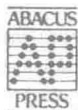


REACTION MECHANISMS IN ORGANIC CHEMISTRY

REACTION MECHANISMS IN ORGANIC CHEMISTRY

Florin Badea



ABACUS PRESS
Tunbridge Wells, Kent

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Revised, up-dated English version of the third edition of
MECANISME DE REACTIE IN CHIMIA ORGANICĂ
published in Romanian by Editura Științifică, Bucharest, in 1974

ABACUS PRESS,

Abacus House, Speldhurst Road, Tunbridge Wells, Kent, England

British Library Cataloguing in Publication Data

Badea, Florin

Reaction mechanisms in organic chemistry.

Bibl. — Index.

ISBN 0-85626-002-9

1. Title 2. Constantinescu, Take V

3. Hammel, John

547.1'39

QD258

Chemistry, Organic

Chemical reactions

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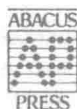
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1. Title 2. Constantinescu, Take V

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QD258

Chemistry, Organic

Chemical reactions

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