Mitsuhiko Miyazaki

List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/8121947/publications.pdf

Version: 2024-02-01

69 papers 2,084 citations

257450 24 h-index 243625 44 g-index

72 all docs 72 docs citations

72 times ranked 1445 citing authors

#	Article	IF	CITATIONS
1	Real-time observation of photoionization-induced water migration dynamics in 4-methylformanilide–water by picosecond time-resolved infrared spectroscopy and ⟨i⟩ab initio⟨ i⟩ molecular dynamics simulations. Physical Chemistry Chemical Physics, 2021, 24, 73-85.	2.8	8
2	Revealing the role of excited state proton transfer (ESPT) in excited state hydrogen transfer (ESHT): systematic study in phenol–(NH ₃) _n clusters. Chemical Science, 2021, 12, 3836-3856.	7.4	18
3	Isomer-Selective Spectroscopy and Dynamics of Phenol–Ar _{<i>n</i>} (<i>n</i> â‰ುち) Clusters. Journal of Physical Chemistry A, 2021, 125, 9969-9981.	2.5	2
4	Excited state hydrogen transfer dynamics in phenol–(NH ₃) ₂ studied by picosecond UV-near IR-UV time-resolved spectroscopy. Physical Chemistry Chemical Physics, 2020, 22, 5740-5748.	2.8	4
5	lonization-Induced π → H Site Switching in Resorcinol–Ar _{<i>n</i>} (<i>n</i> = 1 and 2) Clusters Probed by Infrared Spectroscopy. Journal of Physical Chemistry A, 2019, 123, 6828-6839.	2.5	2
6	Effect of alkali ions on optical properties of flavins: vibronic spectra of cryogenic M ⁺ lumiflavin complexes (M = Li–Cs). Faraday Discussions, 2019, 217, 256-275.	3.2	17
7	Sequential microhydration of cationic 5-hydroxyindole (5Hl ⁺): infrared photodissociation spectra of 5Hl ⁺ –W _n clusters (W = H ₂ O,) Tj ETQq1	1 0 2.8 431	.4 r gB T /Overlo
8	Electronâ€Proton Transfer Mechanism of Excitedâ€State Hydrogen Transfer in Phenolâ€(NH ₃ (sub> <i>n</i> (<i>n=</i>)3 and 5). Chemistry - A European Journal, 2018, 24, 881-890.	3.3	8
9	A theoretical study on the size-dependence of ground-state proton transfer in phenol–ammonia clusters. Physical Chemistry Chemical Physics, 2018, 20, 3265-3276.	2.8	8
10	Real-time observation of the photoionization-induced water rearrangement dynamics in the 5-hydroxyindole–water cluster by time-resolved IR spectroscopy. Physical Chemistry Chemical Physics, 2018, 20, 3079-3091.	2.8	16
11	Effect of alkali ions on optical properties of flavins: vibronic spectra of cryogenic M ⁺ lumichrome ions (M = Li–Cs) in the gas phase. Physical Chemistry Chemical Physics, 2018, 20, 22148-22158.	2.8	15
12	Stepwise microhydration of aromatic amide cations: water solvation networks revealed by the infrared spectra of acetanilide ⁺ –(H ₂ O) _n clusters (<i>n</i> â‰ು). Physical Chemistry Chemical Physics, 2018, 20, 3148-3164.	2.8	15
13	Electron-proton transfer mechanism of excited-state hydrogen transfer in phenolâ^'(NH3) (nâ€=â€⁻5) studied by delayed ionization detected femtosecond time-resolved NIR spectroscopy. Chemical Physics, 2018, 515, 580-585.	1.9	6
14	A structural study on the excimer state of an isolated benzene dimer using infrared spectroscopy in the skeletal vibration region. Physical Chemistry Chemical Physics, 2017, 19, 22759-22776.	2.8	20
15	Deciphering environment effects in peptide bond solvation dynamics by experiment and theory. Physical Chemistry Chemical Physics, 2017, 19, 22564-22572.	2.8	11
16	Photoionization-induced π ↔ H site switching dynamics in phenol ⁺ –Rg (Rg = Ar, Kr) dimers probed by picosecond time-resolved infrared spectroscopy. Physical Chemistry Chemical Physics, 2016, 18, 24746-24754.	2.8	19
17	Theoretical Study on the Size Dependence of Ground-State Proton Transfer in 1-Naphthol–Ammonia Clusters. Journal of Physical Chemistry A, 2016, 120, 7167-7174.	2.5	6
18	Revealing Single Molecular Solvent Reorientation Dynamics by Complementary Use of Picosecond Time Resolved IR Spectroscopy and MD Simulation. Molecular Science, 2016, 10, A0087.	0.2	0

#	Article	IF	CITATIONS
19	Mass analyzed threshold ionization detected infrared spectroscopy: isomerization activity of the phenol–Ar cluster near the ionization threshold. Physical Chemistry Chemical Physics, 2015, 17, 2494-2503.	2.8	12
20	Trapped Hydronium Radical Produced by Ultraviolet Excitation of Substituted Aromatic Molecule. Journal of Physical Chemistry A, 2015, 119, 12730-12735.	2.5	6
21	Single water solvation dynamics in the 4-aminobenzonitrile–water cluster cation revealed by picosecond time-resolved infrared spectroscopy. Physical Chemistry Chemical Physics, 2015, 17, 29969-29977.	2.8	20
22	Real time observation of the excimer formation dynamics of a gas phase benzene dimer by picosecond pump–probe spectroscopy. Physical Chemistry Chemical Physics, 2015, 17, 25989-25997.	2.8	27
23	Electron–Proton Decoupling in Excitedâ€ S tate Hydrogen Atom Transfer in the Gas Phase. Angewandte Chemie - International Edition, 2015, 54, 15089-15093.	13.8	20
24	The mechanism of excited-state proton transfer in 1-naphthol–piperidine clusters. Physical Chemistry Chemical Physics, 2015, 17, 25393-25402.	2.8	4
25	Theoretical Study on the Size Dependence of Excited State Proton Transfer in 1-Naphthol–Ammonia Clusters. Journal of Physical Chemistry B, 2015, 119, 2415-2424.	2.6	9
26	Microhydrated aromatic cluster cations: Binding motifs of 4-aminobenzonitrile-(H2O)n cluster cations with n ≤. Journal of Chemical Physics, 2014, 141, 214301.	3.0	29
27	lonization-induced π → H site-switching in phenol–CH ₄ complexes studied using IR dip spectroscopy. Physical Chemistry Chemical Physics, 2014, 16, 110-116.	2.8	13
28	Microsolvation of the acetanilide cation (AA ⁺) in a nonpolar solvent: IR spectra of AA ⁺ –L _n clusters (L = He, Ar, N ₂ ; n ≶0). Physical Chemistry Chemical Physics, 2014, 16, 7980-7995.	2.8	26
29	Solvation Dynamics of a Single Water Molecule Probed by Infrared Spectra—Theory Meets Experiment. Angewandte Chemie - International Edition, 2014, 53, 14601-14604.	13.8	31
30	Solvent Migration in Microhydrated Aromatic Aggregates: Ionizationâ€Induced Site Switching in the 4â€Aminobenzonitrile–Water Cluster. Chemistry - A European Journal, 2014, 20, 2031-2039.	3.3	21
31	Structure of 1-naphthol–water clusters in the S1 state studied by UV–IR fluorescence dip spectroscopy and ab initio molecular orbital calculations. Chemical Physics Letters, 2013, 557, 19-25.	2.6	4
32	Unusual Behavior in the First Excited State Lifetime of Catechol. Journal of Physical Chemistry Letters, 2013, 4, 3819-3823.	4.6	23
33	Conformationally resolved spectra of acetaminophen by UV-UV hole burning and IR dip spectroscopy in the gas phase. Physical Chemistry Chemical Physics, 2013, 15, 957-964.	2.8	18
34	IR Spectroscopy of the 4â€Aminobenzonitrile–Ar Cluster in the S O , S 1 Neutral and D O Cationic States. ChemPhysChem, 2013, 14, 741-745.	2.1	13
35	Ground State Proton Transfer in Phenol–(NH ₃) _{<i>n</i>} (<i>n</i> ≶1) Clusters Studied by Mid-IR Spectroscopy in 3–10 μm Range. Journal of Physical Chemistry A, 2013, 117, 1522-1530.	2.5	30
36	MODE-specific deactivation of adenine at the singlet excited states. Journal of Chemical Physics, 2013, 139, 124311.	3.0	4

3

#	Article	IF	Citations
37	Microsolvation of the 4â€Aminobenzonitrile Cation (ABN +) in a Nonpolar Solvent: IR Spectra of ABN + L n (L=Ar and N 2 , n â‰ #). ChemPhysChem, 2013, 14, 728-740.	2.1	17
38	IR–UV Double Resonance Spectroscopy as Implemented by Polarized Laser Schemes: Probing Orientations of Vibrational Transition Dipole Moments. Chemistry Letters, 2013, 42, 1070-1072.	1.3	0
39	IR spectroscopy of monohydrated tryptamine cation: Rearrangement of the intermolecular hydrogen bond induced by photoionization. Journal of Chemical Physics, 2012, 137, 224311.	3.0	23
40	Ionizationâ€Induced Solvent Migration in Acetanilideâ€Methanol Clusters Inferred from Isomerâ€Selective Infrared Spectroscopy. ChemPhysChem, 2012, 13, 3875-3881.	2.1	14
41	Watching Water Migration around a Peptide Bond. Angewandte Chemie - International Edition, 2012, 51, 6604-6607.	13.8	63
42	Isomer selective IR-UV depletion spectroscopy of 4-fluorotoluene-NH3: evidence for π-proton-acceptor and linear hydrogen-bonded complexes. Physical Chemistry Chemical Physics, 2011, 13, 15633.	2.8	7
43	Photoionization-induced large-amplitude pendular motion in phenol ⁺ –Kr. Physical Chemistry Chemical Physics, 2011, 13, 2744-2747.	2.8	32
44	Conformational reduction of DOPA in the gas phase studied by laser desorption supersonic jet laser spectroscopy. Physical Chemistry Chemical Physics, 2011, 13, 7812.	2.8	23
45	Gas-Phase Spectroscopy of Synephrine by Laser Desorption Supersonic Jet Technique. Journal of Physical Chemistry A, 2011, 115, 10363-10369.	2.5	29
46	lonization-induced π → H site switching dynamics in phenol–Ar ₃ . Physical Chemistry Chemical Physics, 2011, 13, 2409-2416.	2.8	37
47	Gas-phase Infrared Spectroscopy of Monopeptides from 10 to 3 µm. Chemistry Letters, 2011, 40, 1157-1158.	1.3	3
48	Isomerization reaction in high-n Rydberg states of phenol–Ar/Kr clusters measured by autoionization detected infrared spectroscopy. Chemical Physics Letters, 2011, 513, 208-211.	2.6	16
49	Holeâ€Burning Spectra of <i>m</i> â€Fluorophenol/Ammonia (1:3) Clusters and Their Excited State Hydrogen Transfer Dynamics. ChemPhysChem, 2011, 12, 1928-1934.	2.1	10
50	Measurement of adiabatic ionization energies of the rotational isomers of n-propylbenzene and m-fluorophenol by direct VUV laser photoionization. Chemical Physics Letters, 2010, 485, 31-35.	2.6	1
51	Excited state hydrogen transfer dynamics in substituted phenols and their complexes with ammonia: ππâ^—πσâ^— energy gap propensity and ortho-substitution effect. Journal of Chemical Physics, 2010, 133, 124	330 313.	123
52	Evidence for Catechol Ring- Induced Conformational Restriction in Neurotransmitters. Journal of Physical Chemistry Letters, 2010, 1, 1130-1133.	4.6	39
53	IR spectra of resorcinol+–Ar cluster cations (n= 1, 2): Evidence for photoionization-induced π → H isomerization. Chemical Physics Letters, 2009, 474, 7-12.	2.6	18
54	Isomer selective infrared spectroscopy of supersonically cooled cis- and trans-N-phenylamides in the region from the amide band to NH stretching vibration. Physical Chemistry Chemical Physics, 2009, 11, 6098.	2.8	41

#	Article	IF	CITATIONS
55	High-energy, broadly tunable, narrow-bandwidth mid-infrared optical parametric system pumped by quasi-phase-matched devices. Optics Letters, 2008, 33, 1699.	3.3	42
56	Vibrational Signature of the Conformers in Tyramine Studied by IR Dip and Dispersed Fluorescence Spectroscopies. Journal of Physical Chemistry A, 2008, 112, 13463-13469.	2.5	27
57	IR spectra of phenol+–Krn cluster cations (n=1,2): Evidence for photoionization-induced π→H isomerization. Chemical Physics Letters, 2007, 443, 227-231.	2.6	34
58	IR laser manipulation of cis↔trans isomerization of 2-naphthol and its hydrogen-bonded clusters. Journal of Chemical Physics, 2006, 124, 054315.	3.0	12
59	Electronic and Infrared Spectroscopy of [Benzeneâ^'(Methanol)n]+(n= 1â^'6). Journal of Physical Chemistry A, 2005, 109, 9471-9480.	2.5	13
60	Morphology of Protonated Methanol Clusters:  An Infrared Spectroscopic Study of Hydrogen Bond Networks of H+(CH3OH)n (n = 4â^'15). Journal of Physical Chemistry A, 2005, 109, 138-141.	2.5	51
61	Infrared Spectroscopic Evidence for Protonated Water Clusters Forming Nanoscale Cages ChemInform, 2004, 35, no.	0.0	3
62	Electronic spectroscopy of benzene–water cluster cations, [C6H6–(H2O)n]+ (n=1–4): spectroscopic evidence for phenyl radical formation through size-dependent intracluster proton transfer reactions. Chemical Physics Letters, 2004, 399, 412-416.	2.6	35
63	Infrared Spectroscopic Evidence for Protonated Water Clusters Forming Nanoscale Cages. Science, 2004, 304, 1134-1137.	12.6	493
64	Infrared Spectroscopy of Size-Selected Benzeneâ "Water Cluster Cations [C6H6â (H2O)n]+ (n = 1â 23): Hydrogen Bond Network Evolution and Microscopic Hydrophobicity. Journal of Physical Chemistry A, 2004, 108, 10656-10660.	2.5	55
65	A Molecular Cluster Study on Activated CH/Ï€ Interactions:Â Infrared Spectroscopy of Aromatic Moleculeâ^'Acetylene Clusters. Journal of Physical Chemistry A, 2004, 108, 2652-2658.	2.5	67
66	Binding Energy of the Benzeneâ^'Water Cluster Cation:Â An Ar-Mediated IR Photodissociation Study. Journal of Physical Chemistry A, 2004, 108, 8269-8272.	2.5	35
67	Infrared spectroscopy of hydrated benzene cluster cations, [C6H6-(H2O)n]+ (n = 1–6): Structural changes upon photoionization and proton transfer reactions. Physical Chemistry Chemical Physics, 2003, 5, 1137-1148.	2.8	79
68	Infrared spectroscopy of the benzene–H2O cluster cation: experimental study on the drastic structural change upon photoionization. Chemical Physics Letters, 2001, 349, 431-436.	2.6	63
69	Infrared spectroscopy of the phenol-N2 cluster in S0 and D0: Direct evidence of the in-plane structure of the cluster. Journal of Chemical Physics, 1999, 110, 11125-11128.	3.0	61