Miroslav Urban

List of Publications by Year in descending order

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| | | 933447 | 794594 |
|----------|----------------|--------------|----------------|
| 20 | 1,195 | 10 | 19 |
| papers | citations | h-index | g-index |
| | | | |
| | | | |
| 21 | 21 | 21 | 960 |
| all docs | docs citations | times ranked | citing authors |
| | | | |

| # | Article | IF | CITATIONS |
|----|---|-----|-----------|
| 1 | Towards a full CCSDT model for electron correlation. Journal of Chemical Physics, 1985, 83, 4041-4046. | 3.0 | 843 |
| 2 | Relativistic effects in atomic and molecular properties. Acta Physica Slovaca, 2010, 60, . | 1.4 | 58 |
| 3 | Electron affinity of the O ₂ molecule: CCSD(T) calculations using the optimized virtual orbitals space approach. International Journal of Quantum Chemistry, 2008, 108, 2159-2171. | 2.0 | 37 |
| 4 | Optimized virtual orbitals for relativistic calculations: an alternative approach to the basis set construction for correlation calculations. Molecular Physics, 2006, 104, 2277-2292. | 1.7 | 34 |
| 5 | Toward Understanding the Bonding Character in Complexes of Coinage Metals with Lone-Pair Ligands. CCSD(T) and DFT Computations. Journal of Physical Chemistry A, 2013, 117, 4472-4485. | 2.5 | 34 |
| 6 | Theoretical study of molecular properties of low-lying electronic excited states of H ₂ O and H ₂ S. Molecular Physics, 2008, 106, 2333-2344. | 1.7 | 30 |
| 7 | On the accuracy of molecular properties by coupled-cluster methods for some difficult examples: Oxygen atom, iron atom, and cyano radical. International Journal of Quantum Chemistry, 1994, 52, 211-225. | 2.0 | 28 |
| 8 | CASPT2 and CCSD(T) calculations of dipole moments and polarizabilities of acetone in excited states. Molecular Physics, 2012, 110, 2219-2237. | 1.7 | 24 |
| 9 | Accurate electrical and spectroscopic properties ofX 1?+ BeO from coupled-cluster methods. Theoretica Chimica Acta, 1995, 90, 341-355. | 0.8 | 21 |
| 10 | Au _{<i>n</i>} (<i>n</i> = 1,11) Clusters Interacting With Lone-Pair Ligands. Journal of Physical Chemistry A, 2016, 120, 3938-3949. | 2.5 | 18 |
| 11 | Density functional theory modeling of C–Au chemical bond formation in gold implanted polyethylene. Physical Chemistry Chemical Physics, 2017, 19, 28897-28906. | 2.8 | 12 |
| 12 | Highly correlated calculations using optimized virtual orbital space with controlled accuracy. Application to counterpoise corrected interaction energy calculations. International Journal of Quantum Chemistry, 2012, 112, 948-959. | 2.0 | 9 |
| 13 | Benchmark CCSD(T) and DFT study of binding energies in Be _{7 â^ 12} : in search of reliable DFT functional for beryllium clusters. Molecular Physics, 2018, 116, 1259-1274. | 1.7 | 9 |
| 14 | A comparative DFT study of interactions of Au and small gold clusters Aun (n = 2–4) with CH3S and CH2 radicals. Chemical Physics Letters, 2017, 671, 84-91. | 2.6 | 8 |
| 15 | DFT Modeling of Cross-Linked Polyethylene: Role of Gold Atoms and Dispersion Interactions. Journal of Physical Chemistry A, 2018, 122, 1496-1503. | 2.5 | 8 |
| 16 | DFT Functionals for Modeling of Polyethylene Chains Cross-Linked by Metal Atoms. DLPNO–CCSD(T) Benchmark Calculations. Journal of Physical Chemistry A, 2021, 125, 7382-7395. | 2.5 | 8 |
| 17 | Electric properties of hydrogen iodide: Reexamination of correlation and relativistic effects. Theoretical Chemistry Accounts, 2003, 110, 176-184. | 1.4 | 7 |
| 18 | Relativistic effects in low-lying electronic states of iron. Theoretical Chemistry Accounts, 2011, 129, 561-566. | 1.4 | 4 |

| # | Article | IF | CITATIONS |
|----|--|-----|-----------|
| 19 | Core–valence correlation effects in the ground and low-lying excited states of GaN. Molecular Physics, 2010, 108, 467-476. | 1.7 | Ο |
| 20 | Towards the full coupled-cluster model with triples: a retrospective after 25 years. Molecular Physics, 2010, 108, 2933-2940. | 1.7 | 0 |