## List of Publications by Year in descending order

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LINELIN

#	Article	IF	CITATIONS
1	Parallel transport dynamics for mixed quantum states with applications to time-dependent density functional theory. Journal of Computational Physics, 2022, 451, 110850.	3.8	2
2	Pure State <i>v</i> -Representability of Density Matrix Embedding Theory. Journal of Chemical Theory and Computation, 2022, 18, 851-864.	5.3	8
3	Staggered Mesh Method for Correlation Energy Calculations of Solids: Random Phase Approximation in Direct Ring Coupled Cluster Doubles and Adiabatic Connection Formalisms. Journal of Chemical Theory and Computation, 2022, 18, 763-775.	5.3	7
4	Heisenberg-Limited Ground-State Energy Estimation for Early Fault-Tolerant Quantum Computers. PRX Quantum, 2022, 3, .	9.2	40
5	Quantum Linear System Solver Based on Time-optimal Adiabatic Quantum Computing and Quantum Approximate Optimization Algorithm. ACM Transactions on Quantum Computing, 2022, 3, 1-28.	4.3	28
6	Universal approximation of symmetric and anti-symmetric functions. Communications in Mathematical Sciences, 2022, 20, 1397-1408.	1.0	4
7	KSSOLV 2.0: An efficient MATLAB toolbox for solving the Kohn-Sham equations with plane-wave basis set. Computer Physics Communications, 2022, 279, 108424.	7.5	9
8	Towards sharp error analysis of extended Lagrangian molecular dynamics. Journal of Computational Physics, 2022, 466, 111403.	3.8	0
9	86 PFLOPS Deep Potential Molecular Dynamics simulation of 100 million atoms with ab initio accuracy. Computer Physics Communications, 2021, 259, 107624.	7.5	100
10	Learning the Mapping \$\$mathbf {x}mapsto sumlimits _{i=1}^d x_i^2\$\$: the Cost of Finding the Needle in a Haystack. Communications on Applied Mathematics and Computation, 2021, 3, 313-335.	1.7	1
11	Low-Rank Representation of Tensor Network Operators with Long-Range Pairwise Interactions. SIAM Journal of Scientific Computing, 2021, 43, A164-A192.	2.8	3
12	Efficient phase-factor evaluation in quantum signal processing. Physical Review A, 2021, 103, .	2.5	39
13	Numerical solution of large scale Hartree–Fock–Bogoliubov equations. ESAIM: Mathematical Modelling and Numerical Analysis, 2021, 55, 763-787.	1.9	3
14	Random circuit block-encoded matrix and a proposal of quantum LINPACK benchmark. Physical Review A, 2021, 103, .	2.5	8
15	Staggered Mesh Method for Correlation Energy Calculations of Solids: Second-Order MÃ,ller–Plesset Perturbation Theory. Journal of Chemical Theory and Computation, 2021, 17, 4733-4745.	5.3	6
16	Bold Feynman Diagrams and the Luttinger–Ward Formalism Via Gibbs Measures: Non-perturbative Analysis. Archive for Rational Mechanics and Analysis, 2021, 242, 527-579.	2.4	1
17	Bold Feynman Diagrams and the Luttinger–Ward Formalism via Gibbs Measures: Perturbative Approach. Archive for Rational Mechanics and Analysis, 2021, 242, 581-642.	2.4	1
18	Realizing Effective Cubic-Scaling Coulomb Hole Plus Screened Exchange Approximation in Periodic Systems via Interpolative Separable Density Fitting with a Plane-Wave Basis Set. Journal of Physical Chemistry A, 2021, 125, 7545-7557.	2.5	13

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19	Split representation of adaptively compressed polarizability operator. Research in Mathematical Sciences, 2021, 8, 1.	1.0	3
20	Convergence of stochastic-extended Lagrangian molecular dynamics method for polarizable force field simulation. Journal of Computational Physics, 2021, 438, 110338.	3.8	3
21	Reinforcement Learning for Many-Body Ground-State Preparation Inspired by Counterdiabatic Driving. Physical Review X, 2021, 11, .	8.9	29
22	Fast inversion, preconditioned quantum linear system solvers, fast Green's-function computation, and fast evaluation of matrix functions. Physical Review A, 2021, 104, .	2.5	34
23	Deep Density: Circumventing the Kohn-Sham equations via symmetry preserving neural networks. Journal of Computational Physics, 2021, 443, 110523.	3.8	11
24	Systematically Improvable Tensor Hypercontraction: Interpolative Separable Density-Fitting for Molecules Applied to Exact Exchange, Second- and Third-Order MÃ,ller–Plesset Perturbation Theory. Journal of Chemical Theory and Computation, 2020, 16, 243-263.	5.3	44
25	Stochastic Constrained Extended System Dynamics for Solving Charge Equilibration Models. Journal of Chemical Theory and Computation, 2020, 16, 5991-5998.	5.3	5
26	Fast optical absorption spectra calculations for periodic solid state systems. Communications in Applied Mathematics and Computational Science, 2020, 15, 89-113.	1.8	10
27	Sparsity Pattern of the Self-energy for Classical and Quantum Impurity Problems. Annales Henri Poincare, 2020, 21, 2219-2257.	1.7	1
28	Enhancing robustness and efficiency of density matrix embedding theory via semidefinite programming and local correlation potential fitting. Physical Review B, 2020, 102, .	3.2	13
29	Quantum Dynamics with the Parallel Transport Gauge. Multiscale Modeling and Simulation, 2020, 18, 612-645.	1.6	3
30	Efficient hybridization fitting for dynamical mean-field theory via semi-definite relaxation. Physical Review B, 2020, 101, .	3.2	18
31	S <scp>iesta</scp> : Recent developments and applications. Journal of Chemical Physics, 2020, 152, 204108.	3.0	229
32	ELSI — An open infrastructure for electronic structure solvers. Computer Physics Communications, 2020, 256, 107459.	7.5	27
33	Mesh independence of the generalized Davidson algorithm. Journal of Computational Physics, 2020, 409, 109322.	3.8	3
34	Discontinuous Galerkin discretization for quantum simulation of chemistry. New Journal of Physics, 2020, 22, 093015.	2.9	12
35	Influence of point defects on the electronic and topological properties of monolayer <mml:math xmlns:mml="http://www.w3.org/1998/Math/MathML"&gt; <mml:msub> <mml:mi>WTe </mml:mi> <mml:mn>2 Physical Review B, 2020, 102, .</mml:mn></mml:msub></mml:math 	m <b>a.2</b> <td>l<b>:m</b>€ub&gt;</td>	l <b>:m</b> €ub>
36	Semidefinite Relaxation of Multimarginal Optimal Transport for Strictly Correlated Electrons in	2.8	5

Second Quantization. SIAM Journal of Scientific Computing, 2020, 42, B1462-B1489.

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37	Convergence of Adaptive Compression Methods for Hartreeâ€Fockâ€Like Equations. Communications on Pure and Applied Mathematics, 2019, 72, 451-499.	3.1	7
38	Structured Quasi-Newton Methods for Optimization with Orthogonality Constraints. SIAM Journal of Scientific Computing, 2019, 41, A2239-A2269.	2.8	18
39	Projected density matrix embedding theory with applications to the two-dimensional Hubbard model. Journal of Chemical Physics, 2019, 151, .	3.0	22
40	A multiscale neural network based on hierarchical nested bases. Research in Mathematical Sciences, 2019, 6, 1.	1.0	19
41	Numerical methods for Kohn–Sham density functional theory. Acta Numerica, 2019, 28, 405-539.	10.7	23
42	Fast real-time time-dependent hybrid functional calculations with the parallel transport gauge and the adaptively compressed exchange formulation. Computer Physics Communications, 2019, 240, 21-29.	7.5	11
43	Room-temperature magnetism and tunable energy gaps in edge-passivated zigzag graphene quantum dots. Npj 2D Materials and Applications, 2019, 3, .	7.9	25
44	Globally Constructed Adaptive Local Basis Set for Spectral Projectors of Second Order Differential Operators. Multiscale Modeling and Simulation, 2019, 17, 92-116.	1.6	2
45	Variational Formulation for Wannier Functions with Entangled Band Structure. Multiscale Modeling and Simulation, 2019, 17, 167-191.	1.6	11
46	A Multiscale Neural Network Based on Hierarchical Matrices. Multiscale Modeling and Simulation, 2019, 17, 1189-1213.	1.6	24
47	Parallel transport time-dependent density functional theory calculations with hybrid functional on summit. , 2019, , .		7
48	Interpolative Separable Density Fitting through Centroidal Voronoi Tessellation with Applications to Hybrid Functional Electronic Structure Calculations. Journal of Chemical Theory and Computation, 2018, 14, 1311-1320.	5.3	39
49	Variational structure of Luttinger–Ward formalism and bold diagrammatic expansion for Euclidean lattice field theory. Proceedings of the National Academy of Sciences of the United States of America, 2018, 115, 2282-2286.	7.1	5
50	Two-Level Chebyshev Filter Based Complementary Subspace Method: Pushing the Envelope of Large-Scale Electronic Structure Calculations. Journal of Chemical Theory and Computation, 2018, 14, 2930-2946.	5.3	28
51	PSelInv – A distributed memory parallel algorithm for selected inversion: The non-symmetric case. Parallel Computing, 2018, 74, 84-98.	2.1	7
52	ELSI: A unified software interface for Kohn–Sham electronic structure solvers. Computer Physics Communications, 2018, 222, 267-285.	7.5	78
53	Fast Real-Time Time-Dependent Density Functional Theory Calculations with the Parallel Transport Gauge. Journal of Chemical Theory and Computation, 2018, 14, 5645-5652.	5.3	20
54	Disentanglement via Entanglement: A Unified Method for Wannier Localization. Multiscale Modeling and Simulation, 2018, 16, 1392-1410.	1.6	25

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55	Accelerating Optical Absorption Spectra and Exciton Energy Computation via Interpolative Separable Density Fitting. Lecture Notes in Computer Science, 2018, , 604-617.	1.3	12
56	PEXSI-\$Sigma\$: a Green's function embedding method for Kohn–Sham density functional theory. Annals of Mathematical Sciences and Applications, 2018, 3, 441-472.	0.4	4
57	Adaptively Compressed Exchange Operator for Large-Scale Hybrid Density Functional Calculations with Applications to the Adsorption of Water on Silicene. Journal of Chemical Theory and Computation, 2017, 13, 1188-1198.	5.3	38
58	Adaptive local basis set for Kohn–Sham density functional theory in a discontinuous Galerkin framework II: Force, vibration, and molecular dynamics calculations. Journal of Computational Physics, 2017, 335, 426-443.	3.8	29
59	Randomized estimation of spectral densities of large matrices made accurate. Numerische Mathematik, 2017, 136, 183-213.	1.9	9
60	Localized spectrum slicing. Mathematics of Computation, 2017, 86, 2345-2371.	2.1	4
61	SCDM-k: Localized orbitals for solids via selected columns of the density matrix. Journal of Computational Physics, 2017, 334, 1-15.	3.8	36
62	Adaptively Compressed Polarizability Operator for Accelerating Large Scale <i>Ab Initio</i> Phonon Calculations. Multiscale Modeling and Simulation, 2017, 15, 29-55.	1.6	22
63	Highly Efficient Photocatalytic Water Splitting over Edge-Modified Phosphorene Nanoribbons. Journal of the American Chemical Society, 2017, 139, 15429-15436.	13.7	244
64	Robust determination of the chemical potential in the pole expansion and selected inversion method for solving Kohn-Sham density functional theory. Journal of Chemical Physics, 2017, 147, 144107.	3.0	5
65	Interpolative Separable Density Fitting Decomposition for Accelerating Hybrid Density Functional Calculations with Applications to Defects in Silicon. Journal of Chemical Theory and Computation, 2017, 13, 5420-5431.	5.3	53
66	Projected Commutator DIIS Method for Accelerating Hybrid Functional Electronic Structure Calculations. Journal of Chemical Theory and Computation, 2017, 13, 5458-5467.	5.3	21
67	<i>A posteriori</i> error estimates for discontinuous Galerkin methods using non-polynomial basis functions. Part II: Eigenvalue problems. ESAIM: Mathematical Modelling and Numerical Analysis, 2017, 51, 1733-1753.	1.9	4
68	A posteriorierror estimates for discontinuous Galerkin methods using non-polynomial basis functions Part I: Second order linear PDE. ESAIM: Mathematical Modelling and Numerical Analysis, 2016, 50, 1193-1222.	1.9	6
69	Low rank approximation in G 0 W 0 calculations. Science China Mathematics, 2016, 59, 1593-1612.	1.7	14
70	Chebyshev polynomial filtered subspace iteration in the discontinuous Galerkin method for large-scale electronic structure calculations. Journal of Chemical Physics, 2016, 145, 154101.	3.0	28
71	Adaptively Compressed Exchange Operator. Journal of Chemical Theory and Computation, 2016, 12, 2242-2249.	5.3	118
72	Decay estimates of discretized Green's functions for Schrödinger type operators. Science China Mathematics, 2016, 59, 1561-1578.	1.7	5

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73	Approximating Spectral Densities of Large Matrices. SIAM Review, 2016, 58, 34-65.	9.5	84
74	Edge-Modified Phosphorene Nanoflake Heterojunctions as Highly Efficient Solar Cells. Nano Letters, 2016, 16, 1675-1682.	9.1	176
75	DGDFT: A massively parallel method for large scale density functional theory calculations. Journal of Chemical Physics, 2015, 143, 124110.	3.0	55
76	Numerical integration for ab initio many-electron self energy calculations within the GW approximation. Journal of Computational Physics, 2015, 286, 1-13.	3.8	15
77	Edge reconstruction in armchair phosphorene nanoribbons revealed by discontinuous Galerkin density functional theory. Physical Chemistry Chemical Physics, 2015, 17, 31397-31404.	2.8	37
78	Compressed Representation of Kohn–Sham Orbitals via Selected Columns of the Density Matrix. Journal of Chemical Theory and Computation, 2015, 11, 1463-1469.	5.3	72
79	<i>GW</i> 100: Benchmarking <i>G</i> <sub>0</sub> <i>W</i> <sub>0</sub> for Molecular Systems. Journal of Chemical Theory and Computation, 2015, 11, 5665-5687.	5.3	280
80	<i>A posteriori</i> error estimator for adaptive local basis functions to solve Kohn–Sham density functional theory. Communications in Mathematical Sciences, 2015, 13, 1741-1773.	1.0	10
81	Electronic structure and aromaticity of large-scale hexagonal graphene nanoflakes. Journal of Chemical Physics, 2014, 141, 214704.	3.0	40
82	Analysis of Time Reversible Born-Oppenheimer Molecular Dynamics. Entropy, 2014, 16, 110-137.	2.2	24
83	SIESTA-PEXSI: massively parallel method for efficient and accurate <i>ab initio</i> materials simulation without matrix diagonalization. Journal of Physics Condensed Matter, 2014, 26, 305503.	1.8	25
84	Parallel eigenvalue calculation based on multiple shift–invert Lanczos and contour integral based spectral projection method. Parallel Computing, 2014, 40, 195-212.	2.1	25
85	Accelerating atomic orbital-based electronic structure calculation via pole expansion and selected inversion. Journal of Physics Condensed Matter, 2013, 25, 295501.	1.8	50
86	Kantorovich dual solution for strictly correlated electrons in atoms and molecules. Physical Review B, 2013, 87, .	3.2	32
87	Elliptic Preconditioner for Accelerating the Self-Consistent Field Iteration in Kohn–Sham Density Functional Theory. SIAM Journal of Scientific Computing, 2013, 35, S277-S298.	2.8	51
88	Element orbitals for Kohn-Sham density functional theory. Physical Review B, 2012, 85, .	3.2	5
89	Adaptive local basis set for Kohn–Sham density functional theory in a discontinuous Galerkin framework I: Total energy calculation. Journal of Computational Physics, 2012, 231, 2140-2154.	3.8	162
90	Optimized local basis set for Kohn–Sham density functional theory. Journal of Computational Physics, 2012, 231, 4515-4529.	3.8	9

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91	Correlated Tunneling in Hydrogen Bonds. Journal of Statistical Physics, 2011, 145, 365-384.	1.2	45
92	Fast construction of hierarchical matrix representation from matrix–vector multiplication. Journal of Computational Physics, 2011, 230, 4071-4087.	3.8	72
93	Momentum distribution, vibrational dynamics, and the potential of mean force in ice. Physical Review B, 2011, 83, .	3.2	33
94	SellnvAn Algorithm for Selected Inversion of a Sparse Symmetric Matrix. ACM Transactions on Mathematical Software, 2011, 37, 1-19.	2.9	167
95	Displaced Path Integral Formulation for the Momentum Distribution of Quantum Particles. Physical Review Letters, 2010, 105, 110602.	7.8	49
96	Pole-Based approximation of the Fermi-Dirac function. Chinese Annals of Mathematics Series B, 2009, 30, 729-742.	0.4	46
97	Fast algorithm for extracting the diagonal of the inverse matrix with application to the electronic structure analysis of metallic systems. Communications in Mathematical Sciences, 2009, 7, 755-777.	1.0	59
98	Variational Embedding for Quantum Manyâ€Body Problems. Communications on Pure and Applied Mathematics, 0, , .	3.1	3
99	Time-dependent unbounded Hamiltonian simulation with vector norm scaling. Quantum - the Open Journal for Quantum Science, 0, 5, 459.	0.0	23
100	Optimal polynomial based quantum eigenstate filtering with application to solving quantum linear systems. Quantum - the Open Journal for Quantum Science, 0, 4, 361.	0.0	44
101	Near-optimal ground state preparation. Quantum - the Open Journal for Quantum Science, 0, 4, 372.	0.0	64
102	Time-dependent Hamiltonian Simulation of Highly Oscillatory Dynamics and Superconvergence for	0.0	7

Time-dependent Hamiltonian Simulation of Highly Oscillatory Dynamics and Superconvergence for SchrĶdinger Equation. Quantum - the Open Journal for Quantum Science, 0, 6, 690. 102