## Mel Levy

## List of Publications by Year in Descending Order

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The third column is the impact factor (IF) of the journal, and the fourth column is the number of citations of the article.

77	11,671	31	80
papers	citations	h-index	g-index
80	12,428 ext. citations	3.2	6.02
ext. papers		avg, IF	L-index

#	Paper Paper	IF	Citations
77	Simple hydrogenic estimates for the exchange and correlation energies of atoms and atomic ions, with implications for density functional theory. <i>Journal of Chemical Physics</i> , <b>2020</b> , 153, 074114	3.9	3
76	Approximating the Shifted Hartree-Exchange-Correlation Potential in Direct Energy Kohn-Sham Theory. <i>Journal of Chemical Theory and Computation</i> , <b>2018</b> , 14, 684-692	6.4	3
75	Nodal variational principle for excited states. <i>Physical Review A</i> , <b>2018</b> , 98,	2.6	4
74	On the best partitioning of the density functional energy. <i>Journal of Molecular Modeling</i> , <b>2018</b> , 24, 311	2	0
73	Properties of Augmented Kohn-Sham Potential for Energy as Simple Sum of Orbital Energies. Journal of Physical Chemistry A, <b>2017</b> , 121, 342-347	2.8	2
7 <sup>2</sup>	Augmented potential, energy densities, and virial relations in the weak- and strong-interaction limits of DFT. <i>Journal of Chemical Physics</i> , <b>2017</b> , 147, 214107	3.9	14
71	On augmented KohnBham potential for energy as a simple sum of orbital energies. <i>Molecular Physics</i> , <b>2016</b> , 114, 1162-1164	1.7	5
70	Highly Excited States from a Time Independent Density Functional Method. Computation, 2016, 4, 28	2.2	6
69	Mathematical thoughts in DFT. International Journal of Quantum Chemistry, 2016, 116, 802-804	2.1	10
68	Generalized Variational Theorem in Quantum Mechanics <b>2015</b> , 92-97		
67	Ground-state energy as a simple sum of orbital energies in Kohn-Sham theory: a shift in perspective through a shift in potential. <i>Physical Review Letters</i> , <b>2014</b> , 113, 113002	7.4	30
66	Kinetic and electron-electron energies for convex sums of ground state densities with degeneracies and fractional electron number. <i>Journal of Chemical Physics</i> , <b>2014</b> , 140, 18A538	3.9	9
65	Tight constraints on the exchange-correlation potentials of degenerate states. <i>Journal of Chemical Physics</i> , <b>2014</b> , 140, 18A537	3.9	5
64	Time-independent density-functional theory for excited states of Coulomb systems. <i>Physical Review A</i> , <b>2012</b> , 85,	2.6	41
63	On the simple constrained-search reformulation of the Hohenbergkohn theorem to include degeneracies and more (1964¶979). <i>International Journal of Quantum Chemistry</i> , <b>2010</b> , 110, 3140-3144	2.1	20
62	Comment on <b>H</b> unctional derivative of the universal density functional in Fock space <i>Physical Review A</i> , <b>2009</b> , 79,	2.6	16
61	Kinetic energy from a single Kohn-Sham orbital. <i>Physical Review A</i> , <b>2009</b> , 79,	2.6	9

60 Pointwise and generalized virial theorems. *International Journal of Quantum Chemistry*, **2009**, 14, 343-34<u>4</u>.1

59	Properties of the exact universal density and one-matrix functionals. <i>International Journal of Quantum Chemistry</i> , <b>2009</b> , 28, 743-744	2.1	
58	Asymptotic coordinate scaling bound for exchange-correlation energy in density-functional theory. <i>International Journal of Quantum Chemistry</i> , <b>2009</b> , 36, 617-619	2.1	12
57	Average local ionization energies in the Hartree-Fock and Kohn-Sham theories. <i>Journal of Physical Chemistry A</i> , <b>2009</b> , 113, 1384-9	2.8	29
56	Time-independent (static) density-functional theories for pure excited states: Extensions and unification. <i>Physical Review A</i> , <b>2009</b> , 80,	2.6	40
55	Formal expressions and corresponding expansions for the exact Kohn-Sham exchange potential. <i>Physical Review A</i> , <b>2009</b> , 80,	2.6	10
54	Line-integral formulas for exchange and correlation potentials separately. <i>World Scientific Series in 20th Century Physics</i> , <b>2009</b> , 613-617	Ο	
53	On the adiabatic connection method, and scaling of electron lectron interactions in the Thomas Hermi limit. World Scientific Series in 20th Century Physics, 2009, 542-545	Ο	
52	Relation between exchange-only optimized potential and Kohn-Sham methods with finite basis sets, and effect of linearly dependent products of orbital basis functions. <i>Journal of Chemical Physics</i> , <b>2008</b> , 128, 104104	3.9	57
51	Generalized density-functional theory: Conquering theN-representability problem with exact functionals for the electron pair density and the second-order reduced density matrix. <i>Journal of Chemical Sciences</i> , <b>2005</b> , 117, 507-514	1.8	70
50	Connections between ground-state energies from optimized-effective potential exchange-only and Hartreeflock methods. <i>Journal of Chemical Physics</i> , <b>2003</b> , 119, 7087-7093	3.9	26
49	Accurate correlation potentials from integral formulation of density functional perturbation theory. <i>Journal of Chemical Physics</i> , <b>2002</b> , 116, 6924-6929	3.9	23
48	Sum rules for exchange and correlation potentials. <i>Journal of Chemical Physics</i> , <b>2001</b> , 115, 4438-4443	3.9	35
47	Perspective on <b>D</b> ensity functional approach to the frontier-electron theory of chemical reactivity <i>Theoretical Chemistry Accounts</i> , <b>2000</b> , 103, 353-360	1.9	374
46	Strictly correlated electrons in density-functional theory. <i>Physical Review A</i> , <b>1999</b> , 59, 51-54	2.6	123
45	Exact high-density limit of correlation potential for two-electron density. <i>Journal of Chemical Physics</i> , <b>1999</b> , 110, 10262-10268	3.9	49
44	Adiabatic integration formula for the correlation energy functional of the Hartreeflock density. <i>Theoretical Chemistry Accounts</i> , <b>1999</b> , 103, 117-123	1.9	6
43	Variational Density-Functional Theory for an Individual Excited State. <i>Physical Review Letters</i> , <b>1999</b> , 83, 4361-4364	7.4	167

42	Closed-form expression relating the second-order component of the density functional theory correlation energy to its functional derivative. <i>Journal of Chemical Physics</i> , <b>1998</b> , 109, 6280-6286	3.9	7
41	Connections between High-Density Scaling Limits of DFT Correlation Energies and Second-Order Z-1 Quantum Chemistry Correlation Energy. <i>Journal of Physical Chemistry A</i> , <b>1998</b> , 102, 3151-3156	2.8	18
40	Line-integral formulas for exchange and correlation potentials separately. <i>Physical Review A</i> , <b>1997</b> , 55, 1885-1889	2.6	47
39	Hybrid schemes combining the HartreeBock method and density-functional theory: Underlying formalism and properties of correlation functionals. <i>Journal of Chemical Physics</i> , <b>1997</b> , 106, 2675-2680	3.9	85
38	Additive density functional correlation corrections to single particle theories. <i>International Journal of Quantum Chemistry</i> , <b>1997</b> , 61, 281-285	2.1	3
37	Approach to density-functional ionization energy. <i>Physical Review B</i> , <b>1996</b> , 53, 969-972	3.3	10
36	Exact local exchange potential from Fock equations at vanishing coupling constant, and delta Tc/delta n from wave-function calculations at full coupling constant. <i>Physical Review A</i> , <b>1996</b> , 53, 3963-396	5 <sup>2.6</sup>	8
35	Density-functional exchange identity from coordinate scaling. <i>Physical Review A</i> , <b>1996</b> , 53, 3140-3142	2.6	40
34	Improving energies by using exact electron densities. <i>Physical Review A</i> , <b>1996</b> , 53, R2915-R2917	2.6	18
33	New exact relations for improving the exchange and correlation potentials. <i>International Journal of Quantum Chemistry</i> , <b>1995</b> , 56, 385-388	2.1	8
32	DFT ionization formulas and a DFT perturbation theory for exchange and correlation, through adiabatic connection. <i>International Journal of Quantum Chemistry</i> , <b>1995</b> , 56, 93-108	2.1	136
31	Correlation-energy density-functional formulas from correlating first-order density matrices. <i>Physical Review A</i> , <b>1995</b> , 52, R1808-R1810	2.6	70
30	Bounds for the exchange and correlation potentials. <i>Physical Review A</i> , <b>1995</b> , 51, 2851-2856	2.6	15
29	Recent constrained-search advances for approximating density functionals. <i>The Philosophical Magazine: Physics of Condensed Matter B, Statistical Mechanics, Electronic, Optical and Magnetic Properties</i> , <b>1994</b> , 69, 763-769		29
28	Exact Kohn-Sham scheme based on perturbation theory. <i>Physical Review A</i> , <b>1994</b> , 50, 196-204	2.6	468
27	Density functionals for exchange and correlation energies: Exact conditions and comparison of approximations. <i>International Journal of Quantum Chemistry</i> , <b>1994</b> , 49, 539-548	2.1	53
26	Correlation-energy functional and its high-density limit obtained from a coupling-constant perturbation expansion. <i>Physical Review B</i> , <b>1993</b> , 47, 13105-13113	3.3	354
25	Expectation values in density-functional theory, and kinetic contribution to the exchange-correlation energy. <i>Physical Review B</i> , <b>1993</b> , 47, 1167-1173	3.3	45

## (1980-1991)

24	Approximate noninteracting kinetic energy functionals from a nonuniform scaling requirement. <i>International Journal of Quantum Chemistry</i> , <b>1991</b> , 40, 379-388	2.1	46
23	Density-functional exchange correlation through coordinate scaling in adiabatic connection and correlation hole. <i>Physical Review A</i> , <b>1991</b> , 43, 4637-4646	2.6	241
22	THEOREMS FOR EXACT LOCAL EXCHANGE POTENTIAL. <i>Modern Physics Letters B</i> , <b>1991</b> , 05, 1613-1616	1.6	
21	Electron density-functional theory and x-ray structure factors. <i>Physical Review B</i> , <b>1987</b> , 35, 7887-7890	3.3	26
20	Energy differences from electrostatic potentials at nuclei. <i>Journal of Chemical Physics</i> , <b>1987</b> , 87, 5044-5	0 <u>4</u> 6	10
19	Success of quantum mechanical approximations for molecular geometries and electronfluclear attraction expectation values: Gift of the Coulomb potential?. <i>Journal of Chemical Physics</i> , <b>1986</b> , 84, 45	19-452	3 <sup>30</sup>
18	A new functional with homogeneous coordinate scaling in density functional theory: F [ 即 <i>Journal of Chemical Physics</i> , <b>1985</b> , 83, 2334-2336	3.9	44
17	Hellmann-Feynman, virial, and scaling requisites for the exact universal density functionals. Shape of the correlation potential and diamagnetic susceptibility for atoms. <i>Physical Review A</i> , <b>1985</b> , 32, 2010	-2021	814
16	Exact differential equation for the density and ionization energy of a many-particle system. <i>Physical Review A</i> , <b>1984</b> , 30, 2745-2748	2.6	713
15	Physical Content of the Exact Kohn-Sham Orbital Energies: Band Gaps and Derivative Discontinuities. <i>Physical Review Letters</i> , <b>1983</b> , 51, 1884-1887	7.4	1795
14	Direct first principles algorithm for the universal electron density functional. <i>Journal of Chemical Physics</i> , <b>1982</b> , 77, 396-398	3.9	47
13	A discontinuous energydensity functional. <i>Journal of Chemical Physics</i> , <b>1982</b> , 77, 3140-3147	3.9	27
12	Electron densities in search of Hamiltonians. <i>Physical Review A</i> , <b>1982</b> , 26, 1200-1208	2.6	502
11	Density-Functional Theory for Fractional Particle Number: Derivative Discontinuities of the Energy. <i>Physical Review Letters</i> , <b>1982</b> , 49, 1691-1694	7.4	2236
10	Energy-density relations and screening constants in atoms. <i>Journal of Chemical Physics</i> , <b>1980</b> , 73, 5168-	51,73	21
9	Atomic binding energies from fundamental theorems involving the electron density, <rul> <li>zll perturbation expansion. <i>Journal of Chemical Physics</i>, 1980, 72, 3416-3417</li> </rul>	3.9	35
8	The nearest self-adjoint operator. <i>Journal of Chemical Physics</i> , <b>1980</b> , 72, 780-781	3.9	10
7	Rigorous and approximate relations between expectation values of atoms. <i>Journal of Chemical Physics</i> , <b>1980</b> , 72, 4009-4013	3.9	34

6	On approximate energy differences from average electron densities. <i>Journal of Chemical Physics</i> , <b>1979</b> , 70, 1573-1574	3.9	22
5	Electronegativity: The density functional viewpoint. <i>Journal of Chemical Physics</i> , <b>1978</b> , 68, 3801-3807	3.9	2329
4	An energy-density equation for isoelectronic changes in atoms. <i>Journal of Chemical Physics</i> , <b>1978</b> , 68, 5298-5299	3.9	36
3	Variational energy functionals involving one-electron operators. <i>Journal of Chemical Physics</i> , <b>1977</b> , 67, 724-726	3.9	6
2	Unconstrained exchange localization and distant orbital tails. <i>Journal of Chemical Physics</i> , <b>1976</b> , 65, 24	7332947	5 11
1	Method for direct determination of localized orbitals. <i>Journal of Chemical Physics</i> , <b>1975</b> , 63, 316-318	3.9	23