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ANALYSIS AND EVALUATION OF IONIZATION POTENTIALS, ELECTRON AFFINITIES, AND EXCITATION ENERGIES BY THE EQUATIONS OF MOTION— GREEN'S FUNCTION METHOD

MICHAEL F. HERMAN

*Department of Chemistry
Columbia University
New York, New York*

KARL F. FREED

*The James Franck Institute and the Department of Chemistry
The University of Chicago
Chicago, Illinois*

AND

D. L. YEAGER

*Department of Chemistry
Texas A & M University
College Station, Texas*

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KINETIC THEORY OF CHEMICAL REACTIONS IN LIQUIDS

RAYMOND KAPRAL

*Department of Chemistry
University of Toronto
Toronto, Ontario, Canada*

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DIELECTRIC CONSTANTS OF FLUID MODELS: STATISTICAL MECHANICAL THEORY AND ITS QUANTITATIVE IMPLEMENTATION

G. STELL

*Departments of Mechanical Engineering and Chemistry
State University of New York
Stony Brook, New York*

G. N. PATEY

*Department of Chemistry
University of British Columbia
Vancouver, British Columbia, Canada*

AND

J. S. HØYE

*Institutt for Teoretisk Fysikk
Universitetet i Trondheim
Trondheim, Norway*

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LIGHT SCATTERING FROM THIN LIQUID FILMS

A. VRIJ, J. G. H. JOOSTEN, AND H. M. FIJNAUT

*Van't Hoff Laboratory University of Utrecht
Utrecht, The Netherlands*

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INTERACTION POTENTIALS AND GLASS FORMATION: A SURVEY OF COMPUTER EXPERIMENTS

C. A. ANGELL

*Department of Chemistry
Purdue University
West Lafayette, Indiana*

J. H. R. CLARKE

*Department of Chemistry
University of Manchester Institute of Science and Technology
Manchester, England*

AND

L. V. WOODCOCK

*Laboratory for Physical Chemistry
University of Amsterdam
Amsterdam, The Netherlands*

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LIQUIDS, GLASSES, AND THE GLASS TRANSITION: A FREE-VOLUME APPROACH

GARY S. GREEST

*Department of Physics
Purdue University
West Lafayette, Indiana*

AND

MORREL H. COHEN

*James Franck Institute and
Department of Physics
University of Chicago
Chicago, Illinois*

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