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**FIRST PASSAGE TIME PROBLEMS IN
CHEMICAL PHYSICS***

GEORGE H. WEISS, *National Cancer Institute, National
Institutes of Health, Bethesda, Maryland, U.S.A.*

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ALAN Q. ESCHENROEDER, *General Motors Corporation, Santa Barbara, California*

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ELECTRONEGATIVITY AND CHEMICAL BONDING*

RICARDO FERREIRA,[†] *Chemistry Department, Indiana University, Bloomington, Indiana, 47405*

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JAMES C. KECK,* *Massachusetts Institute of Technology,
Cambridge, Mass., U.S.A.*

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PROPERTIES OF A PARTICLE IN A ONE-DIMENSIONAL RANDOM POTENTIAL

BERTRAND I. HALPERIN,* *Laboratoire de Physique,
Ecole Normale Supérieure, Paris, France*

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H. LABHART, *Institute for Physical Chemistry,
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FRANK E. HARRIS, *Department of Chemistry, Stanford
University, Stanford, California*

and

H. H. MICHELS, *Research Laboratories, United Aircraft
Corporation, East Hartford, Connecticut*

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THE LOCAL POTENTIAL APPLIED TO INSTABILITY PROBLEMS

J. G. BALL and D. M. HIMMELBLAU, *Department of
Chemical Engineering, The University of Texas, Austin, Texas*

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LOCAL POTENTIAL METHODS IN THE STUDY OF KINETIC EQUATIONS*

G. NICOLIS, *Faculté des Sciences de l'Université Libre de
Bruxelles, Bruxelles, Belgium*

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THERMAL DIFFUSION OF HALIDES IN AQUEOUS SOLUTION

J. CHANU, *Laboratoire de Thermodynamique des Milieux Ioniques,
Muséum National d'Histoire Naturelle, Paris*

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