## Nikolaj F Stepanov

List of Publications by Year in descending order

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| 101      | 1,006          | 17 h-index   | 27             |
|----------|----------------|--------------|----------------|
| papers   | citations      |              | g-index        |
| 103      | 103            | 103          | 687            |
| all docs | docs citations | times ranked | citing authors |

| #  | Article   | IF  | CITATIONS |
|----|---|-----|-----------|
| 1  | Ar–l2interactions: The models based on the diatomicsâ€inâ€molecule approach. Journal of Chemical Physics, 1996, 104, 9913-9925.   | 3.0 | 88        |
| 2  | Numerical-Analytic Implementation of the Higher-Order Canonical Van Vleck Perturbation Theory for the Interpretation of Medium-Sized Molecule Vibrational Spectra. Journal of Physical Chemistry A, 2012, 116, 3691-3709. | 2.5 | 69        |
| 3  | Theoretical Study of Structure and Electronic Absorption Spectra of Some Schiff Bases and Their Zinc Complexes. Inorganic Chemistry, 2009, 48, 11123-11130.   | 4.0 | 56        |
| 4  | Vibration-rotation problem for triatomic molecules with two large-amplitude coordinates. Journal of Molecular Spectroscopy, 1977, 67, 265-282.  | 1.2 | 40        |
| 5  | Simulation of structured 4T1â†'6A1 emission bands of Mn2+ impurity in Zn2SiO4: A first-principle methodology. Journal of Luminescence, 2012, 132, 2143-2150.  | 3.1 | 39        |
| 6  | Small Charged Water Clusters:  Cations. Journal of Physical Chemistry A, 1999, 103, 3285-3288.  | 2.5 | 38        |
| 7  | Anharmonic Vibrational Analysis of the Gas-Phase Infrared Spectrum of 1,1-Difluoroethylene Using the Operator Van Vleck Canonical Perturbation Theory. Journal of Physical Chemistry A, 2013, 117, 3041-3056.             | 2.5 | 30        |
| 8  | Environmental Broadening of the CTTS Bands: The Hexaammineruthenium(II) Complex in Aqueous Solution. Journal of Physical Chemistry A, 2010, 114, 12804-12812.   | 2.5 | 26        |
| 9  | ArHF vibrational predissociation dynamics using the diatomics-in-molecule potential energy surface. Journal of Chemical Physics, 1999, 111, 2470-2477.  | 3.0 | 25        |
| 10 | Diatomics-in-molecules description of the Rg–Hal2 rare gas–halogen van der Waals complexes with applications to He–Cl2. Journal of Chemical Physics, 1997, 106, 4575-4588.  | 3.0 | 23        |
| 11 | Two-centre model potential energy calculations for the 2Σ and 2Πstates of Li+2, Na+2, K+2, Rb+2 and Cs+2. Chemical Physics Letters, 1979, 60, 421-426.  | 2.6 | 22        |
| 12 | Molecular structure of the boron(III) oxide molecule in the SCF approximation. Journal of Molecular Structure, 1980, 68, 199-202.   | 3.6 | 21        |
| 13 | Interaction potentials and fragmentation dynamics of the Neâc Br2complex in the ground and electronically excited states. Journal of the Chemical Society, Faraday Transactions, 1994, 90, 3229-3236.                     | 1.7 | 21        |
| 14 | Errors in molecular integrals: An influence onRHF energy values. International Journal of Quantum Chemistry, 1977, 11, 1-15.  | 2.0 | 18        |
| 15 | First-order intermolecular diatomics-in-molecule potentials. Potential energy surfaces, spectra, and fragmentation dynamics of the Neâc Cl2 complex. Journal of Chemical Physics, 1997, 106, 10134-10144.                 | 3.0 | 18        |
| 16 | Small Charged Water Clusters:Â Anions. Journal of Physical Chemistry A, 1999, 103, 10975-10980.   | 2.5 | 17        |
| 17 | Simulating the structureless emission bands of Mn2+ ions in ZnCO3 and CaCO3 matrices by means of quantum chemistry. Russian Journal of Physical Chemistry A, 2013, 87, 245-251.   | 0.6 | 17        |
| 18 | The potential surface of the ground state and the force field of the B2O2 molecule in the SCF approximation. Journal of Structural Chemistry, 1983, 24, 252-254.  | 1.0 | 16        |

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|----|--|-----|-----------|
| 19 | Structure and interaction energies of the Ar…Cl2 complex. Application of first-order intermolecular potentials. Chemical Physics Letters, 1996, 261, 591-596.            | 2.6 | 15        |
| 20 | Some aspects of scaling the molecular quantum mechanical force field. Journal of Molecular Structure, 1996, 376, 363-368.  | 3.6 | 15        |
| 21 | Electron hydration: interface shells. Chemical Physics Letters, 2001, 344, 619-624.  | 2.6 | 15        |
| 22 | Nonempirical Description of the Atmospherically Important Anionic Species. I. Water Cluster Anions. Structural Chemistry, 2004, 15, 65-70.                               | 2.0 | 15        |
| 23 | Vibrational predissociation of ArHF: a test of global semiempirical potential energy surfaces. Physical Chemistry Chemical Physics, 2002, 4, 4992-4998.                  | 2.8 | 14        |
| 24 | Vibrationâ€"rotation hamiltonian for nonrigid triatomic molecules with diatomic rigid core. Chemical Physics, 1978, 31, 413-423.   | 1.9 | 13        |
| 25 | Diatomics-in-molecules study of LiOH. International Journal of Quantum Chemistry, 1979, 15, 49-56.   | 2.0 | 13        |
| 26 | Molecular structure of acetyldeyiethylphosphine, MeC(O)PMe2. Journal of Molecular Structure, 1977, 37, 251-259.  | 3.6 | 12        |
| 27 | Vibrational spectra and conformational analysis of five chlorosubstituted buta-1,3-dienes. Journal of Molecular Structure, 1978, 49, 17-27.                              | 3.6 | 12        |
| 28 | Electronic structure and electronic absorption spectra of molybdenum and tungsten oxotetrachlorides. Theoretica Chimica Acta, 1980, 56, 297-306.                         | 0.8 | 12        |
| 29 | The Kirkwood–Buckingham variational method and the boundary value problems for the molecular Schrödinger equation. Journal of Mathematical Physics, 1997, 38, 5626-5633. | 1.1 | 12        |
| 30 | Ab Initio Vibrational Analysis of Cyclopropene, Its Fluoro Derivatives, and Their Deutero Analogues. Journal of Physical Chemistry A, 1998, 102, 2363-2371.              | 2.5 | 12        |
| 31 | Stationary states and dissociation of H3O radical in water clusters. Russian Journal of Physical Chemistry A, 2009, 83, 798-808.   | 0.6 | 12        |
| 32 | Kinematic models of a force field. Journal of Molecular Structure, 1972, 12, 289-297.  | 3.6 | 11        |
| 33 | Diatomics-in-molecules study of alkali hydroxides. Journal of Molecular Structure, 1980, 67, 81-88.  | 3.6 | 11        |
| 34 | Advantages of scaled quantum mechanical molecular force fields. Journal of Molecular Structure, 1995, 348, 413-416.  | 3.6 | 11        |
| 35 | DIM potential energy surfaces for excited states of H3. Molecular Physics, 1980, 41, 377-382.  | 1.7 | 10        |
| 36 | Molecular mechanism of hydrogen bromide addition to olefins. Tetrahedron, 1982, 38, 2585-2589.   | 1.9 | 10        |

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|----|--|-----|-----------|
| 37 | Importance of correlation in LiBO. Theoretica Chimica Acta, 1985, 67, 287-292.   | 0.8 | 9         |
| 38 | On the use of restricted CI for correlation energy calculations: Application to BH. Chemical Physics Letters, 1985, 115, 241-244.  | 2.6 | 9         |
| 39 | Hydrated electron: Nonempirical cluster approach. International Journal of Quantum Chemistry, 2002, 88, 496-506.   | 2.0 | 9         |
| 40 | Convergence problems in the solution of SCF equations. International Journal of Quantum Chemistry, 1974, 8, 107-117.   | 2.0 | 8         |
| 41 | Approximate phaseâ€space transport theory for vibrational predissociation. Journal of Chemical Physics, 1993, 98, 5486-5498.   | 3.0 | 8         |
| 42 | Half- and full-collision VT energy transfer in the Heî—,Br2(B) system. Chemical Physics Letters, 1997, 269, 448-454.   | 2.6 | 8         |
| 43 | Classical vibrational predissociation dynamics: The effects of phase-space bifurcations. Journal of Chemical Physics, 1998, 108, 6282-6290.  | 3.0 | 7         |
| 44 | When and why Hund's cases arise. Journal of Molecular Spectroscopy, 1974, 52, 277-286.   | 1.2 | 6         |
| 45 | A new derivation for the Tellerâ€"Redlich isotopic product rule. Vibrational Spectroscopy, 1994, 7, 191-196.   | 2.2 | 6         |
| 46 | Theoretical study of VRT energy transfer in Ne+I2(B) collisions using a spectroscopic interaction potential. Journal of the Chemical Society, Faraday Transactions, 1998, 94, 2307-2313.                               | 1.7 | 6         |
| 47 | Comparative analysis of the state of lithium and sodium atoms in water clusters. Russian Journal of Physical Chemistry A, 2009, 83, 1134-1144.   | 0.6 | 6         |
| 48 | A quantum-chemical and gas phase electron diffraction study of the structure of formylphosphine and acetyldimethylphosphine. Russian Journal of Physical Chemistry A, 2010, 84, 1745-1751.                             | 0.6 | 6         |
| 49 | Electronic structure and one-electron properties of the MoO2Cl2 molecule. Electronic spectra of the MoO2Cl2, MoO2Br2 and WO2Br2 molecules. Theoretica Chimica Acta, 1982, 61, 369-377.                                 | 0.8 | 5         |
| 50 | Diatomics-in-molecules studies of the Li + H2O interaction. Computational and Theoretical Chemistry, 1983, 104, 403-409.   | 1.5 | 5         |
| 51 | Some Aspects of the Implementation of Scaling Quantum Mechanical Molecular Force Fields.<br>Structural Chemistry, 2004, 15, 95-101.  | 2.0 | 5         |
| 52 | The determination of the equilibrium geometry of a molecule with the use of microwave data and theoretical rotational-vibrational interaction constants. Russian Journal of Physical Chemistry A, 2006, 80, 1448-1455. | 0.6 | 5         |
| 53 | Scale factors as effective parameters for correcting nonempirical force fields. Russian Journal of Physical Chemistry A, 2007, 81, 585-592.  | 0.6 | 5         |
| 54 | The applicability of TD-DFT methods to calculations of the electronic absorption spectrum of hexaamminoruthenium(II) in aqueous solution. Russian Journal of Physical Chemistry A, 2010, 84, 39-43.                    | 0.6 | 5         |

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|----|--|-----|-----------|
| 55 | Nuclear charge changes: Influence on the energy of highly symmetrical molecules. Chemical Physics Letters, 1977, 45, 589-591.  | 2.6 | 4         |
| 56 | The extreme values of mean-square amplitudes of molecular vibrations. Journal of Molecular Structure, 1978, 43, 101-108.   | 3.6 | 4         |
| 57 | CNDO/2 calculations of intermolecular interactions in chlorine-ethylene and bromine-ethylene systems. Advances in Molecular Relaxation and Interaction Processes, 1978, 12, 325-332.                           | 0.5 | 4         |
| 58 | On diatomic state mixing and treatment of overlap in the diatomics-in-molecules theory. Chemical Physics Letters, 1979, 63, 396-400.   | 2.6 | 4         |
| 59 | Calculation of the potential curves of the low-lying electronic states of LiBe and LiBe+ in a mixed orbital basis by the SCF method. Journal of Structural Chemistry, 1983, 24, 321-323.                       | 1.0 | 4         |
| 60 | The vibrational predissociation lifetime of the Heâ $\in$   N+2 (X, ν = 1) complex. Chemical Physics Letters, 1994, 220, 93-96.  | 2.6 | 4         |
| 61 | Possible transformations of the ozone molecule in the presence of water associates. Russian Journal of Physical Chemistry A, 2006, 80, 106-114.  | 0.6 | 4         |
| 62 | Hybrid Ab initio/EFP approach for calculating <i>dâ€d</i> absorption spectrum of hexaammineruthenium(II) ion in aqueous solutions. International Journal of Quantum Chemistry, 2008, 108, 2711-2718.           | 2.0 | 4         |
| 63 | The conformational analysis of 2,3,4,5-tetrachloro-trans-hexa-1,3,5-triene. Journal of Molecular Structure, 1975, 29, 329-334.   | 3.6 | 3         |
| 64 | Concerning the physical solution of the inverse vibrational problem. Journal of Molecular Structure, 1975, 27, 423-431.  | 3.6 | 3         |
| 65 | Electron spectra and one-electron properties of the WOCl 4, WSCl 4 and WOBr 4 molecules. Chemical Physics, 1980, 53, 63-69.  | 1.9 | 3         |
| 66 | Ab initiofinite-field transition moment calculations. Journal of Physics B: Atomic, Molecular and Optical Physics, 1998, 31, 5275-5280.  | 1.5 | 3         |
| 67 | Interpreting the vibrational spectra of uracil molecules and their deuterated isotopomers using a scaled quantum-chemical quadratic force field. Russian Journal of Physical Chemistry A, 2012, 86, 1855-1861. | 0.6 | 3         |
| 68 | A theoretical study of the molecule of boric oxide B2O3. Journal of Structural Chemistry, 1981, 22, 21-28.   | 1.0 | 2         |
| 69 | Influence of the choice of the adiabatic approximation and nonadiabatic effects on molecular structure. Journal of Structural Chemistry, 1983, 24, 17-24.  | 1.0 | 2         |
| 70 | Fragment-localized analysis of the multiconfigurational wavefunctions. Chemical Physics, 1990, 148, 309-314.   | 1.9 | 2         |
| 71 | Effect of scaling of a quantum mechanical force field on the frequencies and forms of molecular vibrations. Journal of Structural Chemistry, 1998, 39, 169-174.  | 1.0 | 2         |
| 72 | Nitrite ion formation: Nonempirical simulation in terms of cluster model. International Journal of Quantum Chemistry, 2004, 100, 460-468.  | 2.0 | 2         |

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|----|---|-----|-----------|
| 73 | <title>Ultraviolet-light absorption and electron localization by ozone in the presence of water: nonempirical consideration</title> ., 2004, , .  |     | 2         |
| 74 | Properties of methanol bound to a defect of zeolitic structure. International Journal of Quantum Chemistry, 2005, 104, 214-222.   | 2.0 | 2         |
| 75 | On the kashiwagi-sasaki generalization of the löwdin orthogonalization and the inverse vibrational problem. International Journal of Quantum Chemistry, 1977, 12, 317-326.                                | 2.0 | 1         |
| 76 | Possible generalization of the optimized diatomics-in molecules theory: Further studies of the diatomic state mixing parameter adjustment. International Journal of Quantum Chemistry, 1980, 17, 679-687. | 2.0 | 1         |
| 77 | A study of the potential surface of the groud state of HCN by the method of diatomic fragments in molecules. Journal of Structural Chemistry, 1981, 22, 147-152.  | 1.0 | 1         |
| 78 | Unitary transformations with the unitary operators depending on projection operators. International Journal of Quantum Chemistry, 1981, 19, 793-803.  | 2.0 | 1         |
| 79 | A study of the potential surface of LiOB by the method of diatomic fragments in molecules. Journal of Structural Chemistry, 1983, 24, 339-343.  | 1.0 | 1         |
| 80 | Approximate treatment of the phase-space bottlenecks for vibrational predissociation. Journal of Physics B: Atomic, Molecular and Optical Physics, 1991, 24, L545-L549.                                   | 1.5 | 1         |
| 81 | Approximate Quantal Calculations on the Predissociative Lifetimes of the Ne…Hal <sub>2</sub> ( <i>X,v</i> ) (Hal = Cl,Br,I) Van Der Waals Complexes. Spectroscopy Letters, 1992, 25, 189-200.             | 1.0 | 1         |
| 82 | Three-dimensional quantum calculations on the HeBr2 (B) predissociative linewidths. , 1994, 2205, 178.  |     | 1         |
| 83 | Calculations of spectra and dynamics of rare-gas halogen molecule complexes using the semi-empirical potential energy surfaces. , 1997, , .   |     | 1         |
| 84 | Nonempirical estimation of the ionization conditions of water and amorphous ice. Russian Journal of Physical Chemistry A, 2007, 81, 941-948.  | 0.6 | 1         |
| 85 | Rate constants of atomic hydrogen formation in H3O+(H2O) n + e â†' H + (H2O) n gas-phase processes.<br>Russian Journal of Physical Chemistry A, 2009, 83, 1502-1510.                                      | 0.6 | 1         |
| 86 | Potential magnetic properties of nanotubes (n, 0) with Klein and Fujita edges. Russian Journal of Physical Chemistry A, 2012, 86, 1261-1267.  | 0.6 | 1         |
| 87 | Study of the geometric structure of the molecules of H2O, Li2O, and LiOH by the method of diatomic fragments in molecules. Journal of Structural Chemistry, 1979, 19, 665-671.                            | 1.0 | 0         |
| 88 | Low-temperature properties of wide-cut diesel fuels. Chemistry and Technology of Fuels and Oils, 1983, 19, 429-431.   | 0.5 | 0         |
| 89 | Bonding energy in (Li2OH)� complex ions. Journal of Structural Chemistry, 1987, 28, 19-22.  | 1.0 | 0         |
| 90 | Theoretical models of vibrational predissociation for van der Waals complexes., 1994,,.   |     | 0         |

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| 91  | Modeling the electronic structure of graphite intercalation compounds with lithium by metal complexes with polybenzene systems. Journal of Structural Chemistry, 1996, 37, 392-397.                             | 1.0 | 0         |
| 92  | Application of the non-Hermitean effective Hamiltonian method to metastable van der Waals complexes. , 1997, , .  |     | 0         |
| 93  | Structural peculiarities and the possibility of the existence of small water cluster anions (H2O) n â <sup>^</sup> withnâ‰ <b>4</b> . Russian Chemical Bulletin, 1997, 46, 36-41.                               | 1.5 | 0         |
| 94  | The possibility of the existence of (H2O) n â° anions withn=5, 6. Russian Chemical Bulletin, 1997, 46, 42-48.   | 1.5 | 0         |
| 95  | Ab-initio-based model for the charge transfer mechanisms in Ar+ + H2O collisions. International Journal of Mass Spectrometry, 2000, 203, 19-29.   | 1.5 | 0         |
| 96  | Title is missing!. Protection of Metals, 2002, 38, 78-82.   | 0.2 | 0         |
| 97  | <title>State of the art in quantum chemistry today</title> ., 2004, 5311, 39.   |     | O         |
| 98  | Simple molecules on a zeolite acceptor center: A quantum-chemical approach. Russian Journal of Physical Chemistry A, 2007, 81, 1365-1370.   | 0.6 | 0         |
| 99  | The equilibrium structures of the Li[C n ]1 (n = $7\hat{a} \in 12$ ) complexes and their alternation depending on n. Russian Journal of Physical Chemistry A, 2009, 83, 77-80.                                  | 0.6 | 0         |
| 100 | 9th International Conference on Fundamental and Applied Aspects of Physical Chemistry: PHYSICAL CHEMISTRY 2008 (Belgrade, September 24–26, 2008). Russian Journal of Physical Chemistry A, 2009, 83, 1429-1429. | 0.6 | 0         |
| 101 | Calculations of predissociative lifetimes of RGHal 2 Van der Waals complexes., 1992,,.  |     | 0         |