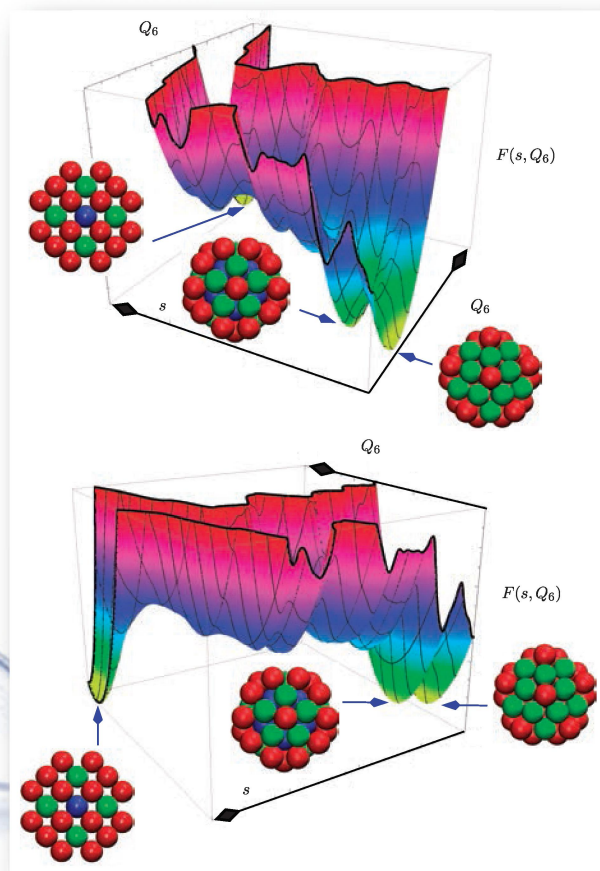


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Perspective: Insight into reaction coordinates and dynamics from the potential energy landscape

PERSPECTIVES

Perspective: Insight into reaction coordinates and dynamics from the potential energy landscape

D. J. Wales *J. Chem. Phys.* **142**, 130901 (2015)

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Communication: Unraveling the He droplet-mediated soft-landing from ab initio-assisted and time-resolved density functional simulations: Au@He₃₀₀/TiO₂(110)

Maria Pilar de Lara-Castells, Néstor F. Aguirre, Hermann Stoll, Alexander O. Mitrushchenkov, David Mateo and Martí Pi *J. Chem. Phys.* **142**, 131101 (2015);

Communication: A simple full range analytical potential for H₂b₃Σ⁺u, H–He Σ, and He₂1Σ⁺g
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Communication: Note on detailed balance in symmetrical quasi-classical models for electronically non-adiabatic dynamics William H. Miller and Stephen J. Cotton

J. Chem. Phys. **142**, 131103 (2015)

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Comments

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Michael Galperin, Mark A. Ratner and Abraham Nitzan

J. Chem. Phys. **142**, 137101 (2015);

Response to “Comment on ‘Frequency-domain stimulated and spontaneous light emission signals at molecular junctions’” [J. Chem. Phys. 142, 137101 (2015)]

Upendra Harbola, Bijay Kumar Agarwalla and Shaul Mukamel

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