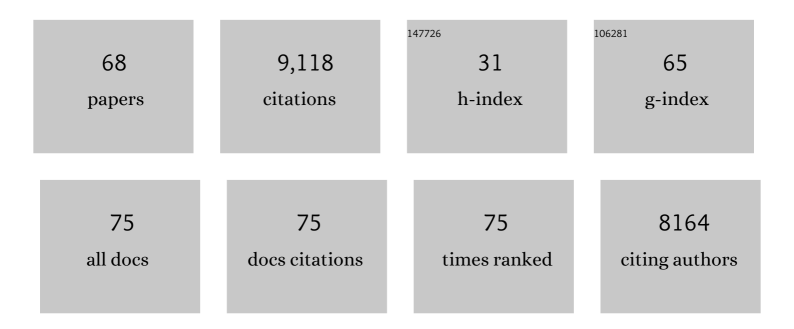
Viviana Consonni

List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/599612/publications.pdf Version: 2024-02-01



#	Article	IF	CITATIONS
1	Predicting molecular activity on nuclear receptors by multitask neural networks. Journal of Chemometrics, 2022, 36, e3325.	0.7	13
2	Expanding Antineoplastic Drugs Surface Monitoring Profiles: Enhancing of Zwitterionic Hydrophilic Interaction Methods. Separations, 2022, 9, 34.	1.1	1
3	Application of DNA mini-barcoding and infrared spectroscopy for the authentication of the Italian product "bottarga― LWT - Food Science and Technology, 2021, 139, 110603.	2.5	9
4	CATMoS: Collaborative Acute Toxicity Modeling Suite. Environmental Health Perspectives, 2021, 129, 47013.	2.8	63
5	A MATLAB toolbox for multivariate regression coupled with variable selection. Chemometrics and Intelligent Laboratory Systems, 2021, 213, 104313.	1.8	22
6	Parsimonious Optimization of Multitask Neural Network Hyperparameters. Molecules, 2021, 26, 7254.	1.7	10
7	Chemometrics for QSAR Modeling. , 2020, , 599-634.		6
8	Deep Ranking Analysis by Power Eigenvectors (DRAPE): A polypharmacology case study. Chemometrics and Intelligent Laboratory Systems, 2020, 203, 104001.	1.8	2
9	CoMPARA: Collaborative Modeling Project for Androgen Receptor Activity. Environmental Health Perspectives, 2020, 128, 27002.	2.8	120
10	Consensus versus Individual QSARs in Classification: Comparison on a Large-Scale Case Study. Journal of Chemical Information and Modeling, 2020, 60, 1215-1223.	2.5	26
11	Geographical identification of Chianti red wine based on ICP-MS element composition. Food Chemistry, 2020, 315, 126248.	4.2	37
12	On the Misleading Use of for QSAR Model Comparison. Molecular Informatics, 2019, 38, e1800029.	1.4	31
13	Structural alerts for the identification of bioaccumulative compounds. Integrated Environmental Assessment and Management, 2019, 15, 19-28.	1.6	13
14	Machine Learning Consensus To Predict the Binding to the Androgen Receptor within the CoMPARA Project. Journal of Chemical Information and Modeling, 2019, 59, 1839-1848.	2.5	36
15	Integrated QSAR Models to Predict Acute Oral Systemic Toxicity. Molecular Informatics, 2019, 38, e1800124.	1.4	30
16	Acceptableâ€byâ€design QSARs to predict the dietary biomagnification of organic chemicals in fish. Integrated Environmental Assessment and Management, 2019, 15, 51-63.	1.6	7
17	Mapping of Activity through Dichotomic Scores (MADS): A new chemoinformatic approach to detect activityâ€rich structural regions. Journal of Chemometrics, 2018, 32, e2994.	0.7	1
18	Impact of Molecular Descriptors on Computational Models. Methods in Molecular Biology, 2018, 1825, 171-209.	0.4	34

VIVIANA CONSONNI

#	Article	IF	CITATIONS
19	Detecting the bioaccumulation patterns of chemicals through data-driven approaches. Chemosphere, 2018, 208, 273-284.	4.2	14
20	Molecular Descriptors for Structure–Activity Applications: A Hands-On Approach. Methods in Molecular Biology, 2018, 1800, 3-53.	0.4	28
21	Scaffold hopping from natural products to synthetic mimetics by holistic molecular similarity. Communications Chemistry, 2018, 1, .	2.0	42
22	Molecular Descriptors. , 2017, , 2065-2093.		30
23	Matrixâ€based Molecular Descriptors for Prospective Virtual Compound Screening. Molecular Informatics, 2017, 36, 1600091.	1.4	18
24	A QSTR-Based Expert System to Predict Sweetness of Molecules. Frontiers in Chemistry, 2017, 5, 53.	1.8	41
25	In Silico Prediction of Cytochrome P450-Drug Interaction: QSARs for CYP3A4 and CYP2C9. International Journal of Molecular Sciences, 2016, 17, 914.	1.8	50
26	Expert QSAR system for predicting the bioconcentration factor under the REACH regulation. Environmental Research, 2016, 148, 507-512.	3.7	24
27	Mixtures, metabolites, ionic liquids: a new measure to evaluate similarity between complex chemical systems. Journal of Cheminformatics, 2016, 8, 49.	2.8	10
28	A new concept of higher-order similarity and the role of distance/similarity measures in local classification methods. Chemometrics and Intelligent Laboratory Systems, 2016, 157, 50-57.	1.8	27
29	Quantitative structure–activity relationships to predict sweet and non-sweet tastes. Theoretical Chemistry Accounts, 2016, 135, 1.	0.5	27
30	Investigating the mechanisms of bioconcentration through QSAR classification trees. Environment International, 2016, 88, 198-205.	4.8	32
31	Molecular Descriptors. , 2016, , 1-29.		13
32	Impact of medium-distance pollution sources in a Galician suburban site (NW Iberian peninsula). Science of the Total Environment, 2015, 512-513, 114-124.	3.9	11
33	QSAR models for bioconcentration: Is the increase in the complexity justified by more accurate predictions?. Chemosphere, 2015, 127, 171-179.	4.2	41
34	N3 and BNN: Two New Similarity Based Classification Methods in Comparison with Other Classifiers. Journal of Chemical Information and Modeling, 2015, 55, 2365-2374.	2.5	32
35	How to weight Hasse matrices and reduce incomparabilities. Chemometrics and Intelligent Laboratory Systems, 2015, 147, 95-104.	1.8	12
36	Prediction of Acute Aquatic Toxicity toward <i>Daphnia Magna</i> by using the GA- <i>k</i> NN Method. ATLA Alternatives To Laboratory Animals, 2014, 42, 31-41.	0.7	59

VIVIANA CONSONNI

#	Article	IF	CITATIONS
37	Towards Global QSAR Model Building for Acute Toxicity: Munro Database Case Study. International Journal of Molecular Sciences, 2014, 15, 18162-18174.	1.8	36
38	QSPR STUDY OF RHEOLOGICAL AND MECHANICAL PROPERTIES OF CHLOROPRENE RUBBER ACCELERATORS. Rubber Chemistry and Technology, 2014, 87, 219-238.	0.6	7
39	QSAR Modeling: Where Have You Been? Where Are You Going To?. Journal of Medicinal Chemistry, 2014, 57, 4977-5010.	2.9	1,401
40	A novel variable reduction method adapted from space-filling designs. Chemometrics and Intelligent Laboratory Systems, 2014, 136, 147-154.	1.8	60
41	Assessing the Validity of QSARs for Ready Biodegradability of Chemicals: An Applicability Domain Perspective. Current Computer-Aided Drug Design, 2014, 10, 137-147.	0.8	24
42	Classification tools in chemistry. Part 1: linear models. PLS-DA. Analytical Methods, 2013, 5, 3790.	1.3	893
43	Locally centred Mahalanobis distance: A new distance measure with salient features towards outlier detection. Analytica Chimica Acta, 2013, 787, 1-9.	2.6	60
44	Quantitative Structure–Activity Relationship Models for Ready Biodegradability of Chemicals. Journal of Chemical Information and Modeling, 2013, 53, 867-878.	2.5	160
45	Defining a novel k-nearest neighbours approach to assess the applicability domain of a QSAR model for reliable predictions. Journal of Cheminformatics, 2013, 5, 27.	2.8	65
46	Comparison of Different Approaches to Define the Applicability Domain of QSAR Models. Molecules, 2012, 17, 4791-4810.	1.7	370
47	Similarity Coefficients for Binary Chemoinformatics Data: Overview and Extended Comparison Using Simulated and Real Data Sets. Journal of Chemical Information and Modeling, 2012, 52, 2884-2901.	2.5	155
48	Assessing bioaccumulation of polybrominated diphenyl ethers for aquatic species by QSAR modeling. Chemosphere, 2012, 89, 433-444.	4.2	28
49	Relationships between apple texture and rheological parameters by means of multivariate analysis. Chemometrics and Intelligent Laboratory Systems, 2012, 111, 28-33.	1.8	22
50	Genetic Algorithms for architecture optimisation of Counter-Propagation Artificial Neural Networks. Chemometrics and Intelligent Laboratory Systems, 2011, 105, 56-64.	1.8	66
51	Structure –Activity Relationships by Autocorrelation Descriptors and Genetic Algorithms. , 2011, , 60-94.		4
52	Canonical Measure of Correlation (CMC) and Canonical Measure of Distance (CMD) between sets of data. Part 3. Variable selection in classification. Analytica Chimica Acta, 2010, 657, 116-122.	2.6	7
53	Evaluation of model predictive ability by external validation techniques. Journal of Chemometrics, 2010, 24, 194-201.	0.7	290
54	Molecular Descriptors. Challenges and Advances in Computational Chemistry and Physics, 2010, , 29-102.	0.6	62

4

VIVIANA CONSONNI

#	Article	IF	CITATIONS
55	The Kohonen and CP-ANN toolbox: A collection of MATLAB modules for Self Organizing Maps and Counterpropagation Artificial Neural Networks. Chemometrics and Intelligent Laboratory Systems, 2009, 98, 115-122.	1.8	111
56	Canonical Measure of Correlation (CMC) and Canonical Measure of Distance (CMD) between sets of data. Analytica Chimica Acta, 2009, 648, 52-59.	2.6	7
57	Comments on the Definition of the <i>Q</i> ² Parameter for QSAR Validation. Journal of Chemical Information and Modeling, 2009, 49, 1669-1678.	2.5	483
58	Classification of multiway analytical data based on MOLMAP approach. Analytica Chimica Acta, 2007, 605, 134-146.	2.6	17
59	New QSAR Modelling Approach Based on Ranking Models by Genetic Algorithms - Variable Subset Selection (GA-VSS). , 2006, , 181-217.		8
60	A distance measure between models: a tool for similarity/diversity analysis of model populations. Chemometrics and Intelligent Laboratory Systems, 2004, 70, 55-61.	1.8	28
61	Detecting "bad―regression models: multicriteria fitness functions in regression analysis. Analytica Chimica Acta, 2004, 515, 199-208.	2.6	156
62	MobyDigs: software for regression and classification models by genetic algorithms. Data Handling in Science and Technology, 2003, 23, 141-167.	3.1	34
63	Structure/Response Correlations and Similarity/Diversity Analysis by GETAWAY Descriptors. 1. Theory of the Novel 3D Molecular Descriptors. Journal of Chemical Information and Computer Sciences, 2002, 42, 682-692.	2.8	402
64	Structure/Response Correlations and Similarity/Diversity Analysis by GETAWAY Descriptors. 2. Application of the Novel 3D Molecular Descriptors to QSAR/QSPR Studies. Journal of Chemical Information and Computer Sciences, 2002, 42, 693-705.	2.8	278
65	QSAR study on the tropospheric degradation of organic compounds. Chemosphere, 1999, 38, 1371-1378.	4.2	67
66	Toward an in Vitro Test for the Diagnosis of Allergy to Penicillins. Synthesis, Characterization, and Use of β-Lactam and β-Lactam Metabolite Poly-l-lysines Which Recognize Human IgE Antibodies. Bioconjugate Chemistry, 1999, 10, 332-337.	1.8	7
67	High-performance size-exclusion chromatographic behaviour of substituted benzoylpoly-l-lysines by principal component analysis and molecular dynamics simulation. Journal of Chromatography A, 1998, 813, 255-265.	1.8	5
68	Qualitative consensus of QSAR ready biodegradability predictions. Toxicological and Environmental Chemistry, 0, , 1-24.	0.6	14