## Silvia Salerno

## List of Publications by Year in descending order

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331538 377752 1,224 49 21 34 citations h-index g-index papers 51 51 51 1671 docs citations times ranked citing authors all docs

#	Article	lF	Citations
1	Cancer Immunotherapy: An Overview on Small Molecules as Inhibitors of the Immune Checkpoint PD-1/PD-L1 (2015-2021). Mini-Reviews in Medicinal Chemistry, 2022, 22, .	1.1	3
2	Carbonic Anhydrase Activators for Neurodegeneration: An Overview. Molecules, 2022, 27, 2544.	1.7	17
3	Multiple Topoisomerase I (Topol), Topoisomerase II (Topoll) and Tyrosyl-DNA Phosphodiesterase (TDP) inhibitors in the development of anticancer drugs. European Journal of Pharmaceutical Sciences, 2021, 156, 105594.	1.9	31
4	Novel positive allosteric modulators of A <sub>2B</sub> adenosine receptor acting as bone mineralisation promoters. Journal of Enzyme Inhibition and Medicinal Chemistry, 2021, 36, 287-295.	2.5	12
5	An update into the medicinal chemistry of translocator protein (TSPO) ligands. European Journal of Medicinal Chemistry, 2021, 209, 112924.	2.6	31
6	Carbonic anhydrase activation profile of indole-based derivatives. Journal of Enzyme Inhibition and Medicinal Chemistry, 2021, 36, 1783-1797.	2.5	3
7	Tetrahydroquinazole-based secondary sulphonamides as carbonic anhydrase inhibitors: synthesis, biological evaluation against isoforms I, II, IV, and IX, and computational studies. Journal of Enzyme Inhibition and Medicinal Chemistry, 2021, 36, 1874-1883.	2.5	4
8	The Alpha Keto Amide Moiety as a Privileged Motif in Medicinal Chemistry: Current Insights and Emerging Opportunities. Journal of Medicinal Chemistry, 2021, 64, 3508-3545.	2.9	51
9	Inhibition studies on carbonic anhydrase isoforms I, II, IV and IX with N-arylsubstituted secondary sulfonamides featuring a bicyclic tetrahydroindazole scaffold. European Journal of Medicinal Chemistry, 2021, 220, 113490.	2.6	9
10	Drug Repurposing Meets DNA Independent Pathways: Targeting Alternative Substrates for Anticancer Therapy. Current Topics in Medicinal Chemistry, 2021, 21, 2767-2770.	1.0	0
11	Enriching the Arsenal of Pharmacological Tools against MICAL2. Molecules, 2021, 26, 7519.	1.7	1
12	Targeting the KRAS oncogene: Synthesis, physicochemical and biological evaluation of novel G-Quadruplex DNA binders. European Journal of Pharmaceutical Sciences, 2020, 149, 105337.	1.9	15
13	Discovery of Pyrido[3′,2′:5,6]thiopyrano[4,3- <i>d</i> ]pyrimidine-Based Antiproliferative Multikinase Inhibitors. ACS Medicinal Chemistry Letters, 2019, 10, 457-462.	1.3	3
14	Benzothiopyranoindole- and pyridothiopyranoindole-based antiproliferative agents targeting topoisomerases. European Journal of Medicinal Chemistry, 2019, 165, 46-58.	2.6	5
15	New insights in the structure-activity relationships of 2-phenylamino-substituted benzothiopyrano[4,3-d]pyrimidines as kinase inhibitors. European Journal of Medicinal Chemistry, 2018, 150, 446-456.	2.6	7
16	Novel fluorescent triazinobenzimidazole derivatives as probes for labelling human A1 and A2B adenosine receptor subtypes. Bioorganic and Medicinal Chemistry, 2018, 26, 5885-5895.	1.4	6
17	4-Substituted Benzenesulfonamides Incorporating Bi/Tricyclic Moieties Act as Potent and Isoform-Selective Carbonic Anhydrase II/IX Inhibitors. Journal of Medicinal Chemistry, 2018, 61, 5765-5770.	2.9	18
18	1,2-Benzisothiazole Derivatives Bearing 4-, 5-, or 6-Alkyl/arylcarboxamide Moieties Inhibit Carbonic Anhydrase Isoform IX (CAIX) and Cell Proliferation under Hypoxic Conditions. Journal of Medicinal Chemistry, 2016, 59, 6547-6552.	2.9	20

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19	Sulfonamides incorporating heteropolycyclic scaffolds show potent inhibitory action against carbonic anhydrase isoforms I, II, IX and XII. Bioorganic and Medicinal Chemistry, 2016, 24, 921-927.	1.4	18
20	Deepening the Topology of the Translocator Protein Binding Site by Novel <i>N</i> , <i>N</i> ,Dialkyl-2-arylindol-3-ylglyoxylamides. Journal of Medicinal Chemistry, 2015, 58, 6081-6092.	2.9	31
21	Investigation of new 2-aryl substituted Benzothiopyrano[4,3-d]pyrimidines as kinase inhibitors targeting vascular endothelial growth factor receptor 2. European Journal of Medicinal Chemistry, 2015, 103, 29-43.	2.6	17
22	Structure-Based Optimization of Tyrosine Kinase Inhibitor <b>CLM3</b> . Design, Synthesis, Functional Evaluation, and Molecular Modeling Studies Journal of Medicinal Chemistry, 2014, 57, 1225-1235.	2.9	18
23	Phenylpyrazolo[1,5- <i>a</i> )quinazolin-5(4 <i>H</i> )-one: A Suitable Scaffold for the Development of Noncamptothecin Topoisomerase I (Top1) Inhibitors. Journal of Medicinal Chemistry, 2013, 56, 7458-7462.	2.9	43
24	Arylsulfonamide inhibitors of aggrecanases as potential therapeutic agents for osteoarthritis: Synthesis and biological evaluation. European Journal of Medicinal Chemistry, 2013, 62, 379-394.	2.6	38
25	Medicinal Chemistry of Indolylglyoxylamide TSPO High Affinity Ligands with Anxiolytic-Like Effects. Current Topics in Medicinal Chemistry, 2012, 12, 333-351.	1.0	6
26	Medicinal Chemistry of Indolylglyoxylamide GABAA/BzR High Affinity Ligands: Identification of Novel Anxiolytic/Non Sedative Agents. Current Topics in Medicinal Chemistry, 2012, 12, 286-311.	1.0	8
27	Benzofuroxane Derivatives as Multi-Effective Agents for the Treatment of Cardiovascular Diabetic Complications. Synthesis, Functional Evaluation, and Molecular Modeling Studies. Journal of Medicinal Chemistry, 2012, 55, 10523-10531.	2.9	24
28	Tricyclic Sulfonamides Incorporating Benzothiopyrano [4,3-c]pyrazole and Pyridothiopyrano [4,3-c]pyrazole Effectively Inhibit $\hat{l}_{\pm}$ - and $\hat{l}^2$ -Carbonic Anhydrase: X-ray Crystallography and Solution Investigations on 15 Isoforms. Journal of Medicinal Chemistry, 2012, 55, 9619-9629.	2.9	35
29	3-Aryl-[1,2,4]triazino[4,3- <i>a</i> )benzimidazol-4(10 <i>H</i> )-one: A Novel Template for the Design of Highly Selective A <sub>2B</sub> Adenosine Receptor Antagonists. Journal of Medicinal Chemistry, 2012, 55, 1490-1499.	2.9	28
30	Novel Irreversible Fluorescent Probes Targeting the 18 kDa Translocator Protein: Synthesis and Biological Characterization. Journal of Medicinal Chemistry, 2010, 53, 4085-4093.	2.9	25
31	Novel <i>N</i> <sup>2</sup> -Substituted Pyrazolo[3,4- <i>d</i> ]pyrimidine Adenosine A <sub>3</sub> Receptor Antagonists: Inhibition of A <sub>3</sub> -Mediated Human Glioblastoma Cell Proliferation <sup>â€</sup> . Journal of Medicinal Chemistry, 2010, 53, 3954-3963.	2.9	50
32	Identification of Anxiolytic/Nonsedative Agents among Indol-3-ylglyoxylamides Acting as Functionally Selective Agonists at the $\hat{I}^3$ -Aminobutyric Acid-A (GABA $<$ sub $>$ A $<$ sub $>$ ) $\hat{I}_{\pm}<$ sub $>$ 2 $<$ /sub $>$ Benzodiazepine Receptor. Journal of Medicinal Chemistry, 2009, 52, 3723-3734.	2.9	27
33	Benzothiopyranoindole-Based Antiproliferative Agents: Synthesis, Cytotoxicity, Nucleic Acids Interaction, and Topoisomerases Inhibition Properties. Journal of Medicinal Chemistry, 2009, 52, 5429-5441.	2.9	30
34	Exploiting the Pyrazolo[3,4-d]pyrimidin-4-one Ring System as a Useful Template To Obtain Potent Adenosine Deaminase Inhibitors. Journal of Medicinal Chemistry, 2009, 52, 1681-1692.	2.9	44
35	Synthesis andin vitroantiproliferative activity of new substituted benzo[3′,2′:5,6]thiopyrano[4,3-d]pyrimidines. Journal of Heterocyclic Chemistry, 2008, 45, 745-749.	1.4	14
36	Acetic Acid Aldose Reductase Inhibitors Bearing a Five-Membered Heterocyclic Core with Potent Topical Activity in a Visual Impairment Rat Model. Journal of Medicinal Chemistry, 2008, 51, 3182-3193.	2.9	47

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37	Anxiolytic-like Effects of <i>N</i> , <i>N</i> -Dialkyl-2-phenylindol-3-ylglyoxylamides by Modulation of Translocator Protein Promoting Neurosteroid Biosynthesis. Journal of Medicinal Chemistry, 2008, 51, 5798-5806.	2.9	80
38	Pyrido[1,2- <i>a</i> )]pyrimidin-4-one Derivatives as a Novel Class of Selective Aldose Reductase Inhibitors Exhibiting Antioxidant Activity. Journal of Medicinal Chemistry, 2007, 50, 4917-4927.	2.9	130
39	5-Amino-2-phenyl[1,2,3]triazolo[1,2-a][1,2,4]benzotriazin-1-one:  A Versatile Scaffold To Obtain Potent and Selective A3 Adenosine Receptor Antagonists. Journal of Medicinal Chemistry, 2007, 50, 5676-5684.	2.9	22
40	Mitochondrial permeability transition induced by novel pyridothiopyranopyrimidine derivatives: Potential new antimitochondrial antitumour agents. Biochemical Pharmacology, 2006, 72, 1657-1667.	2.0	16
41	Spirohydantoin derivatives of thiopyrano [2,3-b] pyridin-4(4H)-one as potent in vitro and in vivo aldose reductase inhibitors. Bioorganic and Medicinal Chemistry, 2005, 13, 491-499.	1.4	34
42	High Affinity Central Benzodiazepine Receptor Ligands:Â Synthesis and Biological Evaluation of a Series of Phenyltriazolobenzotriazindione Derivatives. Journal of Medicinal Chemistry, 2005, 48, 2936-2943.	2.9	9
43	Synthesis of novel 1,4-dihydropyrido[3′,2′:5,6]thiopyrano[4,3-c]-pyrazoles and 5H-pyrido[3′,2′:5,6]thiopyrano[4,3-d]pyrimidines as potential antiproliferative agents. Journal of Heterocyclic Chemistry, 2003, 40, 783-788.	1.4	14
44	Synthesis of novel pyrido $[3\hat{a}\in^2,2\hat{a}\in^2:5,6]$ thiopyrano $[3,2-b]$ indol-5(6H)-ones and 6H-pyrido $[3\hat{a}\in^2,2\hat{a}\in^2:5,6]$ thiopyrano $[4,3-b]$ quinolines, two new heterocyclic ring systems. Journal of Heterocyclic Chemistry, 2002, 39, 1001-1006.	1.4	19
45	Facile synthesis of 3â€substituted [1,2,4]triazino[3,4â€ <i>f</i> ]purineâ€4,6,8â€trione derivatives. Journal of Heterocyclic Chemistry, 2001, 38, 607-612.	1.4	5
46	Synthesis, in vitro antiproliferative activity and DNA-interaction of benzimidazoquinazoline derivatives as potential anti-tumor agents. Il Farmaco, 2001, 56, 159-167.	0.9	70
47	Synthesis of novel 5 <i>H</i> , 11 <i>H</i> ê€pyrido[2′,3′:2,3]thiopyrano[4,3â€ <i>b</i> ]â€indoles by fischer cyclization. Journal of Heterocyclic Chemistry, 2000, 37, 379-382.	â€jndole 1.4	21
48	Synthesis of purinobenzodiazepine and purinobenzotriazocine derivatives, two new heterocyclic ring systems. Journal of Heterocyclic Chemistry, 1999, 36, 639-642.	1.4	8
49	Synthesis, DNA binding and in vitro antiproliferative activity of purinoquinazoline, pyridopyrimidopurine and pyridopyrimidobenzimidazole derivatives as potential antitumor agents. European Journal of Medicinal Chemistry, 1998, 33, 685-696.	2.6	57