

Bill Poirier

List of Publications by Year in descending order

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100
papers

2,395
citations

172207

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h-index

243296

44
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101
all docs

101
docs citations

101
times ranked

767
citing authors

#	ARTICLE	IF	CITATIONS
1	Full-dimensional Schrödinger wavefunction calculations using tensors and quantum computers: the Cartesian component-separated approach. <i>Physical Chemistry Chemical Physics</i> , 2022, 24, 4437-4454.	1.3	1
2	A Non-relativistic Approach to Relativistic Quantum Mechanics: The Case of the Harmonic Oscillator. <i>Foundations of Physics</i> , 2022, 52, 1.	0.6	7
3	Quantum Trajectory Description of the Time-Independent (Inverse) Fermi Accelerator. <i>Brazilian Journal of Physics</i> , 2021, 51, 193-203.	0.7	2
4	Calculation of rovibrational eigenstates of H3+ using ScallIT. <i>AIP Advances</i> , 2021, 11, 045033.	0.6	3
5	On neglecting Coriolis and related couplings in first-principles rovibrational spectroscopy: Considerations of symmetry, accuracy, and simplicity. II. Case studies for H2O isotopologues, O3, and NH3. <i>Spectrochimica Acta - Part A: Molecular and Biomolecular Spectroscopy</i> , 2021, 250, 119164.	2.0	9
6	Making relativistic quantum mechanics simple. <i>European Journal of Physics</i> , 2021, 42, 055404.	0.3	15
7	Hitting the Trifecta: How to Simultaneously Push the Limits of Schrödinger Solution with Respect to System Size, Convergence Accuracy, and Number of Computed States. <i>Journal of Chemical Theory and Computation</i> , 2021, 17, 7732-7744.	2.3	7
8	Plumbing Potentials for Molecules with Up To Tens of Atoms: How to Find Saddle Points and Fix Leaky Holes. <i>Journal of Physical Chemistry Letters</i> , 2020, 11, 6468-6474.	2.1	11
9	Trajectory-based conservation laws for massive spin-zero relativistic quantum particles in 1 + 1 spacetime. <i>Journal of Physics: Conference Series</i> , 2020, 1612, 012022.	0.3	0
10	An algorithm to find (and plug) "holes" in multi-dimensional surfaces. <i>Journal of Chemical Physics</i> , 2020, 152, 214102.	1.2	14
11	On neglecting Coriolis and related couplings in first-principles rovibrational spectroscopy: considerations of symmetry, accuracy, and simplicity. <i>Scientific Reports</i> , 2020, 10, 4872.	1.6	8
12	Comment on "Calculated vibrational states of ozone up to dissociation". <i>J. Chem. Phys.</i> 144, 074302 (2016). <i>Journal of Chemical Physics</i> , 2020, 152, 177101.	1.2	4
13	Using phase-space Gaussians to compute the vibrational states of OCHCO+. <i>Journal of Chemical Physics</i> , 2019, 151, 014114.	1.2	10
14	The J-dependent rotational Hamiltonian method for analyzing rovibrational spectra: Application to HO2, H2O, and O3. <i>Chemical Physics Letters</i> , 2019, 733, 136700.	1.2	3
15	Exact bound rovibrational spectra of the neon tetramer. <i>Journal of Chemical Physics</i> , 2019, 151, 174304.	1.2	9
16	Isotope shifts and band progressions in SO ₂ rovibrational energy levels: using quantum theory to extract rotational constants. <i>Molecular Physics</i> , 2019, 117, 2456-2469.	0.8	3
17	Accurate characterization of the lowest triplet potential energy surface of SO2 with a coupled cluster method. <i>Journal of Chemical Physics</i> , 2019, 150, 144303.	1.2	2
18	Exact matrix elements for general two-body central-force interactions, expressed as sums of products. <i>Molecular Physics</i> , 2019, 117, 1264-1275.	0.8	2

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19	Two-body Schrödinger wave functions in a plane-wave basis via separation of dimensions. Journal of Chemical Physics, 2018, 148, 104101.	1.2	10
20	Critical evaluation of measured rotational-vibrational transitions of four sulphur isotopologues of S16O2. Journal of Quantitative Spectroscopy and Radiative Transfer, 2018, 208, 152-163.	1.1	32
21	Accurate rovibrational energies of ozone isotopologues up to $J = 10$ utilizing artificial neural networks. Journal of Chemical Physics, 2018, 149, 024307.	1.2	17
22	Communication: Adiabatic quantum trajectory capture for cold and ultra-cold chemical reactions. Journal of Chemical Physics, 2018, 149, 021101.	1.2	8
23	A quantum dynamical study of the rotation of the dihydrogen ligand in the Fe(H)2(H2)(PEtPh2)3 coordination complex. Journal of Chemical Physics, 2018, 148, 154303.	1.2	4
24	Photoabsorption Assignments for the $\tilde{X}^1B_2 \leftarrow \tilde{X}^1A_1$ Vibronic Transitions of SO2, Using New Ab Initio Potential Energy and Transition Dipole Surfaces. Journal of Physical Chemistry A, 2017, 121, 1012-1021.	1.1	18
25	Final State Resolved Quantum Predissociation Dynamics of SO_2 ($\tilde{X}^1A_1 \leftarrow \tilde{X}^1B_2$) and Its Isotopomers via a Crossing with a Singlet Repulsive State. Journal of Physical Chemistry A, 2017, 121, 4930-4938.	1.1	9
26	First-principles C band absorption spectra of SO_2 and its isotopologues. Journal of Chemical Physics, 2017, 146, 154305.	1.2	7
27	Exploring the propagation of relativistic quantum wavepackets in the trajectory-based formulation. Journal of Physics: Conference Series, 2016, 701, 012013.	0.3	7
28	New <i>ab initio</i> adiabatic potential energy surfaces and bound state calculations for the singlet ground \tilde{X}^1A_1 and excited $\tilde{C}^1B_2(21A''^2)$ states of SO2. Journal of Chemical Physics, 2016, 144, 174301.	1.2	17
29	Publisher's Note: New <i>ab initio</i> adiabatic potential energy surfaces and bound state calculations for the singlet ground \tilde{X}^1A_1 and excited $\tilde{C}^1B_2(21A''^2)$ states of SO2 [J. Chem. Phys. 144, 174301 (2016)]. Journal of Chemical Physics, 2016, 144, 209901.		0
30	Rovibrational bound states of SO2 isotopologues. II: Total angular momentum $J = 11 \leftarrow 20$. Chemical Physics, 2015, 461, 34-46.	0.9	8
31	Communication: The H2@C60 inelastic neutron scattering selection rule: Expanded and explained. Journal of Chemical Physics, 2015, 143, 101104.	1.2	21
32	Development of new ferritic alloys reinforced by nano titanium nitrides. Journal of Nuclear Materials, 2015, 456, 449-454.	1.3	7
33	Rovibrational bound states of SO2 isotopologues. I: Total angular momentum $J = 0 \leftarrow 10$. Chemical Physics, 2015, 450-451, 59-73.	0.9	10
34	Large scale exact quantum dynamics calculations: Ten thousand quantum states of acetonitrile. Chemical Physics Letters, 2015, 624, 37-42.	1.2	34
35	One Million Quantum States of Benzene. Journal of Physical Chemistry A, 2015, 119, 12417-12433.	1.1	38
36	Accurate calculations of bound rovibrational states for argon trimer. Journal of Chemical Physics, 2014, 141, 034302.	1.2	13

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37	Calculation of exact vibrational spectra for P2O and CH2NH using a phase space wavelet basis. Journal of Chemical Physics, 2014, 140, 204112.	1.2	26
38	Comparison of J-shifting models for rovibrational spectra as applied to the HO2 molecule. Chemical Physics Letters, 2014, 605-606, 16-21.	1.2	10
39	Using ScalT for Performing Accurate Rovibrational Spectroscopy Calculations for Triatomic Molecules: A Practical Guide. Applied Mathematics, 2014, 05, 2756-2763.	0.1	13
40	Quantum Dynamical Calculation of Bound Rovibrational States of HO ₂ up to Largest Possible Total Angular Momentum, $J \leq 130$. Journal of Physical Chemistry A, 2013, 117, 7280-7297.	1.1	18
41	ROVIBRATIONAL BOUND STATES OF THE Ar_2Ne COMPLEX. Journal of Theoretical and Computational Chemistry, 2013, 12, 1250107.	1.8	6
42	Quantum dynamical calculation of rovibrational bound states of Ne ₂ Ar. Journal of Physics B: Atomic, Molecular and Optical Physics, 2012, 45, 135102.	0.6	10
43	Accurate quantum dynamics calculations using symmetrized Gaussians on a doubly dense Von Neumann lattice. Journal of Chemical Physics, 2012, 137, 224101.	1.2	30
44	Sech wave packets, their Wigner functions and Bohmian trajectories. Journal of Physics A: Mathematical and Theoretical, 2012, 45, 405302.	0.7	2
45	Communication: Quantum mechanics without wavefunctions. Journal of Chemical Physics, 2012, 136, 031102.	1.2	56
46	Classical-like trajectory simulations for accurate computation of quantum reactive scattering probabilities. Computational and Theoretical Chemistry, 2012, 990, 3-17.	1.1	23
47	An action principle for complex quantum trajectories. Molecular Physics, 2012, 110, 897-908.	0.8	5
48	Quantum dynamics of hydrogen interacting with single-walled carbon nanotubes: Multiple H-atom adsorbates. Journal of Chemical Physics, 2011, 134, 074308.	1.2	10
49	Bipolar Reaction Path Hamiltonian Approach for Reactive Scattering Problems. Journal of Chemical Theory and Computation, 2011, 7, 3484-3504.	2.3	11
50	Rovibrational bound states of neon trimer: Quantum dynamical calculation of all eigenstate energy levels and wavefunctions. Journal of Chemical Physics, 2011, 135, 094306.	1.2	21
51	Bohmian mechanics without pilot waves. Chemical Physics, 2010, 370, 4-14.	0.9	47
52	Parallel implementation of an efficient preconditioned linear solver for grid-based applications in chemical physics. III: Improved parallel scalability for sparse matrix-vector products. Journal of Parallel and Distributed Computing, 2010, 70, 779-782.	2.7	20
53	QUANTUM DYNAMICS ON MASSIVELY PARALLEL COMPUTERS: EFFICIENT NUMERICAL IMPLEMENTATION FOR PRECONDITIONED LINEAR SOLVERS AND EIGENSOLVERS. Journal of Theoretical and Computational Chemistry, 2010, 09, 825-846.	1.8	22
54	QUANTUM DYNAMICAL CALCULATION OF ALL ROVIBRATIONAL STATES OF HO ₂ FOR TOTAL ANGULAR MOMENTUM $J = 0 \leq 10$. Journal of Theoretical and Computational Chemistry, 2010, 09, 435-469.	1.8	19

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55	QUANTUM TRAJECTORY CALCULATIONS FOR BIPOLAR WAVEPACKET DYNAMICS IN ONE DIMENSION: SYNTHETIC SINGLE-WAVEPACKET PROPAGATION. <i>Journal of Theoretical and Computational Chemistry</i> , 2010, 09, 711-734.	1.8	11
56	Quantum dynamics of hydrogen interacting with single-walled carbon nanotubes. <i>Journal of Chemical Physics</i> , 2009, 130, 064701.	1.2	13
57	Reconciling semiclassical and Bohmian mechanics. VI. Multidimensional dynamics. <i>Journal of Chemical Physics</i> , 2008, 129, 084103.	1.2	30
58	Reconciling semiclassical and Bohmian mechanics. V. Wavepacket dynamics. <i>Journal of Chemical Physics</i> , 2008, 128, 164115.	1.2	34
59	Quantum trajectory calculations for bipolar wavepacket dynamics in one dimension. <i>Journal of Chemical Physics</i> , 2008, 129, 194112.	1.2	20
60	Flux continuity and probability conservation in complexified Bohmian mechanics. <i>Physical Review A</i> , 2008, 77, .	1.0	37
61	DEVELOPMENT AND NUMERICAL ANALYSIS OF "BLACK-BOX" COUNTERPROPAGATING WAVE ALGORITHM FOR EXACT QUANTUM SCATTERING CALCULATIONS. <i>Journal of Theoretical and Computational Chemistry</i> , 2007, 06, 99-125.	1.8	10
62	USING DISCRETE VARIABLE REPRESENTATION PATH INTEGRAL MONTE CARLO WITH METROPOLIS SAMPLING TO COMPUTE GROUND STATE WAVEFUNCTIONS. <i>Journal of Theoretical and Computational Chemistry</i> , 2007, 06, 309-321.	1.8	1
63	Reconciling Semiclassical and Bohmian Mechanics: IV. Multisurface Dynamics. <i>Journal of Physical Chemistry A</i> , 2007, 111, 10400-10408.	1.1	46
64	Fully Quantum Rovibrational Calculation of the He(H ₂) Bound and Resonance States. <i>Journal of Physical Chemistry A</i> , 2006, 110, 5475-5480.	1.1	10
65	Parallel implementation of efficient preconditioned linear solver for grid-based applications in chemical physics. II: QMR linear solver. <i>Journal of Computational Physics</i> , 2006, 219, 198-209.	1.9	37
66	Parallel implementation of efficient preconditioned linear solver for grid-based applications in chemical physics. I: Block Jacobi diagonalization. <i>Journal of Computational Physics</i> , 2006, 219, 185-197.	1.9	30
67	Reconciling semiclassical and Bohmian mechanics. III. Scattering states for continuous potentials. <i>Journal of Chemical Physics</i> , 2006, 124, 034116.	1.2	29
68	Reconciling semiclassical and Bohmian mechanics. II. Scattering states for discontinuous potentials. <i>Journal of Chemical Physics</i> , 2006, 124, 034115.	1.2	30
69	Improving the accuracy of Weyl-Heisenberg wavelet and symmetrized Gaussian representations using customized phase-space-region operators. <i>Physical Review E</i> , 2006, 74, 036705.	0.8	15
70	Rovibrational spectroscopy calculations of neon dimer using a phase space truncated Weyl-Heisenberg wavelet basis. <i>Journal of Chemical Physics</i> , 2006, 124, 144107.	1.2	13
71	PARALLEL SUBSPACE ITERATION METHOD FOR THE SPARSE SYMMETRIC EIGENVALUE PROBLEM. <i>Journal of Theoretical and Computational Chemistry</i> , 2006, 05, 801-818.	1.8	4
72	Analytical treatment of Coriolis coupling for three-body systems. <i>Chemical Physics</i> , 2005, 308, 305-315.	0.9	18

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73	Accurate, two-stateab initio study of the ground and first-excited states of He ₂ ⁺ , including exact treatment of all Bornâ€“Oppenheimer correction terms. Journal of Chemical Physics, 2005, 122, 184310.	1.2	35
74	Accurate quantum calculation of the bound and resonant rovibrational states of Liâˆ“(H ₂). Journal of Chemical Physics, 2005, 122, 124318.	1.2	13
75	A Quantum Dynamical Treatment of Symmetry-Induced Kinetic Isotope Effects in the Formation of He ₂ ⁺ . Journal of the American Chemical Society, 2005, 127, 16969-16975.	6.6	3
76	Multidimensional quantum trajectories: Applications of the derivative propagation method. Journal of Chemical Physics, 2005, 122, 164104.	1.2	39
77	Quantum dynamics calculations using symmetrized, orthogonal Weyl-Heisenberg wavelets with a phase space truncation scheme. III. Representations and calculations. Journal of Chemical Physics, 2004, 121, 1704-1724.	1.2	36
78	Accurate and highly efficient calculation of the highly excited pure OH stretching resonances of O(1D)HCl, using a combination of methods. Journal of Chemical Physics, 2004, 121, 4467-4478.	1.2	31
79	Reconciling semiclassical and Bohmian mechanics. I. Stationary states. Journal of Chemical Physics, 2004, 121, 4501-4515.	1.2	83
80	Quantum dynamics calculations using symmetrized, orthogonal Weyl-Heisenberg wavelets with a phase space truncation scheme. II. Construction and optimization. Journal of Chemical Physics, 2004, 121, 1690-1703.	1.2	43
81	Exploiting both C _{3v} symmetry and sparsity in vibrational calculations for methanelike molecules. Journal of Chemical Physics, 2003, 119, 90-93.	1.2	9
82	Semiclassically optimized complex absorbing potentials of polynomial form. II. Complex case. Journal of Chemical Physics, 2003, 119, 77-89.	1.2	35
83	A quantum dynamical study of the He ⁺⁺ +2Heâ†’He ₂ ⁺⁺ +He reaction. Journal of Chemical Physics, 2003, 119, 10678-10686.	1.2	14
84	Eigenspectra calculations using Cartesian coordinates and a rotational symmetry adapted Lanczos method. Journal of Chemical Physics, 2003, 119, 6609-6619.	1.2	15
85	Semiclassically optimized complex absorbing potentials of polynomial form. I. Pure imaginary case. Journal of Chemical Physics, 2003, 118, 17-28.	1.2	57
86	USING WAVELETS TO EXTEND QUANTUM DYNAMICS CALCULATIONS TO TEN OR MORE DEGREES OF FREEDOM. Journal of Theoretical and Computational Chemistry, 2003, 02, 65-72.	1.8	71
87	ACCURATE AND HIGHLY EFFICIENT CALCULATION OF THE O(1D)HCl VIBRATIONAL BOUND STATES, USING A COMBINATION OF METHODS. Journal of Theoretical and Computational Chemistry, 2003, 02, 583-597.	1.8	47
88	A general framework for discrete variable representation basis sets. Journal of Chemical Physics, 2002, 116, 8691-8703.	1.2	129
89	A preconditioned inexact spectral transform method for calculating resonance energies and widths, as applied to HCO. Journal of Chemical Physics, 2002, 116, 1215-1227.	1.2	74
90	Phase Space Optimization of Quantum Representations: Non-Cartesian Coordinate Spaces. Foundations of Physics, 2001, 31, 1581-1610.	0.6	27

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91	Accelerating the calculation of energy levels and wave functions using an efficient preconditioner with the inexact spectral transform method. <i>Journal of Chemical Physics</i> , 2001, 114, 9254-9264.	1.2	78
92	Phase space optimization of quantum representations: Three-body systems and the bound states of HCO. <i>Journal of Chemical Physics</i> , 2001, 114, 6562-6571.	1.2	45
93	Efficient preconditioning scheme for block partitioned matrices with structured sparsity. <i>Numerical Linear Algebra With Applications</i> , 2000, 7, 715-726.	0.9	82
94	Algebraically Self-Consistent Quasiclassical Approximation on Phase Space. <i>Foundations of Physics</i> , 2000, 30, 1191-1226.	0.6	26
95	Efficient distributed Gaussian basis for rovibrational spectroscopy calculations. <i>Journal of Chemical Physics</i> , 2000, 113, 211-217.	1.2	54
96	Wigner-Weyl correspondence and semiclassical quantization in spherical coordinates. <i>Journal of Mathematical Physics</i> , 1999, 40, 6302-6318.	0.5	5
97	Phase space optimization of quantum representations: Direct-product basis sets. <i>Journal of Chemical Physics</i> , 1999, 111, 4869-4885.	1.2	79
98	Quantum reactive scattering for three-body systems via optimized preconditioning, as applied to the O+HCl reaction. <i>Journal of Chemical Physics</i> , 1998, 108, 5216-5224.	1.2	70
99	Optimal separable bases and series expansions. <i>Physical Review A</i> , 1997, 56, 120-130.	1.0	42
100	Optimized preconditioners for Green function evaluation in quantum reactive scattering calculations. <i>Chemical Physics Letters</i> , 1997, 265, 77-83.	1.2	43