

# Edward F C Byrd

## List of Publications by Year in descending order

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64  
papers

6,363  
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186265  
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docs citations

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times ranked

5603  
citing authors

#	ARTICLE	IF	CITATIONS
1	1,5-Diaminotetrazole-4 <i>i</i> >N <i>&lt;/i&gt;-oxide (SYX-9): a new high-performing energetic material with a calculated detonation velocity over 10 km s<sup>â˜’1</sup>. Journal of Materials Chemistry A, 2022, 10, 1876-1884.</i>	10.3	21
2	General quantitative structureâ€“property relationships and machine learning correlations to energetic material sensitivities. Theoretical and Computational Chemistry, 2022, , 139-156.	0.4	1
3	Energetic triazinium salts from N-amination of 3,5-diamino-6-nitro-1,2,4-triazine. Energetic Materials Frontiers, 2022, 3, 128-136.	3.2	3
4	3-Methyl-1,2,3-triazolium-1N-dinitromethyllylide and the strategy of zwitterionic dinitromethyl groups in energetic materials design. RSC Advances, 2021, 11, 17710-17714.	3.6	3
5	Methyl sydnone imine and its energetic salts. New Journal of Chemistry, 2021, 45, 2228-2236.	2.8	6
6	Synthesis and Characterization of the Energetic 3â€“Azidoâ€“5â€“aminoâ€“6â€“nitroâ€“1,2,4â€“triazine. Propellants, Explosives, Pyrotechnics, 2021, 46, 214-221.	1.6	11
7	Heterocyclic Nitrilimines and Their Use in the Synthesis of Complex High-Nitrogen Materials. Inorganic Chemistry, 2021, 60, 7607-7611.	4.0	12
8	Synthesis and Characterization of the Potential Meltâ€“Castable Explosive 3â€“(1,2,4â€“Oxadiazolyl)â€“5â€“Nitratomethyl Isoxazole. ChemPlusChem, 2021, 86, 875-878.	2.8	5
9	1,3,4,5-Tetraamino-1,2,4-triazolium Cation: An Energetic Moiety. Inorganic Chemistry, 2021, 60, 9645-9652.	4.0	9
10	4,4â€“Dinitrimino-5,5â€“diamino-3,3â€“azo-bis-1,2,4-triazole: A High-Performing Zwitterionic Energetic Material. Inorganic Chemistry, 2021, 60, 16204-16212.	4.0	9
11	Heuristics for chemical species identification in dense systems. Journal of Chemical Physics, 2020, 153, 064102.	3.0	8
12	4,4â€“5,5â€“Tetraamino-3,3â€“azo-bis-1,2,4-triazole and the electrosynthesis of high-performing insensitive energetic materials. Journal of Materials Chemistry A, 2020, 8, 19337-19347.	10.3	43
13	Tailoring Energetic Sensitivity and Classification through Regiosomerism. Organic Letters, 2020, 22, 9114-9117.	4.6	29
14	Sensitive Energetics from the <i>N</i>â€“Amination of 4â€“Nitroâ€“1,2,3â€“Triazole. ChemistryOpen, 2020, 9, 806-8119	11	
15	Tetrazole Azasydnone ( $C_{2}N_7O_2H$ ) And Its Salts: Highâ€“Performing Zwitterionic Energetic Materials Containing A Unique Explosophore. Chemistry - A European Journal, 2020, 26, 14530-14535.	3.3	53
16	Synthesis of Erythritol Tetranitrate Derivatives: Functional Group Tuning of Explosive Sensitivity. Journal of Organic Chemistry, 2020, 85, 4619-4626.	3.2	28
17	Bis(Nitroxymethylisoxazolyl) Furoxan: A Promising Standalone Meltâ€“Castable Explosive. ChemPlusChem, 2020, 85, 237-239.	2.8	34
18	Impact of Stereo- and Regiochemistry on Energetic Materials. Journal of the American Chemical Society, 2019, 141, 12531-12535.	13.7	92

#	ARTICLE	IF	CITATIONS
19	Synthesis and Characterization of Salts of the 3,6-dinitro-[1,2,4]triazolo[4,3- <i>b</i> ][1,2,4]triazolate Anion: In-sensitive Energetic Materials Available From Economical Precursors. <i>Zeitschrift Fur Anorganische Und Allgemeine Chemie</i> , 2019, 645, 1197-1204.	1.2	16
20	Bis(1,2,4-oxadiazolyl) Furoxan: A Promising Melt-Castable Eutectic Material of Low Sensitivity. <i>ChemPlusChem</i> , 2019, 84, 319-322.	2.8	27
21	Toward a Predictive Hierarchical Multiscale Modeling Approach for Energetic Materials. <i>Challenges and Advances in Computational Chemistry and Physics</i> , 2019, , 229-282.	0.6	16
22	Ray tracing calculations in simulated propellant flames with detailed chemistry. <i>Applied Optics</i> , 2019, 58, 1451.	1.8	2
23	Effect of a core-softened O-O interatomic interaction on the shock compression of fused silica. <i>Journal of Chemical Physics</i> , 2018, 148, .	3.0	3
24	Density Functional Theory and Experimental Studies of the Molecular, Vibrational, and Crystal Structure of Bis-Oxadiazole-Bis-Methylene Dinitrate (BODN). <i>Journal of Physical Chemistry A</i> , 2018, 122, 9043-9053.	2.5	9
25	Challenges and opportunities in integration of 2D materials on 3D substrates: Materials and device perspectives. , 2018, , .		2
26	Bis(1,2,4-oxadiazole)bis(methylene) Dinitrate: A High-Energy Melt-Castable Explosive and Energetic Propellant Plasticizing Ingredient. <i>Organic Process Research and Development</i> , 2018, 22, 736-740.	2.7	90
27	Harmonic Vibrational Frequencies: Approximate Global Scaling Factors for TPSS, M06, and M11 Functional Families Using Several Common Basis Sets. <i>Journal of Physical Chemistry A</i> , 2017, 121, 2265-2273.	2.5	141
28	Synthesis of Biisoxazolotetrakis(methyl nitrate): A Potential Nitrate Plasticizer and Highly Explosive Material. <i>European Journal of Organic Chemistry</i> , 2017, 2017, 1765-1768.	2.4	29
29	Simple and Efficient Synthesis of Explosive Cocrystals containing 3,5-dimethylpyrazol-1-yl-substituted-1,2,4,5-tetrazines. <i>Chemistry - A European Journal</i> , 2017, 23, 16466-16471. <sup>2,3</sup>		21
30	Structural and electrical analysis of epitaxial 2D/3D vertical heterojunctions of monolayer MoS <sub>2</sub> on GaN. <i>Applied Physics Letters</i> , 2017, 111, .	3.3	27
31	Synthesis of bis-isoxazole-bis-methylene Dinitrate: A Potential Nitrate Plasticizer and Melt-Castable Energetic Material. <i>ChemPlusChem</i> , 2017, 82, 195-198.	2.8	37
32	Theoretical Study of Shocked Formic Acid: Born-Oppenheimer MD Calculations of the Shock Hugoniot and Early-Stage Chemistry. <i>Journal of Physical Chemistry B</i> , 2016, 120, 1711-1719.	2.6	5
33	Parameterizing Complex Reactive Force Fields Using Multiple Objective Evolutionary Strategies (MOES): Part 2: Transferability of ReaxFF Models to H-N-O Energetic Materials. <i>Journal of Chemical Theory and Computation</i> , 2015, 11, 392-405.	5.3	36
34	Parameterizing Complex Reactive Force Fields Using Multiple Objective Evolutionary Strategies (MOES). Part 1: ReaxFF Models for Cyclotrimethylene Trinitramine (RDX) and 1,1-Diamino-2,2-dinitroethene (FOX-7). <i>Journal of Chemical Theory and Computation</i> , 2015, 11, 381-391.	5.3	36
35	Development of quantitative structure property relationships for predicting the melting point of energetic materials. <i>Journal of Molecular Graphics and Modelling</i> , 2015, 62, 190-201.	2.4	11
36	Special quasirandom structures of alon. <i>Computational Materials Science</i> , 2015, 96, 312-318.	3.0	22

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37	Assessing the Performances of Dispersion-Corrected Density Functional Methods for Predicting the Crystallographic Properties of High Nitrogen Energetic Salts. <i>Journal of Chemical Theory and Computation</i> , 2014, 10, 4982-4994.	5.3	22
38	Evaluation of electrostatic descriptors for predicting crystalline density. <i>Journal of Computational Chemistry</i> , 2013, 34, 2146-2151.	3.3	112
39	Shock Hugoniot calculations of polymers using quantum mechanics and molecular dynamics. <i>Journal of Chemical Physics</i> , 2012, 137, 204901.	3.0	46
40	Assessment of Dispersion Corrected Atom Centered Pseudopotentials: Application to Energetic Molecular Crystals. <i>Journal of Physical Chemistry B</i> , 2011, 115, 803-810.	2.6	32
41	A Comparison of Methods To Predict Solid Phase Heats of Formation of Molecular Energetic Salts. <i>Journal of Physical Chemistry A</i> , 2009, 113, 345-352.	2.5	64
42	An Investigation of KS-DFT Electron Densities used in Atoms-in-Molecules Studies of Energetic Molecules. <i>Journal of Physical Chemistry A</i> , 2009, 113, 6166-6171.	2.5	38
43	Development of quantitative structure–property relationships for predictive modeling and design of energetic materials. <i>Journal of Molecular Graphics and Modelling</i> , 2008, 27, 349-355.	2.4	38
44	Computational Aspects of Nitrogen-Rich HEDMs. , 2007, , 153-194.		48
45	Accurate Predictions of Crystal Densities Using Quantum Mechanical Molecular Volumes. <i>Journal of Physical Chemistry A</i> , 2007, 111, 10874-10879.	2.5	175
46	Ab Initio Study of Compressed 1,3,5,7-Tetranitro-1,3,5,7-tetraazacyclooctane (HMX), Cyclotrimethylenetrinitramine (RDX), 2,4,6,8,10,12-Hexanitrohexaazaisowurzitane (CL-20), 2,4,6-Trinitro-1,3,5-benzenetriamine (TATB), and Pentaerythritol Tetranitrate (PETN). <i>Journal of Physical Chemistry C</i> , 2007, 111, 2787-2796.	3.1	150
47	Advances in methods and algorithms in a modern quantum chemistry program package. <i>Physical Chemistry Chemical Physics</i> , 2006, 8, 3172-3191.	2.8	2,597
48	Improved Prediction of Heats of Formation of Energetic Materials Using Quantum Mechanical Calculations. <i>Journal of Physical Chemistry A</i> , 2006, 110, 1005-1013.	2.5	494
49	Theoretical chemical characterization of energetic materials. <i>Journal of Materials Research</i> , 2006, 21, 2444-2452.	2.6	35
50	Theoretical Chemical Characterization of Energetic Materials. <i>Materials Research Society Symposia Proceedings</i> , 2005, 896, 11.	0.1	1
51	An ab Initio Study of Solid Nitromethane, HMX, RDX, and CL20: Successes and Failures of DFT. <i>Journal of Physical Chemistry B</i> , 2004, 108, 13100-13106.	2.6	122
52	Quadratic Coupled-Cluster Doubles: Implementation and Assessment of Perfect Pairing Optimized Geometries. <i>Journal of Physical Chemistry B</i> , 2002, 106, 8070-8077.	2.6	22
53	Coupled Cluster Methods for Bond-Breaking. <i>ACS Symposium Series</i> , 2002, , 93-108.	0.5	6
54	A perturbative correction to the quadratic coupled-cluster doubles method for higher excitations. <i>Chemical Physics Letters</i> , 2002, 353, 359-367.	2.6	79

#	ARTICLE		IF	CITATIONS
55	The Theoretical Prediction of Molecular Radical Species: a Systematic Study of Equilibrium Geometries and Harmonic Vibrational Frequencies. <i>Journal of Physical Chemistry A</i> , 2001, 105, 9736-9747.		2.5	142
56	Q-Chem 2.0: a high-performance ab initio electronic structure program package. <i>Journal of Computational Chemistry</i> , 2000, 21, 1532-1548.		3.3	617
57	Complete basis set extrapolations for low-lying triplet electronic states of acetylene and vinylidene. <i>Journal of Chemical Physics</i> , 2000, 113, 1447-1454.		3.0	24
58	Q-Chem 2.0: a high-performance ab initio electronic structure program package. , 2000, 21, 1532.			2
59	Energies and analytic gradients for a coupled-cluster doubles model using variational Brueckner orbitals: Application to symmetry breaking in O <sub>4+</sub> . <i>Journal of Chemical Physics</i> , 1998, 109, 4171-4181.		3.0	228
60	Size-consistent wave functions for nondynamical correlation energy: The valence active space optimized orbital coupled-cluster doubles model. <i>Journal of Chemical Physics</i> , 1998, 109, 10669-10678.		3.0	222
61	The inclusion of correlation in the calculation of phosphorus NMR chemical shieldings. <i>Heteroatom Chemistry</i> , 1996, 7, 307-312.		0.7	20
62	The use of locally dense basis sets in correlated NMR chemical shielding calculations. <i>Chemical Physics</i> , 1996, 213, 153-158.		1.9	65
63	Title is missing!. <i>Journal of Computational Chemistry</i> , 1996, 17, 1431.		3.3	5
64	Titanium Superoxide for the Oxidation of Amines: Synthesis of bis(3-nitro-1H-1,2,4-triazol-5-yl)methane and its Metal Salts. <i>Zeitschrift Fur Anorganische Und Allgemeine Chemie</i> , 0, .		1.2	2