


Web of Science | InCites | Journal Citation Reports | Essential Science Indicators | EndNote | Sign In ▾ | Help | English ▾

Web of Science 

Search | Search Results | My Tools ▾ | Search History | Marked List

 [Look Up Full Text](#)



Save to EndNote online ▾

[Add to Marked List](#)

◀ 1 of 1 ▶

Synthesis of new 3-(2-mercapto-4-oxo-4H-quinazolin-3-yl)-benzenesulfonamides with strong inhibition properties against the tumor associated carbonic anhydrases IX and XII

By: [Bozdag, M](#) (Bozdag, Murat)^[1,4]; [Alafeefy, AM](#) (Alafeefy, Ahmed Mahmoud)^[2]; [Altamimi, AM](#) (Altamimi, Abdul Malik); [Carta, F](#) (Carta, Fabrizio)^[1]; [Supuran, CT](#) (Supuran, Claudiu T.)^[1]; [Vullo, D](#) (Vullo, Daniela)^[4]

BIOORGANIC & MEDICINAL CHEMISTRY

Volume: 25 Issue: 10 Pages: 2782-2788

DOI: 10.1016/j.bmc.2017.03.054

Published: MAY 15 2017

[View Journal Impact](#)

Abstract

We report a series of novel metanilamide-based derivatives 3a-q bearing the 2-mercapto-4-oxo-4H-quinazolin-3-yl moiety as tail. All compounds were synthesized by means of straightforward condensation procedures and were investigated in vitro for their inhibition potency against the human (h) carbonic anhydrase (CA; EC 4.2.1.1.1) isoforms I, II, IX and XII. Among all compounds tested the 6-iodo 3g and the 7-fluoro 3i derivatives were the most potent inhibitors against the tumor associated CA IX and XII isoform (K(1)s 1.5 and 2.7 nM respectively for the hCA IX and K1s 0.57 and 1.9 nM respectively for the hCA XII).

The kinetic data reported here strongly support compounds of this type for their future development as radiotracers in tumor pathologies which are strictly dependent on the enzymatic activity of the hCA IX and XII isoforms. (C) 2017 Elsevier Ltd. All rights reserved.

Keywords

Author Keywords: Carbonic anhydrase inhibitors (CAIs); Quinazolines; Tumors; Imaging

KeyWords Plus: RAY CRYSTALLOGRAPHIC INVESTIGATIONS; ISOFORM-SELECTIVE INHIBITORS; THERAPEUTIC APPLICATIONS; ANTIGLAUCOMA ACTION; PHENOLIC-COMPOUNDS; IN-VIVO; COUMARINS; SULFONAMIDES; DERIVATIVES; POTENT

Author Information

Reprint Address: Supuran, CT (reprint author)

+ Univ Firenze, Dipartimento Neurofarba, Sez Sci Farmaceut, Polo Sci, Via U Schiff 6, I-50019 Florence, Italy.

Reprint Address: Vullo, D (reprint author)

+ Univ Firenze, Dipartimento Chim, Lab Chim Bioinorgan, Rm 188, Via Lastruccia 3, Florence, Italy.

Addresses:

+ [1] Univ Firenze, Dipartimento Neurofarba, Sez Sci Farmaceut, Polo Sci, Via U Schiff 6, I-50019 Florence, Italy

+ [2] Int Islamic Univ, Kulliyah Sci, Chem Dept, Kuantan, Malaysia

+ [3] Prince Sattam Bin Abdulaziz Univ, Coll Pharm, Pharmaceut Chem Dept, Al Kharj, Saudi Arabia

+ [4] Univ Firenze, Dipartimento Chim, Lab Chim Bioinorgan, Rm 188, Via Lastruccia 3, Florence, Italy

Citation Network

0 Times Cited

[65 Cited References](#)

[View Related Records](#)

 [Create Citation Alert](#)

(data from Web of Science Core Collection)

All Times Cited Counts

0 in All Databases

0 in Web of Science Core Collection

0 in BIOSIS Citation Index

0 in Chinese Science Citation Database

0 in Data Citation Index

0 in Russian Science Citation Index

0 in SciELO Citation Index

Usage Count

Last 180 Days: 2

Since 2013: 2

[Learn more](#)

This record is from:

Web of Science Core Collection
- Science Citation Index Expanded

Suggest a correction

If you would like to improve the quality of the data in this record, please [suggest a correction](#).