VIII-W  Vibrational Energy Transfer from Solute to Solvent: An Analysis Based upon Path Integral Influence Functional Theory and Mixed Quantum-Classical Molecular Dynamics Method

MIKAMI, Taiji; OKAZAKI, Susumu

Path integral influence functional theory and mixed quantum–classical molecular dynamics method have been applied to the analysis of molecular mechanism of solute vibrational energy relaxation in solution. A numerical example is CN⁻ ion in water.

First, quantum effect of the solvent has been examined by taking a zero limit of Planck constant relevant to the solvent degrees of freedom. The effect is unexpectedly great. Classical approximation for the solvent significantly overestimates the relaxation time. The degree of the overestimation depends much upon the process, e.g. single-phonon, two-phonon, and three-phonon processes. However, it is interesting to find that it affects little energy dissipation path. Second, in order to obtain a microscopic picture of the relaxation in terms of molecular motion, couplings between normal mode and the solute were divided into contributions from each water molecule. Couplings spread over the normal modes are not localized within particular modes. However, in molecular language, molecules in the first hydration shell are mostly responsible for the energy flow from the solute. Third, the couplings actually found in the solution have also been analyzed as a function of time. The solvent does not show delta-function-like coupling. This indicates that the system does not relax by collision which is usually found in the gas state. The coupling in the solution oscillates very rapidly. Then, matching of both transition frequency and phase between the interaction and the wave function of the system were found to be important. In a short time scale, we cannot find any systematic behavior of the coupling. It looks like random. However, integrating the coupling over pico second, the system shows relaxation to the ground state to a certain extent. The relaxation rate changes in hundred femto second or pico second reflecting the structural change of the hydration shell.

VIII-X  Molecular Dynamics Study of Classical Complex Systems

VIII-X-1  A Molecular Dynamics Study of Water Penetration into Biomembrane

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Free energy profile of water penetration into lipid bilayer has been evaluated based upon molecular dynamics calculations combined with thermodynamic integration method. The calculated profile for 25 arbitrarily chosen penetration paths shows that the free energy barrier height has a wide distribution, demonstrating a sort of selectivity of the path by water molecule. Structure of the path or structure of the surrounding lipid molecules was examined in order to extract a factor which determines the barrier height.

VIII-X-2  Molecular Dynamics Study of Mechanical Extension of Polyalanine by AFM Cantilever

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A series of molecular dynamics calculations have been performed mimicking the mechanical extension of α-helical polyalanine molecule in water on the AFM. Force profile has been evaluated as a function of extension. In an initial stage of the extension, ~ 5%, the force was found to follow the Hooke’s law up to about 140 pN. At the extension of about 10%, however, the restoring force diminishes, indicating the rupture of α-helix. Then, α-helix region and β-strand region separate from each other in one molecule. Further extension causes stepwise breakages of hydrogen bonds to give the wholly β-stranded polyalanine.
VIII-Y  Theoretical Studies on Electronic Structure and Dynamics of Electronically Excited States

VIII-Y-1  Chemical Reactions in the O(1D) + HCl System I. Ab Initio Global Potential Energy Surfaces for the 1A', 2A', and 1A'' States

NANBU, Shinkoh; KAMISAKA, Hideyuki1; BIAN, Wensheng; AYOAGI, Mutsumi; TANAKA, Kiyoshi2; NAKAMURA, Hiroki (1GUAS; 2Hokkaido Univ.)

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New global ab initio potential energy surfaces (PES) are presented for the low-lying 1A', 2A', and 1A'' electronic states which are correlated to O(1D) + HCl. These potential energy surfaces are computed by using the multi-reference configuration interaction method with the Davidson correction (MRCI + Q). The reference functions are constructed by the complete active space self-consistent field (CASSCF) calculations using the quadruple zeta + polarization basis set augmented with diffuse functions. The computations are carried out at about 5000 molecular conformations on each three-dimensional potential energy surface. The high accuracy of the computations is confirmed by a comparison with the available most accurate data for the ground state 1A'; thus the present work is the first report of the accurate potential energy surfaces for the two excited states. Three low-lying transition states on the excited surfaces, two (TS2 and TS4) on 1A'' and one (TS3) on 2A', are found. Since TS2 and TS3 are as low as 0.07 eV and 0.28 eV, respectively, and correlated to the electronic states which are correlated to O(1D) + HCl. The computations are carried out with diffuse functions. The computations are carried out between the He atom and the center of mass of HCN along the minimum energy path shows a strong angular dependence: R_m = 4.169 Å and 4.040 Å in the linear and anti-linear forms, respectively, while it is 3.528 Å in a T-shaped configuration. In the first excited internal-rotation state (j = 1), levels with l less than 4 are bound but not for the one with l = 5, according to the energy level diagram calculated from the present potential energy surface, where l denotes the quantum number for the end-over-end rotation of the complex. The energy level diagram is consistent with the millimeter-wave absorption spectroscopy combined with a pulsed-jet expansion technique which was applied to the measurement of the internal-rotation band of the He–HCN in the frequency region of 95–125 GHz. In total 13 rovibrational lines, split into nitrogen nuclear hyperfine structure, were observed for the fundamental internal-rotation band, j = 1–0. The observed transition frequencies were analyzed including their hyperfine splitting to yield an intermolecular potential energy surface, as improved from the one given by a coupled-cluster single double (triple) ab initio calculation. The surface obtained has a global minimum in the linear configuration (He···H–C–N) with a well depth of 30.2 cm\(^{-1}\), and the saddle point located in the anti-linear configuration (H–C–N···He) is higher by 8.174 cm\(^{-1}\) in energy than the global minimum. The distance R_m between the He atom and the center of mass of HCN molecule.


HARADA, Kensuke1; TANAKA, Keiichi1; TANAKA, Takehiko1; NANBU, Shinkoh; AYOAGI, Mutsumi (1Kyushu Univ.)

Millimeter-wave absorption spectroscopy combined with a pulsed-jet expansion technique was applied to the measurement of the internal-rotation band of the He–HCN in the frequency region of 95–125 GHz. In total 13 rovibrational lines, split into nitrogen nuclear hyperfine structure, were observed for the fundamental internal-rotation band, j = 1–0. The observed transition frequencies were analyzed including their hyperfine splitting to yield an intermolecular potential energy surface, as improved from the one given by a coupled-cluster single double (triple) ab initio calculation. The surface obtained has a global minimum in the linear configuration (He···H–C–N) with a well depth of 30.2 cm\(^{-1}\), and the saddle point located in the anti-linear configuration (H–C–N···He) is higher by 8.174 cm\(^{-1}\) in energy than the global minimum. The distance R_m between the He atom and the center of mass of HCN molecule.


NANBU, Shinkoh; ISHIDA, Toshimasa1; GRAY, Stephen K.2; GONZALEZ, Miguel3 (1Shizuoka Univ.; 2Argonne Natl. Lab.; 3Univ. Barcelona)

Computational algorithm for the determination of the global potential energy surfaces of polyatomic systems are developed with using the interpolant moving least squares scheme, which was proposed by Ishida and Schatz [Chem. Phys. Lett. 314, 369 (1999)]. In this scheme, any derivatives in quantum-chemical calculations are not required to construct the surface and in contrast with previously developed schemes based on Shepard interpolation alone. In our new algorithm, the molecular conformations are generated with the Monte Carlo method, while the remaining calculations are carried out on the macromolecular conformations.
Carlo sampling, and then the *ab initio* calculations for all of the conformations are performed by parallel computing. Therefore, we have good advantage for computational time for the serial calculations. Application is made to the tetra-atomic systems, the $2\mathrm{OH} \leftrightarrow \mathrm{H}_2\mathrm{O} + \mathrm{O}$ reaction.

Regarding to the wave packet dynamics, we are also developing the program code based on the MPI-library to make a time-evolution of the wave packet for the tetra-atomic systems.


CHOE, Jong-In\(^1\); CHANG, Suk-Kyu\(^1\); NANBU, Shinkoh

\(^1\)Chung-Ang Univ.


The structures and energies of *p*-tert-butylcalix[4]-crown-6-ether (1) in various conformers and their alkyl ammonium complexes have been calculated by *ab initio* HF/6-31G quantum mechanics method. We have tried to obtain the relative affinity of partial-cone and 1,3-alternate conformers of 1 for alkyl ammonium guests by comparison with its cone-shaped analogue. We have also calculated the relative complexation efficiencies of these host-guest complexes focusing on the binding sites of crown-6-ether moiety or benzene-rings pocket of the host molecule. 1. These calculations revealed that the crown moiety has better complexation efficiency than upper rim part of calix[4]arene that is in similar configuration of the host molecule 1. These calculations revealed that the crown moiety has better complexation efficiency than upper rim part of calix[4]arene that is in similar configuration of the host molecule. These calculations revealed that the crown moiety has better complexation efficiency than upper rim part of calix[4]arene that is in similar configuration of the host molecule.


CHOE, Jong-In\(^1\); CHANG, Suk-Kyu\(^1\); MINAMINO, Satoshi; NANBU, Shinkoh

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The structures and composition energies of penta-O-alkylated 1b and penta-O-tert-butyl ester 1e of *p*-tert-butylcalix[5]arene and its simplified structures (2b and 2e) toward a series of alkyl ammonium guests have been calculated by semi-empirical AM1 quantum mechanics method. For AM1 calculation, complexation efficiencies of the simplified host 2e are very similar to the values of host 1e. The complexes of simplified host 2e with alkyl ammonium ions have been also optimized by *ab initio* HF/6-31G quantum mechanics method. The calculated complexation efficiencies for 2e by *ab initio* method have been found to be bigger in magnitude than the values obtained by AM1 calculations for linear alkyl ammonium guests. Calculation results show that all of the calix[5] are derivatives investigated in this study have much better complexation ability toward ammonium cation without alkyl group over other alkyl ammonium guests. *Ab initio* calculations also well duplicate the molecular discriminating behaviors of calyx[5] arene derivative 2e between butyl ammonium ions: $n$-BuNH\(^+\) > iso-BuNH\(^+\) > sec-BuNH\(^+\) > tert-BuNH\(^+\).

**VIII-Y-6 Formation of HCl*(A2Σ\(^+\)) and HBr*(A2Σ\(^+\)) Resulting from He(2\(^3\)S) Penning Ionization of HCl and HBr**

TOKUE, Ikuo\(^1\); TANAKA, Hiroyuki\(^1\); YAMASAKI, Katsuyoshi\(^1\); NANBU, Shinkoh

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*He*(2\(^3\)S) Penning ionization of HCl and HBr leading to HCl*(A) and HBr*(A) has been studied optically by using a crossed-beam apparatus. The ratios of the vibrational population, $P_{v'}/P_{v}(v' = 2$ and 3) of HCl*(A) and $P_v/P_0$ of HBr*(A), increase with the collision energy in the region of 120–200 meV. The rotational distributions of HCl*(A, $v' = 0$) and HBr*(A, $v' = 0$) can be represented by a double-Boltzmann distribution; the temperatures are 200 ± 50 and 700 ± 80 K for HCl*(A, $v' = 0$) and 250 ± 50 and 1200 ± 200 K for HBr*(A), and are nearly independent of the collision energy. The model potential surface for He*(Li) + HCl as the entrance channel is nearly isotropic and shows a shallow dip of about 20 meV, while the surface for He + HCl*(A) as the exit channel is anisotropic and shows a deep minimum of 250 meV in the He–H–Cl collision direction. These results suggest that at least two processes contribute to formation of these ions; one is the direct Penning ionization and the other is the formation via a temporary complex [HeHCl(A)]\(^+\) by orbiting.

**VIII-Y-7 Theoretical Study of Vibrational States for AINC/AICN**

MINAMINO, Satoshi; NANBU, Shinkoh; AOYAGI, Mutsumi

The character of the low-lying electronic states of AINC strongly depends on the bond distance of AI-NC, because the covalent states are lying closely to the ionic states. Especially, due to the electronic ground state having the ionic character, the bending vibrational motion has a quite strong anharmonicity, and the motion is characterized as a large amplitude motion (LAM). Therefore, in general, the observed spectrum has the complicated vibronic structures and it is difficult to understand the molecular conformation having the LAM. In this study, we determined the global potential energy surfaces of the electronic ground state for the isomerization of AINC/AICN, and we elucidated the molecular structure of the complicated vibronic states by solving the exact Schrödinger equation for the nuclear motion.

**VIII-Y-8 Boundary Expansion in Time-Dependent Nonadiabatic Problems**

TAKAMI, Toshiya
Time-dependent nonadiabatic transitions were studied as an initial value problem in a finite parameter range. We introduced "boundary expansion" by expanding Schrödinger’s equation around end points of the parameter. We have shown that extra transitions emerge at the points, and are explained by the terms of the expansion. We have also shown that Berry’s superadiabatic base is naturally defined through the expansion. By the use of the superadiabatic base, we performed numerical analysis of multi-level nonadiabatic transitions in systems with a random matrix Hamiltonian.

VIII-Y-9 Optimal Control of Random Matrix Systems with a Parameter

TAKAMI, Toshiya; FUJISAKI, Hiroshi

We studied control of chaotic quantum systems by applying Optimal Control Theory (OCT) to random matrix Hamiltonian systems with a parameter. For simple two-state problems, we have shown that the optimized field obtained by OCT can be classified into several types: (i) oscillating one with a frequency which agrees with the energy split between the initial and the final state; (ii) one which mainly uses transition between diabatic states at an avoided crossing; (iii) one which can be seen as nonadiabatic transition, etc. Although the oscillating solution is always obtained even in many level systems if the final time is large enough, it is difficult to distinguish other types of solution in short time.

We have shown that the distribution of the optimized solutions for short time is affected by dynamical properties of the system. In particular, we examined influence of the strength of chaos or complexity of the system. We have also studied systems with banded random elements.