14	2.228
7d	5	13	-1.11	-0.60	316.36	0.10	177.93	2.250
7e	5	13	-1.84	-1.30	330.38	0.36	177.93	2.250

<sup>a</sup>MW: molecular weight; HBa: hydrogen bond acceptors; HBd: hydrogen bond donors; HBtot: total amount of hydrogen bond formers; LogP: logarithmic value of partition coefficient; LogD<sub>7.4</sub>/LogD<sub>8.0</sub>: logarithmic value of distribution coefficient at pH 7.4/8.0; PSA: polar surface area; LogPSA: logarithmic value of polar surface area.

**Table 3.** Calculated permeability ( $P_{app}$ ) values of dorzolamide (1) and compounds **7a–e**.

Compound	Cornea (rabbit)		Cornea (porcine)		Conjunctiva (porcine)	
	$P_{\rm app}$ (cm/s)	% of 1	$P_{\rm app}$ (cm/s)	% of 1	$P_{\rm app}$ (cm/s)	% of 1
1	7.79E — 06	100	1.75E — 07	100	1.86E — 06	100
7a	9.68E — 07	12	1.71E — 07	98	1.83E — 06	98
7b	3.27E — 07	4	1.20E — 07	69	1.47E — 06	79
7c	3.76E — 07	5	1.18E — 07	67	1.45E — 06	78
7d	2.68E — 07	3	8.29E - 08	48	1.17E — 06	63
7e	1.68E — 07	2	8.29E - 08	48	1.17E — 06	63

 Table 4. Formulas for estimating permeability properties of carbonic anhydrase inhibitors.

	Formula	References
Corneal permeability of rabbit (cm/s)	LogPapp = -3.885 - 0.183(HBtot) + 0.277(logD7.4)	19
Corneal permeability of porcine (cm/s)	LogPapp = -4.6823 - 0.7670(logPSA) - 0.1346 (HBd) + 3.0024(Halogen ratio)	20
Conjunctival permeability of porcine (cm/s)	LogPapp = -4.1594 - 0.6121(logPSA) - 0.0792(HBd) + 3.2914(Halogen ratio)	21

LogPapp: logarithmic value of apparent permeability; HBtot: total amount of hydrogen bond formers; LogD7.4: logarithmic value of distribution coefficient at pH 7.4; LogPSA: logarithmic value of polar surface area; HBd: hydrogen bond donors; Halogen ratio: sum of all halogens divided by the sum of all heavy atoms excluding hydrogen.

morpholineamide moiety in the compound 5c was oriented towards the NH-groups of the Trp5 and Asn67. In contrast, the piperazine ring in compound 7a formed a salt bridge with Glu69. As it follows from this analysis, presumably, the ligand–protein interactions displayed by both morpholineamide moiety in 5a and piperazine amide substituent in 7a resulted in the favourable energy for the molecules' binding within the active site of hCA II and thus leading the potent inhibitory action of the compounds against the CA isoform (Figure 3).

In order to test the robustness of the docking poses identified, we performed 120 ns molecular dynamics simulation of ligand **7a** docked in the active site of hCA II in comparison with the clinically used (non-selective) hCA II inhibitor acetazolamide (**3**). The RMSD values of the protein backbone (blue), the ligand relative to hCA II (red) and the ligand relative to its original, pre-simulation docking pose (purple) were found to stabilise to fit the range of 1–3 Å (robust fit) within 23.36 ns for acetazolamide and within

77 ns for ligand **7a** (Figure 4). The longer relaxation time observed for **7a** has likely to do with the greater conformational flexibility of the piperazine carboxamide side chain which took longer to restore the network of critical hydrogen-bonding contacts. Overall, the molecular dynamics simulation demonstrated the robustness of the docking pose presented in Figure 3(B).

The intraocular pressure (IOP) lowering effect of newly developed hydrophilic hCAII inhibitor 7a was tested in normotensive New Zealand White rabbits<sup>19</sup>. The results are shown as percentage changes in Figure 5. Compound 7a (1% eye drop) (tested twice consecutively) showed a clear IOP lowering effect which was comparable to the effect produced by 1 (dorzolamide, administered as 2% eye drops).

For compounds **7a-e**, we have calculated a series of chemical descriptors (Table 2) from which critical ocular permeability parameters can be deduced. It is apparent, that all five compounds are distinctly hydrophilic.

The chemical descriptors presented in Table 2 allowed us to calculate the predicted corneal and conjunctival permeability values for compounds **7a–e** in comparison with dorzolamide (**1**) (Table 3). These calculations are based on the earlier formulas by Kidron et al.<sup>20</sup> and Ramsay et al.<sup>21,22</sup>. It is apparent that the conjunctival permeation route becomes a principal one for hydrophilic compounds **7a–e** in comparison with more lipophilic dorzolamide (**1**) (Table 4).

In summary, we have described next-generation 5-(sulfamoyl)thien-2-yl 1,3-oxazole carbonic anhydrase inhibitors endowed with a primary or secondary amine periphery. The compounds were designed with a dual goal of increasing compounds' hydrophilicity and provide a reactive 'handle' for potential conjugation to sustained-release nanoparticles. Increased hydrophilicity, while desirable for increased drug residence in the intraocular space could be generally viewed as an obstacle for corneal drug absorption. However, hydrophilic compounds may be efficiently absorbed via conjunctiva and thus have greater efficacy which may be expected if corneal absorption alone is considered. Out of the compounds described herein, the lead compound (7a) displayed a potent and selective inhibition of hCA II isoform, a glaucoma target and showed comparable efficacy as 1% eye drops in reducing the intraocular pressure in normotensive rabbit to that of clinically used 2% dorzolamide eye drops. This is despite the fact that the corneal permeability of these hydrophilic compounds was predicted to be significantly lower than that of dorzolamide. The data additionally support the concept of hydrophilic compounds permeating across the conjunctiva and sclera into the ciliary body.

### **Acknowledgements**

The authors are grateful to the Research Centre for Magnetic Resonance and the Centre for Chemical Analysis and Materials Research of Saint Petersburg State University Research Park for the analytical data. Annika Valtari and Marika Ruponen of University of Eastern Finland are thanked for providing the *in vivo* efficacy data associated with a previous publication <sup>18</sup>. Ocular Drug Development Laboratory was used for the *in vivo* studies.

#### **Disclosure statement**

No potential conflict of interest was reported by all author(s) except CTS. C. T. Supuran is Editor-in-Chief of the Journal of Enzyme Inhibition and Medicinal Chemistry. He was not involved in the assessment, peer review, or decision-making process of this paper. The authors have no relevant affiliations of financial involvement with any organisation or entity with a financial interest in or financial conflict with the subject matter or materials discussed in the manuscript. This includes employment, consultancies, honoraria, stock ownership or options, expert testimony, grants or patents received or pending, or royalties.

# **Funding**

This research was supported by the Ministry of Education of the Russian Federation [government contract 073-00077-21-02 'Development of an innovative glaucoma drug based on selective inhibition of carbonic anhydrase II', registry number  $730000\Phi.99.1.\overline{\mathrm{bB}}10\mathrm{A}A00006]$  and by Megagrant of the

Government of Russian Federation [075-15-2021-637] and Academy of Finland.

## **ORCID**

Stanislav Kalinin http://orcid.org/0000-0003-1194-0068
Alessio Nocentini http://orcid.org/0000-0003-3342-702X
Maxim Gureev http://orcid.org/0000-0002-0385-922X
Arto Urtti http://orcid.org/0000-0001-6064-3102
Claudiu T. Supuran http://orcid.org/0000-0003-4262-0323
Mikhail Krasavin http://orcid.org/0000-0002-0200-4772

## References

- Jonas JB, Aung T, Bourne RR, et al. Glaucoma. Lancet 2017; 390:2183–93.
- 2. Mincione F, Nocentini A, Supuran CT. Advances in the discovery of novel agents for the treatment of glaucoma. Expert Opin Drug Discov 2021;16:1209–25.
- 3. Guglielmi P, Carradori S, Campestre C, Poce G. Novel therapies for glaucoma: a patent review (2013–2019). Expert Opin Ther Pat 2019;29:769–80.
- Schmidl D, Schmetterer L, Garhöfer G, Popa-Cherecheanu A. Pharmacotherapy of glaucoma. J Ocul Pharmacol Ther 2015; 31:63–77.
- 5. Kanner E, Tsai JC. Glaucoma medications. Drugs Aging 2006; 23:321–32.
- 6. a) Maren TH. Carbonic anhydrase: chemistry, physiology, and inhibition. Physiol Rev 1967;47:595–781. b) Supuran CT, Casini A, Scozzafava A, Development of sulfonamide carbonic anhydrase inhibitors. In: Supuran CT, Scozzafava A, Conway J, eds. Carbonic anhydrase Its inhibitors and activators. Boca Raton (FL): CRC Press; 2004:67–147. c) Scozzafava A, Menabuoni L, Mincione F, et al. Carbonic anhydrase inhibitors: perfluoroalkyl/aryl-substituted derivatives of aromatic/heterocyclic sulfonamides as topical intraocular pressure-lowering agents with prolonged duration of action. J Med Chem 2000;43:4542–51.
- 7. Maurice DM, Mishima S. Ocular pharmacology. In: Sears M, ed. Handbook of experimental pharmacology. Berlin-Heidelberg: Springer-Verlag; 1984:16–19.
- 8. Ahmed I, Patton TF. Importance of the noncorneal absorption route in topical ophthalmic drug delivery. Invest Ophthalmol Vis Sci 1985;26:584–7.
- Hämäläinen KM, Kananen K, Auriola S, et al. Characterization of paracellular and aqueous penetration routes in cornea, conjunctiva, and sclera. Invest Ophthalmol Vis Sci 1997;38: 627–34.
- Krasavin M, Korsakov M, Dorogov M, et al. Probing the 'bipolar' nature of the carbonic anhydrase active site: aromatic sulfonamides containing 1,3-oxazol-5-yl moiety as picomolar inhibitors of cytosolic CA I and CA II isoforms. Eur J Med Chem 2015;101:334–47.
- 11. Ferraroni M, Lucarini L, Masini E, et al. 1,3-Oxazole-based selective picomolar inhibitors of cytosolic human carbonic anhydrase II alleviate ocular hypertension in rabbits: potency is supported by X-ray crystallography of two leads. Bioorg Med Chem 2017;25:4560–5.

- Broadway DS, Cate H. Pharmacotherapy and adherence issues in treating elderly patients with glaucoma. Drugs Aging 2015;32:569-81.
- Yadav KS, Rajpurohit R, Sharma S. Glaucoma: current treatment and impact of advanced drug delivery systems. Life Sci 2019;221:362-91.
- Del Amo EM, Rimpelä A-K, Heikkinen E, et al. Pharmacokinetic aspects of retinal drug delivery. \*Prog Retin Eye Res 2017;57:134-85.
- Kalinin S, Valtari A, Ruponen M, et al. Highly hydrophilic 1,3-oxazol-5-yl benzenesulfonamide inhibitors of carbonic anhydrase II for reduction of glaucoma-related intraocular pressure. Bioorg Med Chem 2019;27:115086.
- Nocentini A, Angeli A, Carta F, et al. Reconsidering anion inhibitors in the general context of drug design studies of modulators of activity of the classical enzyme carbonic anhydrase. J Enzyme Inhib Med Chem 2021;36:561-80.
- Scozzafava A, Supuran CT. Glaucoma and the applications of carbonic anhydrase inhibitors. Subcell Biochem 2014;75: 349-59.

- Lomelino CL, Andring JT, McKenna R. Crystallography and its impact on carbonic anhydrase research. Int J Med Chem 2018:2018:9419521.
- 19. Pilipenko I, Korzhikov-Vlakh V, Valtari A, et al. Mucoadhesive properties of nanogels based on stimuli-sensitive glycosaminoglycan-graft-pNIPAAm copolymers. Int J Biol Macromol 2021;186:864-72.
- 20. Kidron H, Vellonen K-S, Amo E, et al. Prediction of the corneal permeability of drug-like compounds. Pharm Res 2010; 27:1398-407.
- 21. Ramsay E, Del Amo E, Toropainen E, et al. Corneal and conjunctival drug permeability: systematic comparison and pharmacokinetic impact in the eye. Eur J Pharm Sci 2018; 119:83-9.
- 22. Ramsay E, Ruponen M, Picardat T, et al. Impact of chemical structure on conjunctival drug permeability: adopting porcine conjunctiva and cassette dosing for construction of in silico model. J Pharm Sci 2017;106: 2463-71.