## Maria K Y Chan

## List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/7622769/publications.pdf

Version: 2024-02-01

104 papers 7,418 citations

33 h-index 85 g-index

106 all docs

 $\begin{array}{c} 106 \\ \\ \text{docs citations} \end{array}$ 

106 times ranked 12742 citing authors

#	Article	IF	CITATIONS
1	Data-driven design of novel halide perovskite alloys. Energy and Environmental Science, 2022, 15, 1930-1949.	30.8	26
2	Accelerated screening of functional atomic impurities in halide perovskites using high-throughput computations and machine learning. Journal of Materials Science, 2022, 57, 10736-10754.	3.7	10
3	Universal machine learning framework for defect predictions in zinc blende semiconductors. Patterns, 2022, 3, 100450.	5.9	22
4	Machine learning in scanning transmission electron microscopy. Nature Reviews Methods Primers, 2022, 2, .	21.2	59
5	Machine learning for impurity charge-state transition levels in semiconductors from elemental properties using multi-fidelity datasets. Journal of Chemical Physics, 2022, 156, 114110.	3.0	5
6	Ingrained: An Automated Framework for Fusing Atomicâ€Scale Image Simulations into Experiments. Small, 2022, 18, e2102960.	10.0	12
7	Exploring the Origin of Anionic Redox Activity in Super Li-Rich Iron Oxide-Based High-Energy-Density Cathode Materials. Chemistry of Materials, 2022, 34, 4536-4547.	6.7	10
8	Origin and regulation of oxygen redox instability in high-voltage battery cathodes. Nature Energy, 2022, 7, 808-817.	39.5	55
9	First principles modeling of polymer encapsulant degradation in Si photovoltaic modules. Physical Chemistry Chemical Physics, 2021, 23, 10357-10364.	2.8	2
10	Synthesis of borophane polymorphs through hydrogenation of borophene. Science, 2021, 371, 1143-1148.	12.6	129
11	Unraveling Cu chemical signature in CdTe by Spectral Fluorescence Mapping. , 2021, , .		1
12			
	The Role of Water on the Interfacial Adhesion in Si Solar Modules. , 2021, , .		1
13	The Role of Water on the Interfacial Adhesion in Si Solar Modules. , 2021, , .  Open-Source Tools and Containers for the Production of Large-Scale S/TEM Datasets. Microscopy and Microanalysis, 2021, 27, 62-63.	0.4	1
13 14	Open-Source Tools and Containers for the Production of Large-Scale S/TEM Datasets. Microscopy and	0.4	
	Open-Source Tools and Containers for the Production of Large-Scale S/TEM Datasets. Microscopy and Microanalysis, 2021, 27, 62-63.  4D >Crystal: Deep Learning Crystallographic Information From Electron Diffraction Images.		1
14	Open-Source Tools and Containers for the Production of Large-Scale S/TEM Datasets. Microscopy and Microanalysis, 2021, 27, 62-63.  4D >Crystal: Deep Learning Crystallographic Information From Electron Diffraction Images. Microscopy and Microanalysis, 2021, 27, 2774-2776.  Machine learning on neutron and x-ray scattering and spectroscopies. Chemical Physics Reviews, 2021,	0.4	2
14 15	Open-Source Tools and Containers for the Production of Large-Scale S/TEM Datasets. Microscopy and Microanalysis, 2021, 27, 62-63.  4D >Crystal: Deep Learning Crystallographic Information From Electron Diffraction Images. Microscopy and Microanalysis, 2021, 27, 2774-2776.  Machine learning on neutron and x-ray scattering and spectroscopies. Chemical Physics Reviews, 2021, 2, .	0.4	1 2 49

#	Article	IF	CITATIONS
19	Constructing Self-Labeled Materials Imaging Datasets from Open Access Scientific Journals with EXSCLAIM!. Microscopy and Microanalysis, 2020, 26, 3096-3097.	0.4	1
20	Machine-learned impurity level prediction for semiconductors: the example of Cd-based chalcogenides. Npj Computational Materials, 2020, 6, .	8.7	32
21	Defect Energetics in Pseudo-Cubic Mixed Halide Lead Perovskites from First-Principles. Journal of Physical Chemistry C, 2020, 124, 16729-16738.	3.1	19
22	Direct Observation of Bandgap Oscillations Induced by Optical Phonons in Hybrid Lead Iodide Perovskites. Advanced Functional Materials, 2020, 30, 1907982.	14.9	15
23	Validating first-principles molecular dynamics calculations of oxide/water interfaces with x-ray reflectivity data. Physical Review Materials, 2020, 4, .	2.4	12
24	Cu-Local Structures and Their Relation with Nanoscale Electrical Performance in CdTe. , 2020, , .		2
25	Stabilization of a monolayer tellurene phase at CdTe interfaces. Nanoscale, 2019, 11, 14698-14706.	5.6	10
26	Effect of selenium and chlorine co-passivation in polycrystalline CdSeTe devices. Applied Physics Letters, 2019, 115, .	3.3	33
27	Spectroscopic Comparison of Thermal Transport at Organic–Inorganic and Organic-Hybrid Interfaces Using CsPbBr <sub>3</sub> (FA = Formamidinium) Perovskite Nanocrystals. Nano Letters, 2019, 19, 8155-8160.	9.1	4
28	Study of Effects of Cl and Se in CdSeTe Solar Cells Using Scanning Transmission Electron Microscopy. Microscopy and Microanalysis, 2019, 25, 2150-2151.	0.4	5
29	Fluoroethylene Carbonate Breakdown Mechanisms and Energetics on Two Lithium Silicide Surfaces. Journal of Physical Chemistry C, 2019, 123, 26743-26751.	3.1	4
30	Infrared-pump electronic-probe of methylammonium lead iodide reveals electronically decoupled organic and inorganic sublattices. Nature Communications, 2019, 10, 482.	12.8	25
31	Understanding the Role of Overpotentials in Lithium Ion Conversion Reactions: Visualizing the Interface. ACS Nano, 2019, 13, 7825-7832.	14.6	16
32	Atomistic manipulation of reversible oxidation and reduction in Ag with an electron beam. Nanoscale, 2019, 11, 10756-10762.	5.6	14
33	Comprehensive Computational Study of Partial Lead Substitution in Methylammonium Lead Bromide. Chemistry of Materials, 2019, 31, 3599-3612.	6.7	37
34	Probing Electrochemically Induced Structural Evolution and Oxygen Redox Reactions in Layered Lithium Iridate. Chemistry of Materials, 2019, 31, 4341-4352.	6.7	26
35	Seebeck Tensor Analysis of (p × n)-type Transverse Thermoelectric Materials. MRS Advances, 2019, 4, 491-497.	0.9	2
36	Charge Transfer Dynamics of Phase-Segregated Halide Perovskites: CH <sub>3</sub> NH <sub>3</sub> PbCl <sub>3</sub> and CH <sub>3</sub> NH <sub>3</sub> Pbl <sub>3</sub> or (C <sub>4</sub> H <sub>9</sub> NH <sub>3</sub> NH <sub>3</sub> ) <sub>2</sub> (CH <sub>3</sub> NH <sub>3</sub> ) <sub>&gt;6 Mixtures. ACS Applied Materials &amp; Samp; Interfaces, 2019, 11, 9583-9593.</sub>	8.0 ·na^'1 </td <td>14 :/sub&gt;Pb<i><s< td=""></s<></i></td>	14 :/sub>Pb <i><s< td=""></s<></i>

#	Article	IF	Citations
37	Study of Se and Cl segregation in poly-crystalline CdSeTe. , 2019, , .		O
38	Machine Learning Classical Interatomic Potentials for Molecular Dynamics from First-Principles Training Data. Journal of Physical Chemistry C, 2019, 123, 6941-6957.	3.1	72
39	Operando Observations and Firstâ€Principles Calculations of Reduced Lithium Insertion in Auâ€Coated LiMn 2 O 4. Advanced Materials Interfaces, 2019, 6, 1801923.	3.7	11
40	Identifying the Chemical Origin of Oxygen Redox Activity in Li-Rich Anti-Fluorite Lithium Iron Oxide by Experimental and Theoretical X-ray Absorption Spectroscopy. Journal of Physical Chemistry Letters, 2019, 10, 806-812.	4.6	17
41	Computational prediction of lattice thermal conductivity: A comparison of molecular dynamics and Boltzmann transport approaches. Physical Review Materials, $2019, 3, .$	2.4	29
42	Controlling Interfacial Properties of Lithiumâ€lon Battery Cathodes with Alkylphosphonate Selfâ€Assembled Monolayers. Advanced Materials Interfaces, 2018, 5, 1701292.	3.7	22
43	Imaging Catalytic Activation of CO <sub>2</sub> on Cu <sub>2</sub> O (110): A First-Principles Study. Chemistry of Materials, 2018, 30, 1912-1923.	6.7	56
44	Quantitative Observation of Threshold Defect Behavior in Memristive Devices with <i>Operando</i> X-ray Microscopy. ACS Nano, 2018, 12, 4938-4945.	14.6	12
45	Mechanism of Na accumulation at extended defects in Si from first-principles. Journal of Applied Physics, 2018, 123, 161560.	2.5	12
46	Atomistic determination of the surface structure of Cu <sub>2</sub> O(111): experiment and theory. Physical Chemistry Chemical Physics, 2018, 20, 27456-27463.	2.8	33
47	An Autonomous Microscopy Workflow for Structure Determination from Atomic-Resolution Images. Microscopy and Microanalysis, 2018, 24, 510-511.	0.4	3
48	Ultralow-loading platinum-cobalt fuel cell catalysts derived from imidazolate frameworks. Science, 2018, 362, 1276-1281.	12.6	735
49	Anharmonic stabilization and lattice heat transport in rocksalt $\langle i \rangle \hat{l}^2 \langle  i \rangle$ -GeTe. Applied Physics Letters, 2018, 113, .	3.3	39
50	Atomic-Resolution Study of Grain Boundaries in CdTe Using Scanning Transmission Electron Microscopy. Microscopy and Microanalysis, 2018, 24, 102-103.	0.4	2
51	Two-Dimensional CsAg <sub>5</sub> Te <sub>3–<i>x</i></sub> S <sub><i>x</i></sub> Semiconductors: Multi-anion Chalcogenides with Dynamic Disorder and Ultralow Thermal Conductivity. Chemistry of Materials, 2018, 30, 7245-7254.	6.7	15
52	Lattice thermal transport in group II-alloyed PbTe. Applied Physics Letters, 2018, 112, .	3.3	26
53	Slow thermal equilibration in methylammonium lead iodide revealed by transient mid-infrared spectroscopy. Nature Communications, 2018, 9, 2792.	12.8	25
54	A scripting interface for <i>GSAS-II</i> . Journal of Applied Crystallography, 2018, 51, 1244-1250.	4.5	17

#	Article	IF	CITATIONS
55	Template-Free Vapor-Phase Growth of Patr $\tilde{A}^3$ nite by Atomic Layer Deposition. Chemistry of Materials, 2017, 29, 2864-2873.	6.7	37
56	Evolutionary Optimization of a Charge Transfer Ionic Potential Model for Ta/Ta-Oxide Heterointerfaces. Chemistry of Materials, 2017, 29, 3603-3614.	6.7	22
57	Probing the Release and Uptake of Water in α-MnO <sub>2</sub> Â- <i>x</i> H <sub>2</sub> O. Chemistry of Materials, 2017, 29, 1507-1517.	6.7	31
58	The Role of Water in the Reversible Optoelectronic Degradation of Hybrid Perovskites at Low Pressure. Journal of Physical Chemistry C, 2017, 121, 25659-25665.	3.1	19
59	Theory of Thermal Relaxation of Electrons in Semiconductors. Physical Review Letters, 2017, 119, 136602.	7.8	78
60	Polar Fluctuations in Metal Halide Perovskites Uncovered by Acoustic Phonon Anomalies. ACS Energy Letters, 2017, 2, 2463-2469.	17.4	47
61	Machine learnt bond order potential to model metal–organic (Co–C) heterostructures. Nanoscale, 2017, 9, 18229-18239.	5.6	11
62	Machine Learning Force Field Parameters from Ab Initio Data. Journal of Chemical Theory and Computation, 2017, 13, 4492-4503.	5 <b>.</b> 3	105
63	Enabling the high capacity of lithium-rich anti-fluorite lithium iron oxide by simultaneous anionic and cationic redox. Nature Energy, 2017, 2, 963-971.	39.5	140
64	Leveraging First Principles Modeling and Machine Learning for Microscopy Data Inversion. Microscopy and Microanalysis, 2017, 23, 178-179.	0.4	1
65	Atomic — scale study of model CdTe grain boundaries. , 2017, , .		0
66	Slow Organicâ€toâ€Inorganic Subâ€Lattice Thermalization in Methylammonium Lead Halide Perovskites Observed by Ultrafast Photoluminescence. Advanced Energy Materials, 2016, 6, 1600422.	19.5	32
67	Unraveling the Planar-Globular Transition in Gold Nanoclusters through Evolutionary Search. Scientific Reports, 2016, 6, 34974.	3.3	21
68	Atomic — Scale study of model CdTe grain boundaries. , 2016, , .		0
69	First principles modeling of grain boundaries in CdTe. , 2016, , .		0
70	Enabling direct silicene integration in electronics: First principles study of silicene on NiSi2(111). Applied Physics Letters, 2016, 109, .	3.3	6
71	Atomic and electronic structure of Lomer dislocations at CdTe bicrystal interface. Scientific Reports, 2016, 6, 27009.	3.3	35
72	Atomistic Study of Model CdTe Grain Boundaries. Microscopy and Microanalysis, 2016, 22, 1398-1399.	0.4	0

#	Article	IF	CITATIONS
73	Development of a Modified Embedded Atom Force Field for Zirconium Nitride Using Multi-Objective Evolutionary Optimization. Journal of Physical Chemistry C, 2016, 120, 17475-17483.	3.1	23
74	<i>Ab Initio</i> -Based Bond Order Potential to Investigate Low Thermal Conductivity of Stanene Nanostructures. Journal of Physical Chemistry Letters, 2016, 7, 3752-3759.	4.6	80
75	Describing the Diverse Geometries of Gold from Nanoclusters to Bulk—A First-Principles-Based Hybrid Bond-Order Potential. Journal of Physical Chemistry C, 2016, 120, 13787-13800.	3.1	26
76	Visualizing Redox Dynamics of a Single Ag/AgCl Heterogeneous Nanocatalyst at Atomic Resolution. ACS Nano, 2016, 10, 3738-3746.	14.6	61
77	Electronic transport in VO2—Experimentally calibrated Boltzmann transport modeling. Applied Physics Letters, 2015, 107, .	3.3	7
78	A fundamental study of the effects of grain boundaries on performance of poly-crystalline thin film CdTe solar cells. , $2015$ , , .		0
79	Atomic Scale Study of Lomer-Cottrell and Hirth Lock Dislocations in CdTe. Microscopy and Microanalysis, 2015, 21, 2087-2088.	0.4	2
80	Atomistic simulations of grain boundaries in CdTe. , 2015, , .		3
81	Creation and analysis of atomic structures for CdTe bi-crystal interfaces by the grain boundary genie. , 2015, , .		2
82	Towards First Principles-Based Prediction of Highly Accurate Electrochemical Pourbaix Diagrams. Journal of Physical Chemistry C, 2015, 119, 18177-18187.	3.1	97
83	Understanding Polyol Decomposition on Bimetallic Pt–Mo Catalysts—A DFT Study of Glycerol. ACS Catalysis, 2015, 5, 4942-4950.	11.2	25
84	Edge-terminated molybdenum disulfide with a 9.4-Ã interlayer spacing for electrochemical hydrogen production. Nature Communications, 2015, 6, 7493.	12.8	628
85	In situ high-energy synchrotron X-ray diffraction studies and first principles modeling of α-MnO <sub>2</sub> electrodes in Li–O <sub>2</sub> and Li-ion coin cells. Journal of Materials Chemistry A, 2015, 3, 7389-7398.	10.3	43
86	First-Principles Analysis of Defect Thermodynamics and Ion Transport in Inorganic SEI Compounds: LiF and NaF. ACS Applied Materials & Samp; Interfaces, 2015, 7, 18985-18996.	8.0	191
87	Atomic scale study of polar Lomer–Cottrell and Hirth lock dislocation cores in CdTe. Acta Crystallographica Section A: Foundations and Advances, 2014, 70, 524-531.	0.1	31
88	First-Principles Analysis of Defect-Mediated Li Adsorption on Graphene. ACS Applied Materials & Samp; Interfaces, 2014, 6, 21141-21150.	8.0	120
89	Thermal Stability of Colloidal InP Nanocrystals: Small Inorganic Ligands Boost High-Temperature Photoluminescence. ACS Nano, 2014, 8, 977-985.	14.6	57
90	Polaron Structure and Transport in Fullerene Materials: Insights from First-Principles Calculations. Journal of Physical Chemistry C, 2014, 118, 21785-21797.	3.1	6

#	Article	IF	CITATIONS
91	Defect Evolution in Graphene upon Electrochemical Lithiation. ACS Applied Materials & Samp; Interfaces, 2014, 6, 17626-17636.	8.0	30
92	Atomistic and First Principles: Computational Studies of LiO2 Batteries., 2014, , 159-177.		2
93	Vision for Designing High-Energy, Hybrid Li Ion/Li–O <sub>2</sub> Cells. Journal of Physical Chemistry Letters, 2013, 4, 3607-3611.	4.6	26
94	Surface Coverage and SEI Induced Electrochemical Surface Stress Changes during Li Deposition in a Model System for Li-lon Battery Anodes. Journal of the Electrochemical Society, 2013, 160, A888-A896.	2.9	55
95	Synthesis, Characterization, and Structural Modeling of Highâ€Capacity, Dual Functioning MnO <sub>2</sub> Electrode/Electrocatalysts for Liâ€O <sub>2</sub> Cells. Advanced Energy Materials, 2013, 3, 75-84.	19.5	111
96	UNCERTAINTY CLASSIFICATION AND VISUALIZATION OF MOLECULAR INTERFACES., 2013, 3, 157-169.		10
97	First Principles Simulations of the Electrochemical Lithiation and Delithiation of Faceted Crystalline Silicon. Journal of the American Chemical Society, 2012, 134, 14362-14374.	13.7	221
98	Trends in activity for the water electrolyser reactions on 3d M(Ni,Co,Fe,Mn) hydr(oxy)oxide catalysts. Nature Materials, 2012, 11, 550-557.	27.5	2,423
99	On the Origin of Photoluminescence in Silicon Nanocrystals: Pressure-Dependent Structural and Optical Studies. Nano Letters, 2012, 12, 4200-4205.	9.1	133
100	The First-Cycle Electrochemical Lithiation of Crystalline Ge: Dopant and Orientation Dependence and Comparison with Si. Journal of Physical Chemistry Letters, 2011, 2, 3092-3095.	4.6	48
101	Structure of Lithium Peroxide. Journal of Physical Chemistry Letters, 2011, 2, 2483-2486.	4.6	53
102	Dopant Modulated Li Insertion in Si for Battery Anodes: Theory and Experiment. Journal of Physical Chemistry C, 2011, 115, 18916-18921.	3.1	84
103	Electronic structure of lithium battery interphase compounds: Comparison between inelastic x-ray scattering measurements and theory. Journal of Chemical Physics, 2011, 135, 224513.	3.0	39
104	Efficient Band Gap Prediction for Solids. Physical Review Letters, 2010, 105, 196403.	7.8	398