

CURRICULUM VITAE

FRANK E. HARRIS

University of Utah
Department of Physics
115 S 1400 E Rm 201
Salt Lake City, UT 84112-0830

and

University of Florida
Quantum Theory Project
P. O. Box 118435
Gainesville, FL 32611-8435

Born : August 26, 1929, Boston MA

Education: **A. B.** (Chemistry), Harvard University, Cambridge, MA, 1951.
Ph.D. (Physical Chemistry), University of California, Berkeley, CA, 1954.

Positions:

- 1969-** Professor of Physics and Chemistry, University of Utah.
- 1998-** Resident Adjunct Professor, Quantum Theory Project and Department of Chemistry, University of Florida.
- 1984-87** President, Golden Dawn Computer Systems, Inc.
- 1978-86** Director, College of Science Computer, University of Utah.
- 1977** Visiting Professor of Chemistry, University of Hawaii.
- 1973-75** Dean of the College of Science, University of Utah.
- 1968-69** Professor of Physics, University of Utah.
- 1959-68** Assistant, Associate Professor of Chemistry, Stanford University.
- 1967** Visiting Research Scientist, United Aircraft Research Laboratories.
- 1966-70** Consultant to Quantum Theory Group of the Center for Theoretical Biology, State University of New York at Buffalo.
- 1964-74** Staff Member, International Summer Institute for Quantum Chemistry, Solid State Physics, and Quantum Biology.
- 1957-59** Sloan Foundation Fellow.
- 1956-59** Assistant Professor of Chemistry, University of California, Berkeley.
- 1955** Summer Visitor, General Electric Research Laboratory.
- 1953-56** Instructor in Chemistry, Harvard University.
- 1952-53** National Science Foundation Predoctoral Fellow

Fellow, American Institute of Chemists

Fellow, American Physical Society

Editorial Board: International Journal of Quantum Chemistry, 2003-

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