## Irina Vetter

## List of Publications by Year in descending order

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53794 49909 9,125 167 45 87 citations h-index g-index papers 177 177 177 11916 docs citations times ranked citing authors all docs

#	Article	IF	CITATIONS
1	Novel Neurotoxic Activity in Calliophis intestinalis Venom. Neurotoxicity Research, 2022, 40, 173-178.	2.7	3
2	Multitarget nociceptor sensitization by a promiscuous peptide from the venom of the King Baboon spider. Proceedings of the National Academy of Sciences of the United States of America, 2022, 119, .	7.1	7
3	Polygodial, a drimane sesquiterpenoid dialdehyde purified from <i>Drimys winteri</i> , inhibits voltage-gated sodium channels. Natural Product Research, 2022, 36, 6318-6323.	1.8	5
4	Towards a generic prototyping approach for therapeutically-relevant peptides and proteins in a cell-free translation system. Nature Communications, 2022, 13, 260.	12.8	5
5	Structural and functional insights into the inhibition of human voltage-gated sodium channels by $\hat{l}_{4}$ -conotoxin KIIIA disulfide isomers. Journal of Biological Chemistry, 2022, 298, 101728.	3.4	9
6	A peptide toxin in ant venom mimics vertebrate EGF-like hormones to cause long-lasting hypersensitivity in mammals. Proceedings of the National Academy of Sciences of the United States of America, 2022, 119, .	7.1	15
7	A kinase-dead $\langle i \rangle$ Csf1r $\langle j i \rangle$ mutation associated with adult-onset leukoencephalopathy has a dominant inhibitory impact on CSF1R signalling. Development (Cambridge), 2022, 149, .	2.5	9
8	Low potency inhibition of NaV1.7 by externally applied QX-314 via a depolarizing shift in the voltage-dependence of activation. European Journal of Pharmacology, 2022, , 175013.	<b>3.</b> 5	0
9	Neurotoxic and cytotoxic peptides underlie the painful stings of the tree nettle Urtica ferox. Journal of Biological Chemistry, 2022, 298, 102218.	3.4	5
10	Small cyclic sodium channel inhibitors. Biochemical Pharmacology, 2021, 183, 114291.	4.4	14
11	Convergent evolution of pain-inducing defensive venom components in spitting cobras. Science, 2021, 371, 386-390.	12.6	96
12	Subcutaneous ω-Conotoxins Alleviate Mechanical Pain in Rodent Models of Acute Peripheral Neuropathy. Marine Drugs, 2021, 19, 106.	4.6	13
13	The zebrafish <i>grime</i> mutant uncovers an evolutionarily conserved role for Tmem161b in the control of cardiac rhythm. Proceedings of the National Academy of Sciences of the United States of America, 2021, 118, .	7.1	12
14	Vincristine-induced peripheral neuropathy is driven by canonical NLRP3 activation and IL- $1\hat{l}^2$ release. Journal of Experimental Medicine, 2021, 218, .	8.5	29
15	Production, composition, and mode of action of the painful defensive venom produced by a limacodid caterpillar, <i>Doratifera vulnerans</i> . Proceedings of the National Academy of Sciences of the United States of America, 2021, 118, .	7.1	17
16	Venom chemistry underlying the painful stings of velvet ants (Hymenoptera: Mutillidae). Cellular and Molecular Life Sciences, 2021, 78, 5163-5177.	5 <b>.</b> 4	11
17	Improving the Gastrointestinal Stability of Linaclotide. Journal of Medicinal Chemistry, 2021, 64, 8384-8390.	6.4	14
18	The Allosteric Activation of $\hat{l}\pm7$ nAChR by $\hat{l}\pm$ -Conotoxin MrIC Is Modified by Mutations at the Vestibular Site. Toxins, 2021, 13, 555.	3.4	5

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19	Engineering of a Spider Peptide via Conserved Structure-Function Traits Optimizes Sodium Channel Inhibition In Vitro and Anti-Nociception In Vivo. Frontiers in Molecular Biosciences, 2021, 8, 742457.	3.5	5
20	A pain-causing and paralytic ant venom glycopeptide. IScience, 2021, 24, 103175.	4.1	7
21	Multipurpose peptides: The venoms of Amazonian stinging ants contain anthelmintic ponericins with diverse predatory and defensive activities. Biochemical Pharmacology, 2021, 192, 114693.	4.4	10
22	Evaluation of Efficient Non-reducing Enzymatic and Chemical Ligation Strategies for Complex Disulfide-Rich Peptides. Bioconjugate Chemistry, 2021, 32, 2407-2419.	3.6	4
23	The Tarantula Venom Peptide Eo1a Binds to the Domain II S3-S4 Extracellular Loop of Voltage-Gated Sodium Channel NaV1.8 to Enhance Activation. Frontiers in Pharmacology, 2021, 12, 789570.	3.5	4
24	Inflammatory and Neuropathic Gene Expression Signatures of Chemotherapy-Induced Neuropathy Induced by Vincristine, Cisplatin, and Oxaliplatin in C57BL/6J Mice. Journal of Pain, 2020, 21, 182-194.	1.4	38
25	Enzymatic Ligation of a Pore Blocker Toxin and a Gating Modifier Toxin: Creating Double-Knotted Peptides with Improved Sodium Channel NaV1.7 Inhibition. Bioconjugate Chemistry, 2020, 31, 64-73.	3.6	23
26	Pharmacology and therapeutic potential of venom peptides. Biochemical Pharmacology, 2020, 181, 114207.	4.4	4
27	Discovery, Pharmacological Characterisation and NMR Structure of the Novel Âμ-Conotoxin SxIIIC, a Potent and Irreversible NaV Channel Inhibitor. Biomedicines, 2020, 8, 391.	3.2	12
28	Recombinant production, bioconjugation and membrane binding studies of Pn3a, a selective NaV1.7 inhibitor. Biochemical Pharmacology, 2020, 181, 114148.	4.4	7
29	Australian funnel-web spiders evolved human-lethal Î'-hexatoxins for defense against vertebrate predators. Proceedings of the National Academy of Sciences of the United States of America, 2020, 117, 24920-24928.	7.1	32
30	The NLRP3 Inflammasome: Role and Therapeutic Potential in Pain Treatment. Frontiers in Physiology, 2020, 11, 1016.	2.8	40
31	Neurotoxic peptides from the venom of the giant Australian stinging tree. Science Advances, 2020, 6, .	10.3	16
32	An Integrated Proteomic and Transcriptomic Analysis Reveals the Venom Complexity of the Bullet Ant Paraponera clavata. Toxins, 2020, 12, 324.	3.4	18
33	Mutational analysis of ProTx-I and the novel venom peptide Pe1b provide insight into residues responsible for selective inhibition of the analgesic drug target NaV1.7. Biochemical Pharmacology, 2020, 181, 114080.	4.4	7
34	Animal toxins â€" Nature's evolutionary-refined toolkit for basic research and drug discovery. Biochemical Pharmacology, 2020, 181, 114096.	4.4	97
35	Characterization of Synthetic Tf2 as a NaV1.3 Selective Pharmacological Probe. Biomedicines, 2020, 8, 155.	3.2	8
36	NaV1.1 and NaV1.6 selective compounds reduce the behavior phenotype and epileptiform activity in a novel zebrafish model for Dravet Syndrome. PLoS ONE, 2020, 15, e0219106.	2.5	28

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37	Manipulation of a spider peptide toxin alters its affinity for lipid bilayers and potency and selectivity for voltage-gated sodium channel subtype 1.7. Journal of Biological Chemistry, 2020, 295, 5067-5080.	3.4	13
38	Addition of K22 Converts Spider Venom Peptide Pme2a from an Activator to an Inhibitor of NaV1.7. Biomedicines, 2020, 8, 37.	3.2	6
39	Pharmacological activity and NMR solution structure of the leech peptide HSTX-I. Biochemical Pharmacology, 2020, 181, 114082.	4.4	2
40	It Takes Two: Dimerization Is Essential for the Broad-Spectrum Predatory and Defensive Activities of the Venom Peptide Mp1a from the Jack Jumper Ant Myrmecia pilosula. Biomedicines, 2020, 8, 185.	3.2	12
41	Mapping the Molecular Surface of the Analgesic NaV1.7-Selective Peptide Pn3a Reveals Residues Essential for Membrane and Channel Interactions. ACS Pharmacology and Translational Science, 2020, 3, 535-546.	4.9	16
42	A New Selective Pharmacological Enhancer of the Orail Ca <sup>2+</sup> Channel Reveals Roles for Orail in Smooth and Skeletal Muscle Functions. ACS Pharmacology and Translational Science, 2020, 3, 135-147.	4.9	27
43	High-Throughput Fluorescence Assays for Ion Channels and GPCRs. Advances in Experimental Medicine and Biology, 2020, 1131, 27-72.	1.6	13
44	Characterisation of a Novel A-Superfamily Conotoxin. Biomedicines, 2020, 8, 128.	3.2	9
45	Transcriptomic characterisation of the optimised rat model of Walker 256 breast cancer cellâ€induced bone pain. Clinical and Experimental Pharmacology and Physiology, 2019, 46, 1201-1215.	1.9	2
46	Minocycline Prevents the Development of Mechanical Allodynia in Mouse Models of Vincristine-Induced Peripheral Neuropathy. Frontiers in Neuroscience, 2019, 13, 653.	2.8	30
47	Development of an <i>N</i> -Acyl Amino Acid That Selectively Inhibits the Glycine Transporter 2 To Produce Analgesia in a Rat Model of Chronic Pain. Journal of Medicinal Chemistry, 2019, 62, 2466-2484.	6.4	28
48	Na <sub>V</sub> 1.6 regulates excitability of mechanosensitive sensory neurons. Journal of Physiology, 2019, 597, 3751-3768.	2.9	31
49	Novel conorfamides from Conus austini venom modulate both nicotinic acetylcholine receptors and acid-sensing ion channels. Biochemical Pharmacology, 2019, 164, 342-348.	4.4	12
50	Development of a high-throughput fluorescent no-wash sodium influx assay. PLoS ONE, 2019, 14, e0213751.	2.5	13
51	Missiles of Mass Disruption: Composition and Glandular Origin of Venom Used as a Projectile Defensive Weapon by the Assassin Bug Platymeris rhadamanthus. Toxins, 2019, 11, 673.	3.4	16
52	Antiallodynic effects of the selective NaV1.7 inhibitor Pn3a in a mouse model of acute postsurgical pain: evidence for analgesic synergy with opioids and baclofen. Pain, 2019, 160, 1766-1780.	4.2	35
53	A Centipede Toxin Family Defines an Ancient Class of CSαβ Defensins. Structure, 2019, 27, 315-326.e7.	3.3	17
54	Assessment of the TRPM8 inhibitor AMTB in breast cancer cells and its identification as an inhibitor of voltage gated sodium channels. Life Sciences, 2018, 198, 128-135.	4.3	32

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55	The E15R Point Mutation in Scorpion Toxin Cn2 Uncouples Its Depressant and Excitatory Activities on Human Na <sub>V</sub> 1.6. Journal of Medicinal Chemistry, 2018, 61, 1730-1736.	6.4	9
56	Role of complement anaphylatoxin receptors in a mouse model of acute burn-induced pain. Molecular Immunology, 2018, 94, 68-74.	2.2	4
57	Burn Pain: A Systematic and Critical Review of Epidemiology, Pathophysiology, and Treatment. Pain Medicine, 2018, 19, 708-734.	1.9	61
58	Transcriptomics in pain research: insights from new and old technologies. Molecular Omics, 2018, 14, 389-404.	2.8	22
59	Buzz Kill: Function and Proteomic Composition of Venom from the Giant Assassin Fly Dolopus genitalis (Diptera: Asilidae). Toxins, 2018, 10, 456.	3.4	12
60	Pain-Causing Venom Peptides: Insights into Sensory Neuron Pharmacology. Toxins, 2018, 10, 15.	3.4	27
61	A comprehensive portrait of the venom of the giant red bull ant, <i>Myrmecia gulosa</i> , reveals a hyperdiverse hymenopteran toxin gene family. Science Advances, 2018, 4, eaau4640.	10.3	69
62	Novel analgesic ω-conotoxins from the vermivorous cone snail Conus moncuri provide new insights into the evolution of conopeptides. Scientific Reports, 2018, 8, 13397.	3.3	22
63	An SAR study of hydroxy-trifluoromethylpyrazolines as inhibitors of Orai1-mediated store operated Ca2+ entry in MDA-MB-231 breast cancer cells using a convenient Fluorescence Imaging Plate Reader assay. Bioorganic and Medicinal Chemistry, 2018, 26, 3406-3413.	3.0	9
64	The Somatostatin Receptor-4 Agonist J-2156 Alleviates Mechanical Hypersensitivity in a Rat Model of Breast Cancer Induced Bone Pain. Frontiers in Pharmacology, 2018, 9, 495.	3.5	17
65	Toxins in Neurobiology: New tools from old molecules. Neuroscience Letters, 2018, 679, 1-3.	2.1	1
66	Chemotactic responses of growing neurites to precisely controlled gradients of nerve growth factor. Scientific Data, 2018, 5, 180183.	5.3	8
67	Molecular Pharmacology of Pain-inducing Venom Peptides. Proceedings for Annual Meeting of the Japanese Pharmacological Society, 2018, WCP2018, SY66-4.	0.0	0
68	Pharmacological characterisation of the highly NaV1.7 selective spider venom peptide Pn3a. Scientific Reports, 2017, 7, 40883.	3.3	120
69	The long non-coding RNA NEAT1 is responsive to neuronal activity and is associated with hyperexcitability states. Scientific Reports, 2017, 7, 40127.	3.3	92
70	Role of the NLRP3 inflammasome in a model of acute burn-induced pain. Burns, 2017, 43, 304-309.	1.9	22
71	Multiple sodium channel isoforms mediate the pathological effects of Pacific ciguatoxin-1. Scientific Reports, 2017, 7, 42810.	3.3	67
72	Synthesis of Multivalent [Lys8]-Oxytocin Dendrimers that Inhibit Visceral Nociceptive Responses. Australian Journal of Chemistry, 2017, 70, 162.	0.9	9

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73	Sodium Channels and Venom Peptide Pharmacology. Advances in Pharmacology, 2017, 79, 67-116.	2.0	47
74	The pharmacology of voltage-gated sodium channel activators. Neuropharmacology, 2017, 127, 87-108.	4.1	57
75	Δâ€Myrtoxinâ€Mp1a is a Helical Heterodimer from the Venom of the Jack Jumper Ant that has Antimicrobial, Membraneâ€Disrupting, and Nociceptive Activities. Angewandte Chemie - International Edition, 2017, 56, 8495-8499.	13.8	28
76	Modulatory features of the novel spider toxin μâ€₹RTXâ€Df1a isolated from the venom of the spider <i>Davus fasciatus</i> . British Journal of Pharmacology, 2017, 174, 2528-2544.	5.4	46
77	Pharmacological screening technologies for venom peptide discovery. Neuropharmacology, 2017, 127, 4-19.	4.1	36
78	Lethal effects of an insecticidal spider venom peptide involve positive allosteric modulation of insect nicotinic acetylcholine receptors. Neuropharmacology, 2017, 127, 224-242.	4.1	16
79	The Evolution of Fangs, Venom, and Mimicry Systems in Blenny Fishes. Current Biology, 2017, 27, 1184-1191.	3.9	36
80	NaV1.7 as a pain target – From gene to pharmacology. , 2017, 172, 73-100.		104
81	Conotoxin Φâ€MiXXVIIA from the Superfamily G2 Employs a Novel Cysteine Framework that Mimics Granulin and Displays Antiâ€Apoptotic Activity. Angewandte Chemie, 2017, 129, 15169-15172.	2.0	3
82	Subtle modifications to oxytocin produce ligands that retain potency and improved selectivity across species. Science Signaling, 2017, $10$ , .	3.6	34
83	Evaluation of known and novel inhibitors of Orai1-mediated store operated Ca 2+ entry in MDA-MB-231 breast cancer cells using a Fluorescence Imaging Plate Reader assay. Bioorganic and Medicinal Chemistry, 2017, 25, 440-449.	3.0	17
84	Optimization and In Vivo Profiling of a Refined Rat Model of Walker 256 Breast Cancer Cell-Induced Bone Pain Using Behavioral, Radiological, Histological, Immunohistochemical and Pharmacological Methods. Frontiers in Pharmacology, 2017, 8, 442.	3.5	15
85	Pathophysiology of Chemotherapy-Induced Peripheral Neuropathy. Frontiers in Molecular Neuroscience, 2017, 10, 174.	2.9	403
86	Methods Used to Evaluate Pain Behaviors in Rodents. Frontiers in Molecular Neuroscience, 2017, 10, 284.	2.9	687
87	Discovery and mode of action of a novel analgesic $\hat{l}^2$ -toxin from the African spider Ceratogyrus darlingi. PLoS ONE, 2017, 12, e0182848.	2.5	22
88	Conotoxin Φâ€MiXXVIIA from the Superfamily G2 Employs a Novel Cysteine Framework that Mimics Granulin and Displays Antiâ€Apoptotic Activity. Angewandte Chemie - International Edition, 2017, 56, 14973-14976.	13.8	25
89	The structure, dynamics and selectivity profile of a NaV1.7 potency-optimised huwentoxin-IV variant. PLoS ONE, 2017, 12, e0173551.	2.5	33
90	The Snake with the Scorpion's Sting: Novel Three-Finger Toxin Sodium Channel Activators from the Venom of the Long-Glanded Blue Coral Snake (Calliophis bivirgatus). Toxins, 2016, 8, 303.	3.4	53

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91	Characterization of Three Venom Peptides from the Spitting Spider Scytodes thoracica. PLoS ONE, 2016, 11, e0156291.	2.5	6
92	The Walker 256 Breast Cancer Cell-Induced Bone Pain Model in Rats. Frontiers in Pharmacology, 2016, 7, 286.	3.5	38
93	Analgesic Effects of GpTx-1, PF-04856264 and CNV1014802 in a Mouse Model of NaV1.7-Mediated Pain. Toxins, 2016, 8, 78.	3.4	94
94	Neuronal cell lines as model dorsal root ganglion neurons. Molecular Pain, 2016, 12, 174480691664611.	2.1	81
95	Transcriptomic and behavioural characterisation of a mouse model of burn pain identify the cholecystokinin 2 receptor as an analgesic target. Molecular Pain, 2016, 12, 174480691666536.	2.1	58
96	Crotalphine desensitizes TRPA1 ion channels to alleviate inflammatory hyperalgesia. Pain, 2016, 157, 2504-2516.	4.2	31
97	Rational Design and Synthesis of a Novel Membrane Binding NaV1.8 Selective Inhibitor with in vivo Activity in Pain Models. Biophysical Journal, 2016, 110, 33a.	0.5	0
98	New Insight in Cold Pain: Role of Ion Channels, Modulation, and Clinical Perspectives. Journal of Neuroscience, 2016, 36, 11435-11439.	3.6	52
99	How hot is it Down Under?. Temperature, 2016, 3, 355-357.	3.0	0
100	Interaction of Tarantula Venom Peptide ProTx-II with Lipid Membranes Is a Prerequisite for Its Inhibition of Human Voltage-gated Sodium Channel NaV1.7. Journal of Biological Chemistry, 2016, 291, 17049-17065.	3.4	62
101	Development of a 1¼O-Conotoxin Analogue with Improved Lipid Membrane Interactions and Potency for the Analgesic Sodium Channel NaV1.8. Journal of Biological Chemistry, 2016, 291, 11829-11842.	3.4	37
102	Ciguatoxin and Ciguatera., 2016,, 71-92.		4
103	The thermal probe test: A novel behavioral assay to quantify thermal paw withdrawal thresholds in mice. Temperature, 2016, 3, 199-207.	3.0	45
104	Release of neuropeptides from a neuro-cutaneous co-culture model: A novel inÂvitro model for studying sensory effects of ciguatoxins. Toxicon, 2016, 116, 4-10.	1.6	17
105	Rapid Extraction and Identification of Maitotoxin and Ciguatoxin-Like Toxins from Caribbean and Pacific Gambierdiscus Using a New Functional Bioassay. PLoS ONE, 2016, 11, e0160006.	2.5	59
106	Transcriptome and proteome of <i>Conus planorbis</i> identify the nicotinic receptors as primary target for the defensive venom. Proteomics, 2015, 15, 4030-4040.	2.2	26
107	Vps26Bâ€retromer negatively regulates plasma membrane resensitization of PARâ€2. Cell Biology International, 2015, 39, 1299-1306.	3.0	7
108	Inhibition of N-Type Calcium Channels by Fluorophenoxyanilide Derivatives. Marine Drugs, 2015, 13, 2030-2045.	4.6	11

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109	(-)-Pentylsedinine, a New Alkaloid from the Leaves of Lobelia Tupa with Agonist Activity at Nicotinic Acetylcholine Receptor. Natural Product Communications, 2015, 10, 1934578X1501000.	0.5	3
110	Identification and Characterization of ProTx-III [ $\langle i \rangle \hat{l} \frac{1}{4} \langle i \rangle$ -TRTX-Tp1a], a New Voltage-Gated Sodium Channel Inhibitor from Venom of the Tarantula $\langle i \rangle$ Thrixopelma pruriens $\langle i \rangle$ . Molecular Pharmacology, 2015, 88, 291-303.	2.3	72
111	Therapeutic opportunities for targeting cold pain pathways. Biochemical Pharmacology, 2015, 93, 125-140.	4.4	33
112	A small-molecule inhibitor of the NLRP3 inflammasome for the treatment of inflammatory diseases. Nature Medicine, 2015, 21, 248-255.	30.7	1,967
113	CHAPTER 4. Venoms-Based Drug Discovery: Bioassays, Electrophysiology, High-Throughput Screens andÂTarget Identification. RSC Drug Discovery Series, 2015, , 97-128.	0.3	2
114	Seven novel modulators of the analgesic target <scp>Na<sub>V</sub></scp> 1.7 uncovered using a highâ€throughput venomâ€based discovery approach. British Journal of Pharmacology, 2015, 172, 2445-2458.	5.4	74
115	α-Conotoxin Dendrimers Have Enhanced Potency and Selectivity for Homomeric Nicotinic Acetylcholine Receptors. Journal of the American Chemical Society, 2015, 137, 3209-3212.	13.7	32
116	Î-Conotoxin SuVIA suggests an evolutionary link between ancestral predator defence and the origin of fish-hunting behaviour in carnivorous cone snails. Proceedings of the Royal Society B: Biological Sciences, 2015, 282, 20150817.	2.6	29
117	CHAPTER 1. Seeing the Woods for the Trees: Understanding Venom Evolution as a Guide for Biodiscovery. RSC Drug Discovery Series, 2015, , 1-36.	0.3	13
118	$\hat{l}_{\pm}$ -conotoxin MrIC is a biased agonist at $\hat{l}_{\pm}$ 7 nicotinic acetylcholine receptors. Biochemical Pharmacology, 2015, 94, 155-163.	4.4	16
119	Activation of $\hat{I}^2$ Opioid Receptors in Cutaneous Nerve Endings by Conorphin-1, a Novel Subtype-Selective Conopeptide, Does Not Mediate Peripheral Analgesia. ACS Chemical Neuroscience, 2015, 6, 1751-1758.	3.5	17
120	Feeling hot, feeling cold: TRP channelsâ€"a great story unfolds. Temperature, 2015, 2, 150-151.	3.0	6
121	Comparative Venomics Reveals the Complex Prey Capture Strategy of the Piscivorous Cone Snail <i>Conus catus </i> . Journal of Proteome Research, 2015, 14, 4372-4381.	3.7	62
122	Design, Synthesis and Biological Evaluation of Two Opioid Agonist and Ca <sub>v</sub> 2.2 Blocker Multitarget Ligands. Chemical Biology and Drug Design, 2015, 86, 156-162.	3.2	31
123	Ciguatoxin and Ciguatera., 2015, , 1-19.		0
124	Selenoether oxytocin analogues have analgesic properties in a mouse model of chronic abdominal pain. Nature Communications, 2014, 5, 3165.	12.8	122
125	Ciguatera fish poisoning: A first epidemic in Germany highlights an increasing risk for European countries. Toxicon, 2014, 91, 76-83.	1.6	65
126	Analgesic effects of clinically used compounds in novel mouse models of polyneuropathy induced by oxaliplatin and cisplatin. Neuro-Oncology, 2014, 16, 1324-1332.	1.2	44

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127	2â€Nitroveratryl as a Photocleavable Thiolâ€Protecting Group for Directed Disulfide Bond Formation in the Chemical Synthesis of Insulin. Chemistry - A European Journal, 2014, 20, 9549-9552.	3.3	48
128	Isolation and Structural and Pharmacological Characterization of $\hat{l}_{\pm}$ -Elapitoxin-Dpp2d, an Amidated Three Finger Toxin from Black Mamba Venom. Biochemistry, 2014, 53, 3758-3766.	2.5	23
129	No Gain, No Pain: Na <sub>V</sub> 1.7 as an Analgesic Target. ACS Chemical Neuroscience, 2014, 5, 749-751.	3.5	73
130	Evolution of separate predation- and defence-evoked venoms in carnivorous cone snails. Nature Communications, 2014, 5, 3521.	12.8	275
131	Does Nature do Ion Channel Drug Discovery Better than Us?. RSC Drug Discovery Series, 2014, , 297-319.	0.3	2
132	A ray of venom: Combined proteomic and transcriptomic investigation of fish venom composition using barb tissue from the blue-spotted stingray (Neotrygon kuhlii). Journal of Proteomics, 2014, 109, 188-198.	2.4	29
133	MrIC, a Novel α-Conotoxin Agonist in the Presence of PNU at Endogenous α7 Nicotinic Acetylcholine Receptors. Biochemistry, 2014, 53, 1-3.	2.5	31
134	Ciguatera Toxins: Pharmacology, Toxicology, and Detection. , 2014, , 925-950.		17
135	Analgesic treatment of ciguatoxin-induced cold allodynia. Pain, 2013, 154, 1999-2006.	4.2	51
136	An animal model of oxaliplatin-induced cold allodynia reveals a crucial role for Nav1.6 in peripheral pain pathways. Pain, 2013, 154, 1749-1757.	4.2	144
137	Isolation and characterization of $\hat{l}\pm$ -conotoxin LsIA with potent activity at nicotinic acetylcholine receptors. Biochemical Pharmacology, 2013, 86, 791-799.	4.4	51
138	Chemical Engineering and Structural and Pharmacological Characterization of the $\hat{l}_{\pm}$ -Scorpion Toxin OD1. ACS Chemical Biology, 2013, 8, 1215-1222.	3.4	50
139	Differential Evolution and Neofunctionalization of Snake Venom Metalloprotease Domains. Molecular and Cellular Proteomics, 2013, 12, 651-663.	3.8	83
140	Venom Peptides as a Rich Source of Cav2.2 Channel Blockers. Toxins, 2013, 5, 286-314.	3.4	35
141	Amplified Cold Transduction in Native Nociceptors by M-Channel Inhibition. Journal of Neuroscience, 2013, 33, 16627-16641.	3.6	37
142	Multiple actions of φ-LITX-Lw1a on ryanodine receptors reveal a functional link between scorpion DDH and ICK toxins. Proceedings of the National Academy of Sciences of the United States of America, 2013, 110, 8906-8911.	7.1	35
143	Expression and Pharmacology of Endogenous Cav Channels in SH-SY5Y Human Neuroblastoma Cells. PLoS ONE, 2013, 8, e59293.	2.5	50
144	Ciguatoxins activate specific cold pain pathways to elicit burning pain from cooling. EMBO Journal, 2012, 31, 3795-3808.	7.8	103

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145	Therapeutic Potential of Cone Snail Venom Peptides (Conopeptides). Current Topics in Medicinal Chemistry, 2012, 12, 1546-1552.	2.1	96
146	Pharmacological characterization of $\hat{l}_{\pm}$ -elapitoxin-Al2a from the venom of the Australian pygmy copperhead (Austrelaps labialis): An atypical long-chain $\hat{l}_{\pm}$ -neurotoxin with only weak affinity for $\hat{l}_{\pm}$ 7 nicotinic receptors. Biochemical Pharmacology, 2012, 84, 851-863.	4.4	13
147	ω-Conotoxin GVIA Mimetics that Bind and Inhibit Neuronal Cav2.2 Ion Channels. Marine Drugs, 2012, 10, 2349-2368.	4.6	20
148	Conus Venom Peptide Pharmacology. Pharmacological Reviews, 2012, 64, 259-298.	16.0	372
149	Characterisation of Nav types endogenously expressed in human SH-SY5Y neuroblastoma cells. Biochemical Pharmacology, 2012, 83, 1562-1571.	4.4	64
150	Isolation, characterization and total regioselective synthesis of the novel $\hat{1}\frac{1}{4}$ O-conotoxin MfVIA from Conus magnificus that targets voltage-gated sodium channels. Biochemical Pharmacology, 2012, 84, 540-548.	4.4	54
151	32. Development of High Throughput Calcium ChannelÂAssays to Accelerate the Discovery of NovelÂToxins Targeting Human Cav2.2 Channels. Toxicon, 2012, 60, 111.	1.6	0
152	Development and Optimization of FLIPR High Throughput Calcium Assays for Ion Channels and GPCRs. Advances in Experimental Medicine and Biology, 2012, 740, 45-82.	1.6	30
153	α-Conotoxin ImI Incorporating Stable Cystathionine Bridges Maintains Full Potency and Identical Three-Dimensional Structure. Journal of the American Chemical Society, 2011, 133, 15866-15869.	13.7	81
154	Natural Product Ligands of TRP Channels. Advances in Experimental Medicine and Biology, 2011, 704, 41-85.	1.6	31
155	Expression of the G Protein-Coupled Receptor 68 is Increased by TNF-Mediated Inflammatory Signalling. Gastroenterology, 2011, 140, S-837-S-838.	1.3	0
156	Venomics: a new paradigm for natural products-based drug discovery. Amino Acids, 2011, 40, 15-28.	2.7	172
157	Characterization of endogenous calcium responses in neuronal cell lines. Biochemical Pharmacology, 2010, 79, 908-920.	4.4	90
158	Chemical Synthesis and Structure of the Prokineticin Bv8. ChemBioChem, 2010, 11, 1882-1888.	2.6	22
159	Axon guidance by growth-rate modulation. Proceedings of the National Academy of Sciences of the United States of America, 2010, 107, 5202-5207.	7.1	67
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