Lili Wang

List of Publications by Year in descending order

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Version: 2024-02-01

58	7,455	38	59
	citations	h-index	g-index
papers	CITATIONS	n-maex	g-maex
61	61	61	6078 citing authors
all docs	docs citations	times ranked	

#	Article	IF	CITATIONS
1	Prednisolone reduces the interferon response to AAV in cynomolgus macaques and may increase liver gene expression. Molecular Therapy - Methods and Clinical Development, 2022, 24, 292-305.	4.1	10
2	Helper lipid structure influences protein adsorption and delivery of lipid nanoparticles to spleen and liver. Biomaterials Science, 2021, 9, 1449-1463.	5.4	84
3	Advances and challenges in adeno-associated viral inner-ear gene therapy for sensorineural hearing loss. Molecular Therapy - Methods and Clinical Development, 2021, 21, 209-236.	4.1	20
4	Scalable mRNA and siRNA Lipid Nanoparticle Production Using a Parallelized Microfluidic Device. Nano Letters, 2021, 21, 5671-5680.	9.1	120
5	Long-term stable reduction of low-density lipoprotein in nonhuman primates following inÂvivo genome editing of PCSK9. Molecular Therapy, 2021, 29, 2019-2029.	8.2	42
6	Developing a second-generation clinical candidate AAV vector for gene therapy of familial hypercholesterolemia. Molecular Therapy - Methods and Clinical Development, 2021, 22, 1-10.	4.1	14
7	ITR-Seq, a next-generation sequencing assay, identifies genome-wide DNA editing sites in vivo following adeno-associated viral vector-mediated genome editing. BMC Genomics, 2020, 21, 239.	2.8	35
8	A mutation-independent CRISPR-Cas9–mediated gene targeting approach to treat a murine model of ornithine transcarbamylase deficiency. Science Advances, 2020, 6, eaax5701.	10.3	44
9	Ionizable lipid nanoparticles encapsulating barcoded mRNA for accelerated in vivo delivery screening. Journal of Controlled Release, 2019, 316, 404-417.	9.9	111
10	CRISPR/Cas9-mediated in vivo gene targeting corrects hemostasis in newborn and adult factor IX–knockout mice. Blood, 2019, 133, 2745-2752.	1.4	57
11	Meganuclease targeting of PCSK9 in macaque liver leads to stable reduction in serum cholesterol. Nature Biotechnology, 2018, 36, 717-725.	17.5	95
12	Optimized Adeno-Associated Viral-Mediated Human Factor VIII Gene Therapy in Cynomolgus Macaques. Human Gene Therapy, 2018, 29, 1364-1375.	2.7	18
13	Characterization of Adeno-Associated Viral Vector-Mediated Human Factor VIII Gene Therapy in Hemophilia A Mice. Human Gene Therapy, 2017, 28, 392-402.	2.7	29
14	AAV gene therapy corrects OTC deficiency and prevents liver fibrosis in aged OTC-knock out heterozygous mice. Molecular Genetics and Metabolism, 2017, 120, 299-305.	1.1	39
15	Effects of Self-Complementarity, Codon Optimization, Transgene, and Dose on Liver Transduction with AAV8. Human Gene Therapy Methods, 2016, 27, 228-237.	2.1	15
16	A dual AAV system enables the Cas9-mediated correction of a metabolic liver disease in newborn mice. Nature Biotechnology, 2016, 34, 334-338.	17.5	476
17	Crispr/Cas9-Mediated In Vivo Gene Targeting Corrects Haemostasis in Newborn and Adult FIX-KO Mice. Blood, 2016, 128, 1174-1174.	1.4	9
18	Strategies for Selection of AAV Vectors for Administration to Liver: Studies in Nonhuman Primates. Blood, 2016, 128, 2316-2316.	1.4	1

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19	Development and rescue of human familial hypercholesterolaemia in a xenograft mouse model. Nature Communications, 2015, 6, 7339.	12.8	51
20	Comparative Study of Liver Gene Transfer With AAV Vectors Based on Natural and Engineered AAV Capsids. Molecular Therapy, 2015, 23, 1877-1887.	8.2	94
21	AAV8 Induces Tolerance in Murine Muscle as a Result of Poor APC Transduction, T Cell Exhaustion, and Minimal MHCI Upregulation on Target Cells. Molecular Therapy, 2014, 22, 28-41.	8.2	50
22	Mapping the Structural Determinants Responsible for Enhanced T Cell Activation to the Immunogenic Adeno-Associated Virus Capsid from Isolate Rhesus 32.33. Journal of Virology, 2013, 87, 9473-9485.	3.4	24
23	Hepatic Gene Transfer in Neonatal Mice by Adeno-Associated Virus Serotype 8 Vector. Human Gene Therapy, 2012, 23, 533-539.	2.7	90
24	Adeno-associated virus gene therapy prevents hepatocellular adenoma in murine model of glycogen storage disease type la. Hepatology, 2012, 56, 1593-1595.	7.3	2
25	Preclinical evaluation of a clinical candidate AAV8 vector for ornithine transcarbamylase (OTC) deficiency reveals functional enzyme from each persisting vector genome. Molecular Genetics and Metabolism, 2012, 105, 203-211.	1.1	46
26	Novel AAV-based genetic vaccines encoding truncated dengue virus envelope proteins elicit humoral immune responses in mice. Microbes and Infection, 2012, 14, 1000-1007.	1.9	21
27	Inverse zonation of hepatocyte transduction with AAV vectors between mice and non-human primates. Molecular Genetics and Metabolism, 2011, 104, 395-403.	1.1	58
28	Adeno-associated Viral Vectors for Correction of Inborn Errors of Metabolism: Progressing Towards Clinical Application. Current Pharmaceutical Design, 2011, 17, 2500-2515.	1.9	17
29	AAV8-mediated Hepatic Gene Transfer in Infant Rhesus Monkeys (Macaca mulatta). Molecular Therapy, 2011, 19, 2012-2020.	8.2	85
30	Impact of Pre-Existing Immunity on Gene Transfer to Nonhuman Primate Liver with Adeno-Associated Virus 8 Vectors. Human Gene Therapy, 2011, 22, 1389-1401.	2.7	170
31	Dosage Thresholds for AAV2 and AAV8 Photoreceptor Gene Therapy in Monkey. Science Translational Medicine, 2011, 3, 88ra54.	12.4	179
32	Adeno-Associated Virus Antibody Profiles in Newborns, Children, and Adolescents. Vaccine Journal, 2011, 18, 1586-1588.	3.1	269
33	Evaluation of Adeno-Associated Viral Vectors for Liver-Directed Gene Transfer in Dogs. Human Gene Therapy, 2011, 22, 985-997.	2.7	35
34	Systematic Evaluation of AAV Vectors for Liver directed Gene Transfer in Murine Models. Molecular Therapy, 2010, 18, 118-125.	8.2	110
35	The Pleiotropic Effects of Natural AAV Infections on Liver-directed Gene Transfer in Macaques. Molecular Therapy, 2010, 18, 126-134.	8.2	123
36	Host Immune Responses to Chronic Adenovirus Infections in Human and Nonhuman Primates. Journal of Virology, 2009, 83, 2623-2631.	3.4	61

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37	Sustained transgene expression despite T lymphocyte responses in a clinical trial of rAAV1-AAT gene therapy. Proceedings of the National Academy of Sciences of the United States of America, 2009, 106, 16363-16368.	7.1	295
38	Adeno-Associated Virus-Mediated Gene Transfer to Nonhuman Primate Liver Can Elicit Destructive Transgene-Specific T Cell Responses. Human Gene Therapy, 2009, 20, 930-942.	2.7	88
39	Treatment of Leber Congenital Amaurosis Due to <i>RPE65</i> Mutations by Ocular Subretinal Injection of Adeno-Associated Virus Gene Vector: Short-Term Results of a Phase I Trial. Human Gene Therapy, 2008, 19, 979-990.	2.7	880
40	Cross-Presentation of Adeno-Associated Virus Serotype 2 Capsids Activates Cytotoxic T Cells But Does Not Render Hepatocytes Effective Cytolytic Targets. Human Gene Therapy, 2007, 18, 185-194.	2.7	105
41	Heparin binding directs activation of T cells against adeno-associated virus serotype 2 capsid. Nature Medicine, 2006, 12, 967-971.	30.7	193
42	Inhibition of protein phosphatase 1 by inhibitorâ€2 gene delivery ameliorates heart failure progression in genetic cardiomyopathy. FASEB Journal, 2006, 20, 1197-1199.	0.5	77
43	Biology of AAV Serotype Vectors in Liver-Directed Gene Transfer to Nonhuman Primates. Molecular Therapy, 2006, 13, 77-87.	8.2	161
44	Gene transfer into skeletal muscle using novel AAV serotypes. Journal of Gene Medicine, 2005, 7, 442-451.	2.8	108
45	Sustained correction of disease in naive and AAV2-pretreated hemophilia B dogs: AAV2/8-mediated, liver-directed gene therapy. Blood, 2005, 105, 3079-3086.	1.4	162
46	No Evidence for Tumorigenesis of AAV Vectors in a Large-Scale Study in Mice. Molecular Therapy, 2005, 12, 299-306.	8.2	103
47	Total correction of hemophilia A mice with canine FVIII using an AAV 8 serotype. Blood, 2004, 103, 1253-1260.	1.4	188
48	Adenovirus-mediated overexpression of follistatin enlarges intact liver of adult rats. Hepatology, 2003, 38, 1107-1115.	7.3	38
49	Novel adeno-associated viruses from rhesus monkeys as vectors for human gene therapy. Proceedings of the National Academy of Sciences of the United States of America, 2002, 99, 11854-11859.	7.1	1,398
50	Effect of Adenovirus-Mediated Overexpression of Follistatin and Extracellular Domain of Activin Receptor Type II on Gonadotropin Secretion in Vitro and in Vivo. Endocrinology, 2002, 143, 964-969.	2.8	13
51	Chronic suppression of heart-failure progression by a pseudophosphorylated mutant of phospholamban via in vivo cardiac rAAV gene delivery. Nature Medicine, 2002, 8, 864-871.	30.7	344
52	Noninvasive gene transfer to the lung for systemic delivery of therapeutic proteins. Journal of Clinical Investigation, 2002, 110, 499-504.	8.2	104
53	Noninvasive gene transfer to the lung for systemic delivery of therapeutic proteins. Journal of Clinical Investigation, 2002, 110, 499-504.	8.2	76
54	Hybrid Vectors Based on Adeno-Associated Virus Serotypes 2 and 5 for Muscle-Directed Gene Transfer. Journal of Virology, 2001, 75, 6199-6203.	3.4	203

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55	Sustained Expression of Therapeutic Level of Factor IX in Hemophilia B Dogs by AAV-Mediated Gene Therapy in Liver. Molecular Therapy, 2000, 1, 154-158.	8.2	171
56	Activation of the mec-3 promoter in two classes of stereotyped lineages in Caenorhabditis elegans. Mechanisms of Development, 1996, 56, 165-181.	1.7	6
57	Promoter sequences for the establishment of mec-3 expression in the nematode Caenorhabditis elegans. Mechanisms of Development, 1996, 56, 183-196.	1.7	6
58	Cell polarity and the mechanism of asymmetric cell division. BioEssays, 1994, 16, 925-931.	2.5	23