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A DENSITY MATRIX, BLOCH EQUATION DESCRIPTION OF INFRARED AND MICROWAVE TRANSIENT PHENOMENA*

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CLASSICAL-LIMIT QUANTUM MECHANICS AND THE THEORY OF MOLECULAR COLLISIONS

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SOURCES OF ERROR AND EXPECTED ACCURACY IN *AB INITIO* ONE-ELECTRON OPERATOR PROPERTIES: THE MOLECULAR DIPOLE MOMENT

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ALGEBRAIC VARIATIONAL METHODS IN SCATTERING THEORY*

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