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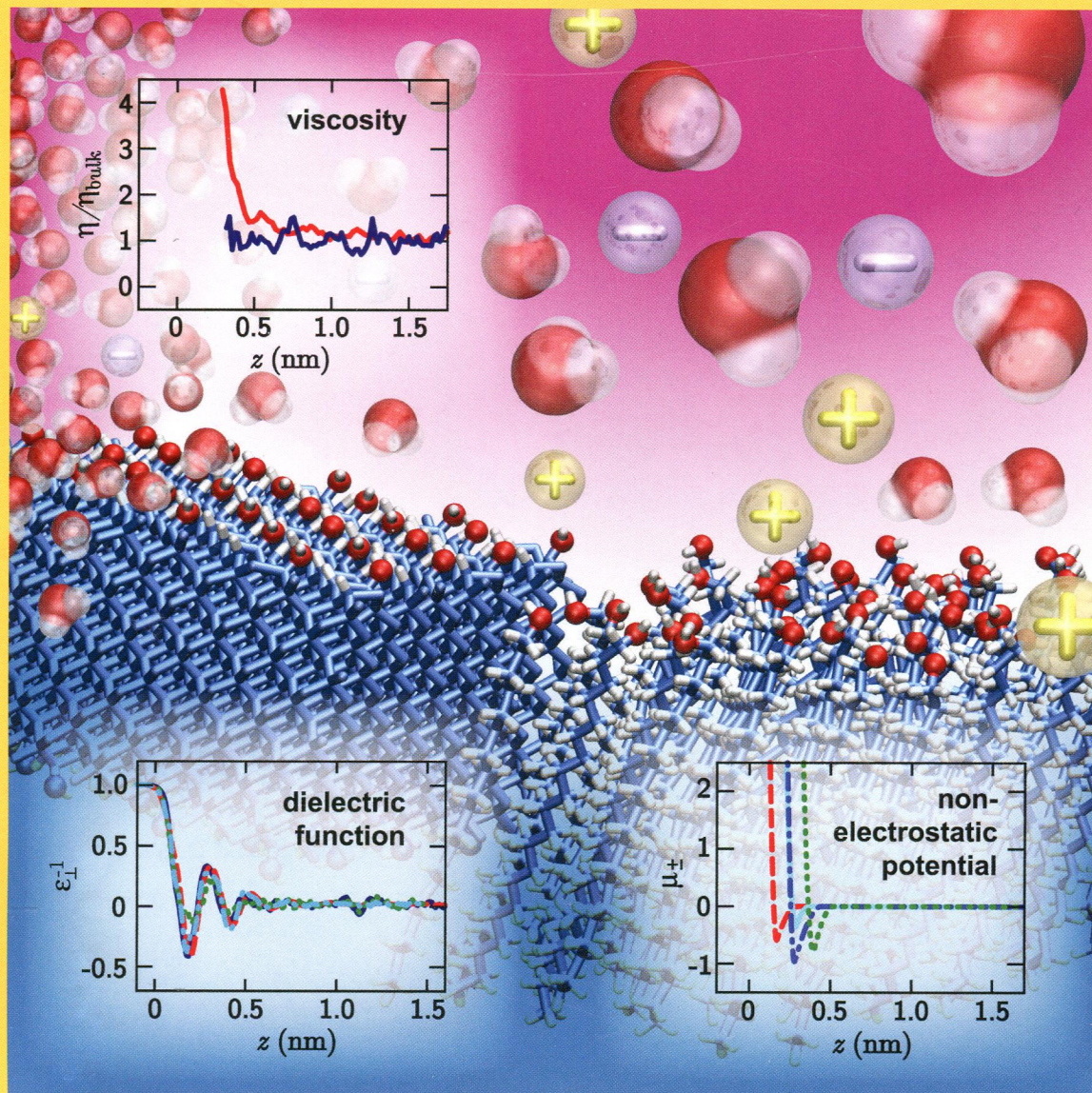
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# THE JOURNAL OF PHYSICAL CHEMISTRY

# B



Molecular Simulations and Continuum Theory Reveal the Microscopic Origins of Electrokinetic Effects at Solid-Electrolyte Interfaces (see page 5A)

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# THE JOURNAL OF PHYSICAL CHEMISTRY **B**

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Pages 11397-11894

## **Feature Article**

### ***Beyond the Continuum: How Molecular Solvent Structure Affects Electrostatics and Hydrodynamics at Solid-Electrolyte Interfaces***

Douwe Jan Bonthuis and Roland R. Netz

pp 11397–11413

**Publication Date (Web):** August 8, 2013 (Feature Article)

**DOI:** 10.1021/jp402482q

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**DOI:** 10.1021/jp406861u

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**DOI:** 10.1021/jp406864e

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**DOI:** 10.1021/jp400655v

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**DOI:** 10.1021/jp402739q

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***Influence of Chirality on Vibrational and Relaxational Properties of (S)- and (R,S)-Ibuprofen/methyl- $\beta$ -cyclodextrin Inclusion Complexes: An INS and QENS Study***

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**DOI:** 10.1021/jp403099a

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**DOI:** 10.1021/jp404582w



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**DOI:** 10.1021/jp404714c



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