

JANUARY 16, 2014

VOLUME 118

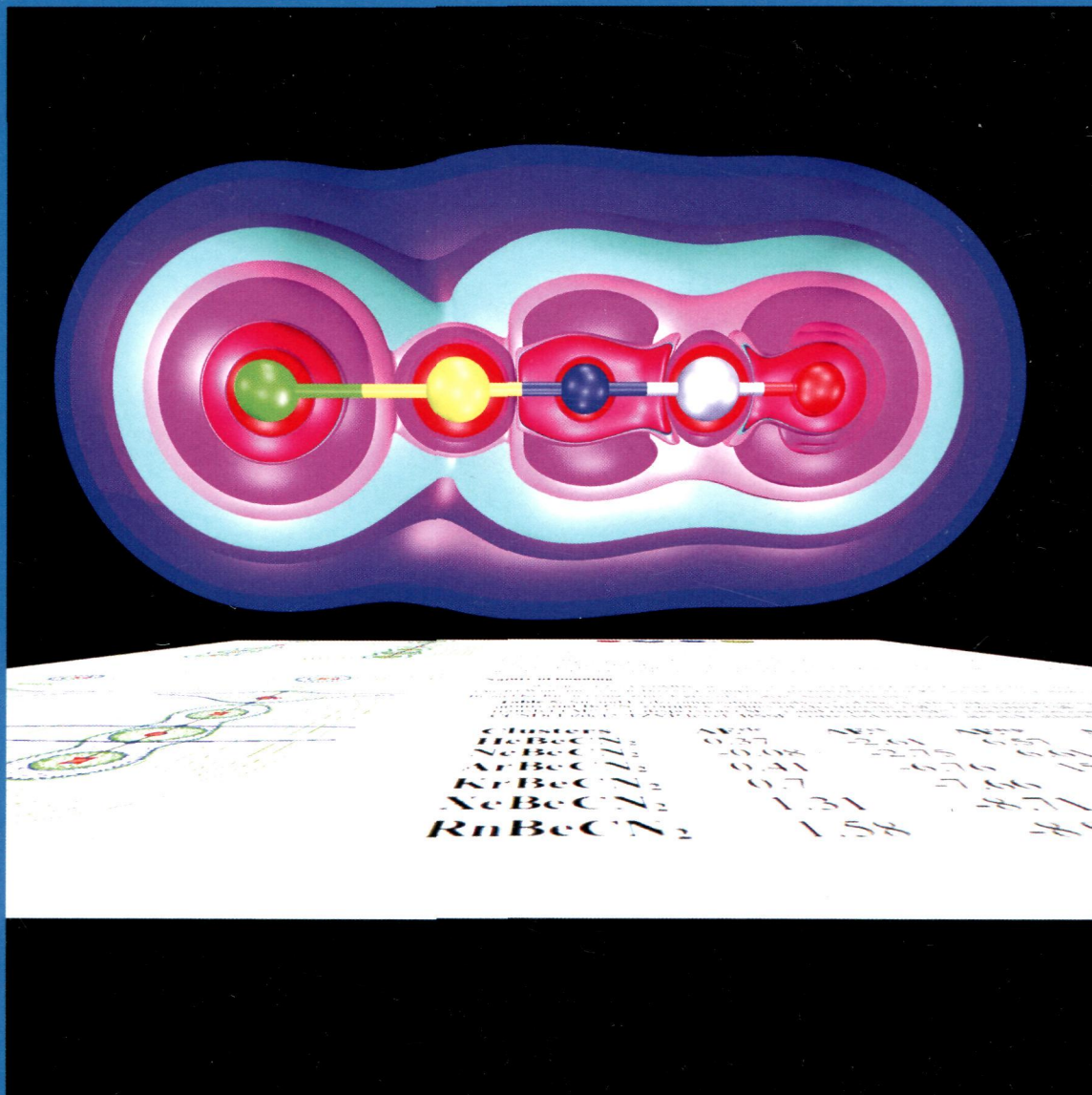
NUMBER 2

pubs.acs.org/JPCA

THE JOURNAL OF PHYSICAL CHEMISTRY

A

A Strong Be–Ng Bond
in the Presence
of an Electron
Withdrawing Group
(see page 5A)



ISOLATED MOLECULES, CLUSTERS, RADICALS, AND IONS; ENVIRONMENTAL CHEMISTRY,
GEOCHEMISTRY, AND ASTROCHEMISTRY; THEORY



ACS Publications

MOST TRUSTED. MOST CITED. MOST READ.

www.acs.org


ON THE COVER: A strong Be–Ng bond in the presence of an electron withdrawing group. See page 487.


Articles

Spectroscopy, Photochemistry, and Excited States


- 333 dx.doi.org/10.1021/jp411609t
Molecular Rotation–Vibration Dynamics of Low-Symmetric Hydrate Crystal in the Terahertz Region
Xiaojuan Fu, Hongya Wu, Xiaoqing Xi, and Ji Zhou*
- 339 dx.doi.org/10.1021/jp4097919
Effect of Amino Group Charge on the Photooxidation Kinetics of Aromatic Amino Acids
Natalya N. Saprygina, Olga B. Morozova, Günter Grampp, and Alexandra V. Yurkovskaya*
- 350 dx.doi.org/10.1021/jp409982f
Light-Induced Opening and Closing of the Intramolecular Hydrogen Bond in Glyoxylic Acid
Adriana Olbert-Majkut,* Jan Lundell, and Maria Wierzejewska
- 358 dx.doi.org/10.1021/jp410740u
Ab initio Theoretical Study on the $4f^2$ and $4f5d$ Electronic Manifolds of Cubic Defects in $\text{CaF}_2\text{:Pr}^{3+}$
Marek Krośnicki,* Andrzej Kędziorski, Luis Seijo, and Zoila Barandiarán
- 369 dx.doi.org/10.1021/jp411149e
Cu(II)- and Mn(III)-Porphyrin-Derived Oligomeric Multianions: Structures and Photoelectron Spectra
Ulrike Schwarz, Matthias Vonderach, Markus K. Armbruster, Karin Fink, Manfred M. Kappes, and Patrick Weis*
- 380 dx.doi.org/10.1021/jp411298p
Vibrational Shifts of HXeCl in Matrix Environments
Keisuke Niimi, Akira Nakayama,* Yuriko Ono, and Tetsuya Taketsugu
- 388 dx.doi.org/10.1021/jp411643e
Multiple Detachment of the SF_6^- Molecular Anion with Shaped Intense Laser Pulses
Yishai Albeck, Durai Murugan Kandhasamy, and Daniel Strasser*

396  dx.doi.org/10.1021/jp409043e
Toward a Better Understanding of Fe(III)–EDDS Photochemistry: Theoretical Stability Calculation and Experimental Investigation of 4-*tert*-Butylphenol Degradation
Yanlin Wu, Marcello Brigante, Wenbo Dong,* Pascal de Sainte-Claire, and Gilles Mailhot*

404  dx.doi.org/10.1021/jp4095886
Imaging and Scattering Studies of the Unimolecular Dissociation of the BrCH₂CH₂O Radical from BrCH₂CH₂ONO Photolysis at 351 nm
Lei Wang, Chow-Shing Lam, Rabi Chhantyal-Pun, Matthew D. Brynteson, Laurie J. Butler,* and Terry A. Miller


417  dx.doi.org/10.1021/jp412142w
Using Terahertz Spectroscopy and Solid-State Density Functional Theory to Characterize a New Polymorph of 5-(4-pyridyl) tetrazole
Steven Pellizzeri, Sean P. Delaney, Timothy M. Korter, and Jon Zubieta*

Environmental and Atmospheric Chemistry, Aerosol Processes, Geochemistry, and Astrochemistry

427  dx.doi.org/10.1021/jp4082905
First Ring Formation by Radical Addition of Propargyl to But-1-ene-3-yne in Combustion. Theoretical Study of the C₄H₇ Radical System
Daniela Trogolo, Andrea Maranzana,* Giovanni Ghigo, and Glauco Tonachini*


441 dx.doi.org/10.1021/jp409946j
Heterogeneous Interaction of H₂O₂ with Arizona Test Dust
Atallah El Zein, Manolis N. Romanias, and Yuri Bedjanian*


449 dx.doi.org/10.1021/jp410077g
New Understanding of the Formation of PCDD/Fs from Chlorophenol Precursors: A Mechanistic and Kinetic Study
Yanfeng Zhang, Dongju Zhang,* Jun Gao, Jinhua Zhan, and Chengbu Liu

457  dx.doi.org/10.1021/jp410345k
Products of the OH Radical-Initiated Reactions of Furan, 2- and 3-Methylfuran, and 2,3- and 2,5-Dimethylfuran in the Presence of NO
Sara M. Aschmann, Noriko Nishino, Janet Arey,* and Roger Atkinson*

467  dx.doi.org/10.1021/jp411134v
Position-Specific and Clumped Stable Isotope Studies: Comparison of the Urey and Path-Integral Approaches for Carbon Dioxide, Nitrous Oxide, Methane, and Propane
Michael A. Webb and Thomas F. Miller III*


Molecular Structure, Quantum Chemistry, and General Theory

475  dx.doi.org/10.1021/jp407854r
Engineering Frontier Energy Levels in Donor–Acceptor Fluoren-9-ylidene Malononitriles versus Fluorenones
Paul J. Homnick, Jonathan S. Tinkham, Raymond Devaughn, and Paul M. Lahti*

487  dx.doi.org/10.1021/jp409941v
In Quest of Strong Be–Ng Bonds among the Neutral Ng–Be Complexes
Sudip Pan, Diego Moreno, José Luis Cabellos, Jonathan Romero, Andres Reyes, Gabriel Merino,* and Pratim K. Chattaraj*

495 dx.doi.org/10.1021/jp410191y
Molecule-Optimized Basis Sets and Hamiltonians for Accelerated Electronic Structure Calculations of Atoms and Molecules
Gergely Gidofalvi and David A. Mazziotti*

503  dx.doi.org/10.1021/jp410458w
Gauche Preference of β -Fluoroalkyl Ammonium Salts
Josué M. Silla, Wesley G. D. P. Silva, Rodrigo A. Cornanich, Roberto Rittner, Cláudio F. Tormena, and Matheus P. Freitas*

508  dx.doi.org/10.1021/jp4109128
Structural Exploration of Water, Nitrate/Water, and Oxalate/Water Clusters with Basin-Hopping Method Using a Compressed Sampling Technique
Yi-Rong Liu, Hui Wen, Teng Huang, Xiao-Xiao Lin, Yan-Bo Gai, Chang-Jin Hu, Wei-Jun Zhang,* and Wei Huang*

517  dx.doi.org/10.1021/jp411039m
Time-Dependent Density Functional Methods for Raman Spectra in Open-Shell Systems
Frey W. Aquino and George C. Schatz*

526  dx.doi.org/10.1021/jp4117003
Lone Pairs: An Electrostatic Viewpoint
Anmol Kumar, Shridhar R. Gadre,* Neetha Mohan, and Cherumuttathu H. Suresh*

Additions and Corrections

533 dx.doi.org/10.1021/jp4123985
Correction to “Near-Infrared Surface-Enhanced Raman Spectroscopy (NIR-SERS) for the Identification of Eosin Y: Theoretical Calculations and Evaluation of Two Different Nanoplasmonic Substrates”
Nathan G. Greeneltch, Amber S. Davis, Nicholas A. Valley, Francesca Casadio, George C. Schatz, Richard P. Van Duyne, and Nilam C. Shah*

 Supporting Information available via online article