General Atomic and Molecular Electronic Structure System

GAMESS User's Guide
Department of Chemistry
Iowa State University
Ames, IA 50011

literature citation:

"General Atomic and Molecular Electronic Structure System"
M.W.Schmidt, K.K.Baldridge, J.A.Boatz, S.T.Elbert,
M.S.Gordon, J.H.Jensen, S.Koseki, N.Matsunaga,
K.A.Nguyen, S.J.Su, T.L.Windus, M.Dupuis, J.A.Montgomery

"Advances in electronic structure theory:
GAMESS a decade later"
M.S.Gordon, M.W.Schmidt
in "Theory and Applications of Computational Chemistry,
the first forty years" C.E.Dykstra, G.Frenking, K.S.Kim,

http://www.msg.ameslab.gov/GAMESS/GAMESS.html

Graphical display of results is possible using MacMolPlt, a
back end visualizer, available for Apple computers only.
MacMolPlt can be downloaded freely at the web site above.

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Capabilities

A wide range of quantum chemical computations are possible using GAMESS, which

1. Calculates RHF, UHF, ROHF, GVB, or MCSCF self-consistent field molecular wavefunctions.

2. Calculates the electron correlation energy correction for these SCF wavefunctions using Density Functional Theory (DFT), Configuration Interaction (CI), Many Body Perturbation Theory (MP2), coupled-cluster (CC) or equation of Motion CC (EOM-CC) methodologies. (see summary table below).

3. Calculates semi-empirical MNDO, AM1, or PM3 models using RHF, UHF, ROHF, or GVB wavefunctions.

4. Calculates analytic energy gradients for any of the SCF or DFT wavefunctions, closed or open shell MP2, or closed shell reference-based CI.

5. Optimizes molecular geometries using the energy gradient, using internal or Cartesian coordinates.

6. Searches for saddle points (transition states) on the potential energy surface.

7. Computes the energy hessian, and thus normal modes, vibrational frequencies, and IR intensities. Raman intensities are a follow-up option.

8. Obtains anharmonic vibrational frequencies and intensities (fundamentals or overtones).

9. Traces the intrinsic reaction path from the saddle point towards products, or back to reactants.

10. Traces gradient extremal curves, which may lead from one stationary point such as a minimum to another, which might be a saddle point.

11. Follows the dynamic reaction coordinate, a classical mechanics trajectory on the potential energy surface.

12. Computes excited state energies, wavefunctions, and transition dipole moments at various levels
   a. SCF (e.g. ROHF or MCSCF)
b. CIS (RHF plus single excitations)
c. more general CI functions
d. Equation of Motion Coupled Cluster
   with analytic gradients for SCF, CIS, and GUGA CI.

13. Evaluates relativistic effects, including
   a. scalar corrections, via 3rd order Douglas-Kroll transformations. Gradients are available.
   b. spin-orbit coupling matrix elements and the resulting spin-mixed wavefunctions.

14. Evaluates the molecular linear polarizability and the first and second order hyperpolarizabilities for all wavefunctions, by applying finite electric fields.

15. Evaluates both the static and frequency dependent polarizabilities for various non-linear optical processes, by analytic means, for RHF wavefunctions. Nuclear derivatives of the polarizabilities lead to analytic Raman spectra, also for RHF wavefunctions. The imaginary frequency dependent polarizabilities can also be obtained.

16. Obtains localized orbitals by the Foster-Boys, Edmiston-Ruedenberg, or Pipek-Mezey methods, with optional SCF or MP2 energy analysis of the LMOs.

17. Calculates the following molecular properties:
   a. dipole, quadrupole, and octopole moments
   b. electrostatic potential
   c. electric field and electric field gradients
   d. electron density and spin density
   e. Mulliken and Lowdin population analysis
   f. virial theorem and energy components
   g. Stone's distributed multipole analysis

18. Models solvent effects by
   a. effective fragment potentials (EFP)
   b. polarizable continuum model (PCM)
   c. surface and simulation of volume polarization for electrostatics (SS(V)PE)
   d. conductor-like screening model (COSMO)
   e. self-consistent reaction field (SCRF)

19. Performs all-electron calculations based on the Fragment Molecular Orbital (FMO) method.

20. When combined with the add-on TINKER molecular mechanics program, performs Surface IMOMM or

22. When combined with the add-on NBO program, performs Natural Bond Order analyses. This program is available at http://www.chem.wisc.edu/~nbo5, for a modest license fee.

Many of these calculations may be performed in parallel!
A quick summary of the current program capabilities is given below.

<table>
<thead>
<tr>
<th>SCFTYP</th>
<th>RHF</th>
<th>ROHF</th>
<th>UHF</th>
<th>GVB</th>
<th>MCSCF</th>
</tr>
</thead>
<tbody>
<tr>
<td>SCF energy</td>
<td>CDFP</td>
<td>CDP</td>
<td>CDP</td>
<td>CDP</td>
<td>CDFP</td>
</tr>
<tr>
<td>SCF analytic gradient</td>
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<td>CDP</td>
<td>CDP</td>
<td>CDP</td>
<td>CDFP</td>
</tr>
<tr>
<td>SCF analytic Hessian</td>
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<td>CDP</td>
<td>-</td>
<td>CDP</td>
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<tr>
<td>CI energy</td>
<td>CDP</td>
<td>CDP</td>
<td>-</td>
<td>CDP</td>
<td>CDP</td>
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<tr>
<td>CI gradient</td>
<td>CD</td>
<td>-</td>
<td>-</td>
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</tr>
<tr>
<td>MP2 energy</td>
<td>CDFP</td>
<td>CDP</td>
<td>CDP</td>
<td>-</td>
<td>CDP</td>
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<tr>
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<td>CDFP</td>
<td>DP</td>
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<tr>
<td>EOMCC excitations</td>
<td>CD</td>
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<td>DFT energy</td>
<td>CDFP</td>
<td>CDP</td>
<td>CDP</td>
<td>-</td>
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<td>CDP</td>
<td>CDP</td>
<td>-</td>
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<tr>
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<td>yes</td>
<td>yes</td>
<td>yes</td>
<td>-</td>
</tr>
<tr>
<td>MOPAC gradient</td>
<td>yes</td>
<td>yes</td>
<td>yes</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

C= conventional storage of AO integrals on disk  
D= direct evaluation of AO integrals whenever needed  
F= Fragment Molecular Orbital enabled  
P= parallel execution

Numerical gradients and fully or partly numerical Hessians are available for any energy or gradient in this table.
History of GAMESS

GAMESS was put together from several existing quantum chemistry programs, particularly HONDO, by the staff of the National Resources for Computations in Chemistry. The NRCC project (1 Oct 77 to 30 Sep 81) was funded by NSF and DOE, and was limited to the field of chemistry. The NRCC staff added new capabilities to GAMESS as well. Besides providing public access to the code on the CDC 7600 at the site of the NRCC (the Lawrence Berkeley Laboratory), the NRCC made copies of the program source code (for a VAX) available to users at other sites. The original citation for this program was

M. Dupuis, D. Spangler, and J. J. Wendoloski
National Resource for Computations in Chemistry
Software Catalog, University of California:
Berkeley, CA (1980), Program QG01

This manual is a completely rewritten version of the original documentation for GAMESS. Any errors found in this documentation, or the program itself, should not be attributed to the original NRCC authors.

The present version of the program has undergone many changes since the NRCC days. This occurred at North Dakota State University from 1982 up to 1992, and now continues at Iowa State University to the present.

It would be difficult to overestimate the contributions Michel Dupuis has made to this program, in its original form, and since. This includes the donation of code from HONDO, and numerous suggestions for other improvements.

The continued development of this program from 1982 on can be directly attributed to the nurturing environment provided by Professor Mark Gordon, at North Dakota State and then Iowa State University.

It is important to also single out Professor Emeritus Klaus Ruedenberg of Iowa State University, whose group is responsible for the determinant technology lying underneath the MCSCF programs in GAMESS.

Even when our students and postdocs leave Iowa State, many continue to make contributions to GAMESS. In addition, we have also included many codes developed in other groups over the years, so that the list of authors of GAMESS is actually much longer than the author list of the
Funding of many of the developments in GAMESS from 1982 to the present time was, and is provided by the Air Force Office of Scientific Research. This has always been the backbone of the support for GAMESS.

In late 1987, NDSU and IBM reached a Joint Study Agreement. One goal of this JSA was the development of a version of GAMESS that was vectorized for the IBM 3090's Vector Facility, which was accomplished by the fall of 1988. This phase of the JSA led to a program which is also considerably faster in scalar mode as well. The second phase of the JSA, which ended in 1990, was to enhance GAMESS' scientific capabilities. These additions include analytic hessians, ECPs and MCPs, MP2, spin-orbit coupling and radiative transitions, and so on. Everyone who uses the current version of GAMESS owes thanks to IBM in general, and Michel Dupuis of IBM Kingston in particular, for their sponsorship of GAMESS during this JSA.

During the first six months of 1990, Digital awarded a Innovator's Program grant to NDSU. The purpose of this grant was to ensure GAMESS would run on the DECstation, and to develop graphical display programs. As a result, the companion programs MOLPLT, PLTORB, DENDIF, and MEPMAP were modernized for the X-windows environment, and interfaced to GAMESS. These programs now run under the X-windows environments, and many other X-windows environments as well. The ability to visualize the molecular structures, orbitals, and electrostatic potentials is a significant improvement. These graphics programs eventually formed the nucleus of the Macintosh program MacMolPlt.

Parallelization of GAMESS began in 1991, with most of the early work and design strategy done by Theresa Windus. This benefited greatly from the ARPA sponsorship of the Touchstone Delta experimental computer. Message passing used the TCGMSG library of Robert Harrison in the early years, up to 1999. Parallelization of GAMESS has turned into a multi-year process as detailed below.

The DoD awarded a CHSSI grant to ISU in 1996 to extend that scalability of existing parallel methods, and more importantly develop new techniques. This brought Graham Fletcher on board as a postdoc, and led to the introduction of the Distributed Data Interface (DDI) programming model.
The first version of DDI, written at ISU, was used from June 1999 to May 2004. Ryan Olson, with help from Alistair Rendell of Australian National University, rewrote DDI entirely in C, adding optimizations for the commonplace SMP nodes, especially System V memory use. Dmitri Fedorov of the National Institute for Advanced Industrial Science and Technology added the concept of subgroups at the same time. This combined new version of DDI has been the message passing support layer for GAMESS since May 2004.

The DoE awarded a SciDAC grant to ISU in 2002 to enable additional scientific capabilities in GAMESS, with emphasis on scalable algorithms. To date, this has supported parallelization of the EFP solvent molecule, and new codes for analytic MCSCF Hessians, and open shell MP2 gradients.

Some summary of these various grants and initiatives is in order. The 1982 version of GAMESS contained roughly 80,000 lines of FORTRAN code, implementing the present five wavefunction types, and analytic nuclear gradients for each, enabling geometry optimization and transition state search, and numerically differentiated frequencies. The only electron correlation method available was GUGA based CI computation. All computations were in the gas phase.

By 2005, GAMESS had grown to roughly 650,000 lines of FORTRAN. Analytic hessian computation is now routine at the SCF levels. Electron correlation is now treated with direct determinant CI codes, and in addition perturbation theory, density functional, or coupled cluster methods (with analytic gradients for some of these) may be used. New AO integral codes, including effective core potentials are used, and direct AO integral computation is possible. Discrete and continuum models for solvated molecules are provided, and there is an associated program for surface chemistry. Additional chemistry runs are provided, such as reaction paths and dynamical trajectories, IR and Raman spectra, anharmonic vibrational corrections, static or frequency dependent polarizabilities, transition moments, and spin-orbit couplings. Scalar relativistic corrections can be applied to any computation. Improvements or complete rewrites have been made for geometry searches, SCF convergers, internal coordinates, ease of use, available basis sets, and so on. The majority of these computations can be run on parallel computers.

The rest of this section gives more specific credit to the sources of various parts of the program.
GAMESS is a synthesis, with many major modifications, of several programs. A large part of the program originate from HONDO 5.

For sp basis functions, modified Gaussian76 s,p,L shell code is used. Both the sp rotated axis integrals and the sp gradient packages were modified in 2001 by Jose Maria Sierra of Synstar Computer Services in Madrid, Spain. The sp integral routines were modified in 2003 and in 2004 by Kazuya Ishimura of the Institute for Molecular Science to use McMurchie-Davidson quadratures for the basic axes-1 integrals, after which they are rotated ala Hehre/Pople. For spd functions, the s,p,d,L shell rotated axis code written by Kazuya Ishimura of the Institute for Molecular Science is used. For integral quartets with higher angular momentum, the s,p,d,f,g Electron Repulsion Integral Calculator (ERIC) code written by Graham Fletcher at NASA/Eloret in 2004 is used, provided the total angular momentum of the quartet is no more than 5. Both rotated axis codes, the sp gradient code, and ERIC share a common, fully accurate evaluation of $F_m(t)$ integrals, and have been tested for very small (down to 0.005) and very large (1.0d+11) Gaussian exponents. The Rys polynomial program of Michel Dupuis is used to handle the general integral case: s,p,d,f,g, or L shells. HONDO 1e- and 2e- Rys routines were redimensioned to handle up to g shells by Theresa Windus at North Dakota State University in 1991.

Any sp gradient integrals are done with Jose Sierra's modified version of the Gaussian80 code due to Schlegel. The spdfg gradient package consists of Michel Dupuis' Rys Polynomial code, and was adapted into GAMESS by Brett Bode at Iowa State University in 1994.

The use of quantum fast multipole methods for avoiding long range integral evaluation in large molecules was programmed by Cheol Choi at Iowa State and at Kyungpook National University, and included in GAMESS in 2001.

The ECP code goes back to Louis Kahn, with gradient modifications originally made by K.Kitaura, S.Obara, and K.Morokuma at IMS in Japan. The code was adapted to HONDO by Stevens, Basch, and Krauss, from whence Kiet Nguyen adapted it to GAMESS at NDSU. Modifications for f functions were made by Drora Cohen and Brett Bode. This code was completely rewritten to use spdfg basis sets, to exploit shell structure during integral evaluation, and to
add the capability of analytic second derivatives by Brett Bode at ISU in 1997-1998. Jose Sierra of Synstar removed the last few bugs from this in 2003.

The MCP code originates from the Huzinaga group at the University of Alberta, and was interfaced to GAMESS in 2003 by Mariusz Klobukowski. A large number of model core potentials with their associated valence basis sets were added by Mariusz in 2005.

Changes in the manner of entering the basis set, and the atomic coordinates (including Z-matrix forms) are due to Jan Jensen at North Dakota State University.

The direct SCF implementation was done at NDSU, guided by a pilot code for the RHF case by Frank Jensen.

The UHF code was taught to do high spin ROHF by John Montgomery at United Technologies, who extended DIIS use to ROHF and the one pair GVB case.

The GVB code is a heavily modified version of GVBONE.

The SCF for Molecular Interactions option was added to GAMESS in 1997 by Antonino Famulari, during a summer visit from the University of Milan. This two fragment code was replaced with a multi-fragment code from Maurizio Sironi of the University of Milan in 2004.

The Direct Inversion in the Iterative Subspace (DIIS) convergence procedure was implemented by Brenda Lam (then at the University of Houston) in 1986, for RHF and UHF functions. Additional GVB-DIIS cases were programmed by Galina Chaban at ISU. The approximate second order SCF converger was implemented by Galina Chaban at Iowa State University in 1995, and was provided for RHF, ROHF, GVB, and MCSCF cases. The FULLNR and FOCAS MCSCF convergers were contributed by Michel Dupuis from his HONDO program. A parallel implementation of the FULLNR converger was written by Graham Fletcher at Eloret in 2002. The Jacobi orbital rotation scheme for MCSCF orbital optimization was written by Joe Ivanic and Klaus Ruedenberg at Iowa State University in 2001.

The Ames Laboratory determinant full CI code was written by Joe Ivanic and Klaus Ruedenberg. As befits code written by an Australian living in Iowa, it was interfaced to GAMESS during an extremely cordial visit to Australia National University in January 1998. An update by Joe in
October 2000 exploits Abelian point group symmetry. A general CI program based on selected determinants was added by Joe and Klaus in July 2001. After moving from Ames Laboratory at ISU to the Advanced Biomedical Computing Center of the National Cancer Institute-Frederick, Fort Detrick, Joe wrote a determinant based program for second order CI, in 2002. In early 2003, Joe added the Occupation Restricted Multiple Active Space determinant CI program, again written at NCI.

The GUGA CI is based on Brooks and Schaefer's unitary group program which was modified to run within GAMESS, using a Davidson eigenvector method written by Steve Elbert.

Programming of the GUGA analytic CI gradient was done by Simon Webb in 1996 at Iowa State University.

The CIS gradient program was written in 2003 by Simon Webb of the Advanced Biomedical Computing Center of the National Cancer Institute in Frederick. Transition moments were added by Simon and Pooja Arora in June 2005.

The sequential MP2 and UMP2 energy code was adapted from HONDO in 1994 by Nikita Matsunaga at ISU. Nikita programmed the RMP open shell energy in 1992. The ZAPT open shell energy was programmed by Rob Bell in 1999. The serial closed shell MP2 gradient code is also from HONDO, and was adapted to GAMESS in 1995 by Simon Webb and Nikita Matsunaga. In 1996, Simon Webb added the frozen core gradient option at ISU. The parallel closed shell MP2 code is a descendant of work for GAMESS-UK by Graham Fletcher, Alistair Rendell, and Paul Sherwood at Daresbury. This was adapted to GAMESS at ISU by Graham Fletcher in 1999.

Serial and parallel codes for the spin unrestricted UMP2 gradient were programmed by Christine Aikens at ISU, in 2002. Christine Aikens added a parallel spin-restricted open shell (ZAPT) gradient code in 2005. Haruyuki Nakano from the University of Tokyo interfaced his multireference MCQDPT code (based on CSFs) to GAMESS during a 1996 visit to ISU. Parallelization of the Tokyo multireference PT code was done by Hiroaki Umeda at Mie University, and included into GAMESS in 2001. A determinant based equivalent to MRMP/MCQDPT was programmed in 2005 by Joe Ivanic of the National Cancer Institute.

The grid-free DFT energy and gradient code was written by Kurt Glaesemann at Iowa State University, starting from the code of Almlof and Zheng, adding four center overlap
integrals, a gradient program, developing the auxiliary basis option, and adding some functionals. This was included in GAMESS in 1999.

The grid based DFT program was written in 2001 at the University of Tokyo, by Takao Tsuneda, Muneaki Kamiya, Susumu Yanagisawa, and Dmitri Fedorov. Many improvements such as use of symmetry and initial coarse grid during the numerical quadrature, functional development and coding, and the ability to run in parallel come from this group. The original program prior to these numerous changes is from Nevin Oliphant, Hideo Sekino, and Rod Bartlett at QTP.

Incorporation of enough MOPAC version 6 routines to run PM3, AM1, and MNDO calculations from within GAMESS was done by Jan Jensen at North Dakota State University.

The numerical force constant computation and normal mode analysis was adapted from Andy Komornicki's GRADSCF program, with decomposition of normal modes in internal coordinates written at NDSU by Jerry Boatz.

The code for the analytic computation of RHF Hessians was contributed by Michel Dupuis of IBM from HONDO 7. High and low spin restricted open shell CPHF code was written at NDSU in 1989. The TCSCF CPHF code is the result of a collaboration between NDSU and John Montgomery, then at United Technologies, in 1990. Analytic IR intensities and polarizabilities (during hessian runs) were programmed by Simon Webb at ISU in 1995. Analytic Hessians for MCSCF wavefunctions based on determinants were coded, and enabled for parallel execution, by Tim Dudley at ISU, and included into GAMESS in April 2004.

Code for Raman intensity prediction was written at Tokyo Metropolitan University in April 2000.

The vibrational SCF and MP2 anharmonic frequency code for fundamental modes and overtones was written by Galina Chaban, Joon Jung, and Benny Gerber at U.California-Irvine and Hebrew University of Jerusalem, and included in GAMESS in 2000. The solver was modified to perform degenerate perturbation theory for more accurate results by Nikita Matsunaga at Long Island University in 2001.

Most geometry search procedures in GAMESS (NR, RFO, QA, and CONOPT) were developed by Frank Jensen of the University of Southern Denmark. These methods are adapted
to use GAMESS symmetry, and Cartesian or internal coordinates.

The non-gradient optimization so aptly described as "trudge" was adapted from HONDO 7 by Mariusz Klobukowski at U.Alberta, who added the option for CI optimizations. Numerical differentiation of the energy to obtain gradients and Hessians was programmed by Ryan Olson at ISU in 2003.

The intrinsic reaction coordinate pathfinder was written at North Dakota State University, and modified later for new integration methods by Kim Baldridge. The Gonzales-Schelegel IRC stepper was incorporated by Shujun Su at Iowa State, based on pilot code from Frank Jensen.

The code for the Dynamic Reaction Coordinate was developed by Tetsuya Taketsugu at Ochanomizu U. and U. of Tokyo, and added to GAMESS by him at ISU in 1994.

The two algorithms for tracing gradient extremals were programmed by Frank Jensen at the University of Southern Denmark.

The program for Monte Carlo generation of trial structures along with a simulated annealing protocol was written by Paul Day at Wright-Patterson Air Force Base. Modifications to this were made by Pradipta Bandyopadhyay at ISU, and the code was included in 2001.

The surface scanning option was implemented by Richard Muller at the University of Southern California.

Static polarizabilities for any type of energy value are based on a code from Henry Kurtz of the University of Memphis. This uses a numerical differentiation based on application of finite electric fields. The program was added in 1992, and was modified by Sanka Ghosh to produce all tensor components in 2005.

Henry Kurtz' program for the fully analytic calculation of static and frequency dependent polarizabilities for NLO properties for closed shell systems was included in 1994, based on a MOPAC implementation by Prakashan Korambath at U. Memphis.

An extended TDHF package for the analytic computation of static and frequency dependent polarizabilities, and also their nuclear derivatives, plus Raman and hyperRaman spectra prediction was written by Olivier Quinet and Benoit
Champagne at the Facultes Universitaires Notre-Dame de la Paix, and coworker Bernard Kirtman at UC-Santa Barbara. Financial support for this was provided by Belgium. This package was added to GAMESS in February 2005.

Ivana Adamovic programmed the imaginary frequency polarizability computation for closed shell functions in 2005, at ISU.

Edmiston-Ruedenberg energy localization is done with a version of the ALIS program "LOCL", modified at NDSU to run inside GAMESS. Foster-Boys localization is based on a highly modified version of QCPE program 354 by D.Boerth, J.A.Hasmall, and A.Streitweiser. John Montgomery implemented the Pipek/Mezey population localization. The LCD SCF decomposition and the MP2 decomposition were written by Jan Jensen at Iowa State in 1994.

Point Determined Charges were implemented by Mark Spackman at the University of New England, Australia.

Delocalized internal coordinates were implemented by Jim Shoemaker at the Air Force Institute of Technology in 1997, and put online in GAMESS by Cheol Choi at ISU after further improvements in 1998.

The Morokuma decomposition was implemented by Wei Chen at Iowa State University, in 1995.

The radiative transition moment and effective nuclear charge spin-orbit coupling modules were written by Shiro Koseki at North Dakota State University in 1990.

The full Breit-Pauli spin-orbit coupling integral package was written by Thomas Furlani. This code was incorporated into GAMESS by Dmitri Fedorov at Iowa State University in 1997, who generalized the spin-orbit coupling matrix element code generously provided by Thomas Furlani (restricted to an active space of two electrons in two orbitals), with assistance from visits to ISU by Thomas Furlani and Shiro Koseki. Dmitri Fedorov has since generalized the full two electron approach to allow for any spins, for more than two spin multiplicities at a time, and a partial treatment of the the two electron terms that runs in time similar to the one electron operator. Space and spin symmetries are exploited to speed up the runs. Dmitri Fedorov programmed the SO-MCQDPT options at the University of Tokyo in 2001.
Inclusion of relativistic effects by means of the Douglas-Kroll transformation was developed to the third order by Takahito Nakajima and Kimihiko Hirao at the University of Tokyo. It was implemented into GAMESS by Takahito Nakajima and Dmitri Fedorov, including energy, semi-analytic nuclear gradient, and spin-orbit coupling (at the first order, DK1). The program was included with GAMESS in November 2003.

Inclusion of relativistic effects by the Relativistic scheme of Elimination of Small Components (RESC) method, was developed by Takahito Nakajima and Kimihiko Hirao at the University of Tokyo. This code was written by Takahito Nakajima and consequently adapted into GAMESS by Dmitri Fedorov, who extended the methodology in March 2000 to the computation of gradients. RESC provides both scalar (spin free) and vector (spin-dependent) relativistic corrections.

The Normalized Elimination of Small Components (NESC) was programmed by Dmitri Fedorov at ISU and the University of Tokyo. Special thanks are due to Kenneth Dyall for his assistance in providing check values. Extension of NESC to include gradient computation was also done by Dmitri.

Development of the EFP method began in the group of Walt Stevens at NIST's Center for Advanced Research in Biotechnology (CARB) in 1988. Walt is the originator of this method, and has provided both guidance and some early financial support to ISU for its continued development. Mark Gordon's group's participation began in 1989-90 as discussions during a year Mark spent in the DC area, and became more serious in 1991 with a visit by Jan Jensen to CARB. At this time the method worked for the energy, and gradient with respect to the ab initio nuclei, for one fragment only. Jan has assisted with most aspects of the multi-fragment development since. Paul Day at NDSU and ISU derived and implemented the gradient with respect to fragments, and programmed EFP geometry optimization, from 1992-1994. Wei Chen at ISU debugged many parts of the EFP energy and gradient, developed the code for following IRCs, improved geometry searches, and fitted much more accurate repulsive potentials, from 1995-1996. Simon Webb at ISU programmed the current self-consistency process for the induced dipoles in 1994. The EFP method was sufficiently developed, tested, and described, to be released in September 1996, with an RHF level potential for water. Code for charge penetration was added by Mark Freitag in 2001. Ivana Adamovic included a DFT level EFP for water in
Parallelization of the EFP codes was done by Heather Netzloff in 2005.

The second EFP theory (called EFP2) was begun in 1996 by Jan Jensen, who programmed an analytic formula for the exchange repulsion. Hui Li replaced this with a faster, more accurate code in 2005. Ivana Adamovic programmed a dispersion term for EFP2 in 2005. Hui Li added the charge transfer term for EFP2 in 2005.

Two other methods using the EFP model are available. A combination of EFP + PCM energies (an onion-like solution model) was programmed by Pradipta Bandyopadhyay in 2000. The use of EFPs to model biological systems, including a boundary across a covalent bond, was coded at the University of Iowa in 2000, by Jan Jensen, Visvaldas Kairys, and Hui Li.

The SCRF solvent model was implemented by Dave Garmer at CARB, and was adapted to GAMESS by Jan Jensen and Simon Webb at Iowa State University.

The COSMO model was developed by Andreas Klamt and Kim Baldridge, at the San Diego Supercomputer Center. It was included into GAMESS by Laura Gregerson in March 2000 during a visit to Ames.

The PCM code originated in the group of Jacopo Tomasi at the University of Pisa. Benedetta Mennucci was instrumental in interfacing the PCM code to GAMESS, in 1997, and answering many technical questions about the code, the methodology, and the documentation. In 2000, Benedetta Mennucci provided code implementing an improved IEF solver for the PCM surface charges. The changes to implement iterative solution of the PCM equations for large molecules, and to provide an accurate nuclear gradient were carried out by Hui Li and Jan Jensen at the University of Iowa in 2001-2004, along with the parallelization. This includes implementation of two new surface tessellation schemes, GEPOL-AS and GEPOL-RT. Hui and Jan also implemented the Conductor-PCM method, and extended the PCM methodology to all types of SCF functions.

The Surface and Volume Polarization for Electrostatics continuum solvation model is written by Dan Chipman of Notre Dame University, using several integral routines written by Michel Dupuis for the SVP model included in HONDO. The SVP model was added to GAMESS in June 2005.
The Coupled-Cluster (CC) and Equation of Motion Coupled-Cluster (EOMCC) programs included in GAMESS are due to Piotr Piecuch, Karol Kowalski, and Marta Wloch of Michigan State University (MSU), and Stanislaw A. Kucharski and Monika Musial of the University of Silesia. Testing was done by Michael McGuire of MSU. Jeffrey Gour of MSU helped test the correctness of the codes by comparisons with independent programs written in collaboration with Marta Wloch, Karol Kowalski, and Piotr Piecuch for general one-particle basis sets. In addition to a number of standard CC and EOMCC methods, including the popular CCSD, CCSD(T), and EOMCCSD approaches, the CC codes incorporated in GAMESS are capable of performing renormalized (R) and completely renormalized (CR) CCSD[T] and CCSD(T) calculations for the ground state, and completely renormalized EOMCCSD(T) (CR-EOMCCSD(T)) calculations for excited states. The combined corrections due to triply and quadruply excited clusters are available in the factorized forms of the CCSD(TQ), renormalized CCSD(TQ), and completely renormalized CCSD(TQ) models. The one-body reduced density matrices, dipole moments, transition dipole moments, and oscillator strengths are available at the CCSD and EOMCCSD levels. The ground-state CC, R-CC, and CR-CC programs were initially incorporated into GAMESS in May 2002. The excited-state EOMCC and CR-EOMCC programs were incorporated in April 2004. Quadruples corrections and CCSD/EOM-CCSD density matrices were added in June 2005. All of these programs were developed with the support of the US Department of Energy, Office of Basic Energy Sciences, SciDAC Computational Chemistry Program and the Chemical Sciences, Geosciences, and Biosciences Division. Additional support has been provided by the NSF's ITR program and the Alfred P. Sloan Foundation.

The GIAO computation of NMR properties for closed shell molecules was programmed by Mark Freitag at Iowa State University, and included in GAMESS in November 2003.

The code for the Fragment Molecular Orbital (FMO) method incorporated and distributed as a part of the standard GAMESS package since May 2004 is being developed at the National Institute of Advanced Industrial Science and Technology (AIST, Japan) by Dmitri Fedorov and Kazuo Kitaura. The FMO method is the successor of the EDA scheme developed by K. Kitaura and K. Morokuma (known in GAMESS as Morokuma-Kitaura decomposition), however, the FMO code was written independently. In GAMESS only the full FMO method is incorporated whereas in the literature one can also find a simplified approach suited for molecular crystals. Since
"FMO" is also used to mean "Frontier Molecular Orbitals" and the concept of fragments is also introduced in the EFP method (see above), it is stressed here that the FMO method bears no relation to either of the two methods, that is to say, it is independent of the two, but might be combined with either of them in the future just as EFPS are used in e.g. RHF.

Many of the options just mentioned have been programmed to run in parallel, on systems ranging from Linux clusters to high-end parallel systems. The same software interface sits between the quantum chemistry in GAMESS and any such hardware, namely the Distributed Data Interface (DDI). This implements a mechanism for using the memory of the entire system to store the large arrays appearing in quantum chemistry codes. The first version of DDI was due to Graham Fletcher and Mike Schmidt, introduced in 1999. The second version of DDI is due to Ryan Olson of ISU, and Alistair Rendell of the Australian National University, and includes optimizations for SMP systems, along with other improvements for some high end systems. The second version also includes the 'group' scheme, presently used only in FMO jobs. This DDI was introduced into GAMESS in April 2004, with public release in June 2004.
**Distribution Policy**

To get a copy, please fill out the application form available at
http://www.msg.ameslab.gov/GAMESS/GAMESS.html

Persons receiving copies of GAMESS are requested to acknowledge that they will not make copies of GAMESS for use at other sites, or incorporate any portion of GAMESS into any other program, without receiving permission to do so from ISU. If you know anyone who wants a copy of GAMESS, please refer them to the web site above, for the most up to date version available.

No large program can ever be guaranteed to be free of bugs, and GAMESS is no exception. If you would like to receive an updated version (fewer bugs, and with new capabilities), simply return to the web site mentioned. You should probably allow a half year or so to pass for enough significant changes to accumulate. The web page always contains a short synopsis of the most recent changes.
Input Philosophy

Input to GAMESS may be in upper or lower case. All input groups begin with a $ sign in column 2, meaning exactly column 2 or else it is not detected, followed by a name identifying that group. There are three types of input groups in GAMESS:

1. A pseudo-namelist, free format, keyword driven group. Almost all input groups fall into this first category.

2. A free format group which does not use keywords. The first line of these will contain only the group name, followed by several lines of positional data usually with no keywords, and a last line containing "$END" only. The only members of this category are $DATA, $ECP, $MCP, $GCILST, $POINTS, $STONE, and the EFP related data $EFRAG, $FRAGNAME, $FRGRPL, and $DAMPGS.

3. Formatted data. This data is NEVER typed by the user, but rather is generated in the correct format by some earlier GAMESS run. Like category 2, the first line contains only the group name, and the last line is a separate $END line.

Type 1 groups may have keyword input on the same line as the group name, and the $END may appear anywhere.

Because each group has a unique name, the groups may be given in any order desired. In fact, multiple occurrences of category 1 groups are permissible.

* * *

Most of the groups can be omitted if the program defaults are adequate. An exception is $DATA, which is always required. A typical free format $DATA group is

```
$DATA
STO-3G test case for water
CNV      2
OXYGEN    8.0
        STO  3
HYDROGEN  1.0   -0.758    0.0    0.545
        STO  3
```
Here, position is important. For example, the atom name must be followed by the nuclear charge and then the x,y,z coordinates. Note that missing values will be read as zero, so that the oxygen is placed at the origin. The zero Y coordinate must be given for the hydrogen, so that the final number is taken as Z.

The free format scanner code used to read $DATA is adapted from the ALIS program, and is described in the documentation for the graphics programs which accompany GAMESS. Note that the characters ;>! mean something special to the free format scanner, and so use of these characters in $DATA and $ECP should probably be avoided.

Because the default type of calculation is a single point (geometry) closed shell SCF, the $DATA group shown is the only input required to do a RHF/STO-3G water calculation.

* * *

As mentioned, the most common type of input is a namelist-like, keyword driven, free format group. These groups must begin with the $ sign in column 2, but have no further format restrictions. You are not allowed to abbreviate the keywords, or any string value they might expect. They are terminated by a $END string, appearing anywhere. The groups may extend over more than one physical card. In fact, you can give a particular group more than once, as multiple occurrences will be found and processed. We can rewrite the STO-3G water calculation using the keyword groups $CONTRL and $BASIS as

$CONTRL SCFTYP=RHF RUNTYP=ENERGY $END
$BASIS GBASIS=STO NGAUSS=3 $END
$DATA
STO-3G TEST CASE FOR WATER
Cnv 2

Oxygen 8.0 0.0 0.0 0.0
Hydrogen 1.0 -0.758 0.0 0.545
$END

Keywords may expect logical, integer, floating point, or string values. Group names and keywords never exceed 6 characters. String values assigned to keywords never
exceed 8 characters. Spaces or commas may be used to separate items:

$\text{CONTRL } \text{MULT}=3 \text{ SCFTYP=UHF,TIMLIM}=30.0 \text{ $END$}

Floating point numbers need not include the decimal, and may be given in exponential form, i.e. \text{TIMLIM}=30, \text{TIMLIM}=3.E1, and \text{TIMLIM}=3.0D+01 are all equivalent.

Numerical values follow the FORTRAN variable name convention. All keywords which expect an integer value begin with the letters I-N, and all keywords which expect a floating point value begin with A-H or O-Z. String or logical keywords may begin with any letter.

Some keyword variables are actually arrays. Array elements are entered by specifying the desired subscript:

$\text{SCF NO}(1)=1 \text{ NO}(2)=1 \text{ $END$}

When contiguous array elements are given this may be given in a shorter form:

$\text{SCF NO}(1)=1,1 \text{ $END$}

When just one value is given to the first element of an array, the subscript may be omitted:

$\text{SCF NO}=1 \text{ NO}(2)=1 \text{ $END$}

Logical variables can be .\text{TRUE.} \text{ or .FALSE.} \text{ or .T. or .F.} \text{ The periods are required.}

The program rewinds the input file before searching for the namelist group it needs. This means that the order in which the namelist groups are given is immaterial, and that comment cards may be placed between namelist groups.

Furthermore, the input file is read all the way through for each free-form namelist so multiple occurrences will be processed, although only the LAST occurrence of a variable will be accepted. Comment fields within a free-form namelist group are turned on and off by an exclamation point (!). Comments may also be placed after the $\text{END}'s$ of free format namelist groups. Usually, comments are placed in between groups,

$\text{CONTRL } \text{SCFTYP=RHF RUNTYP=GRADIENT $END}$
The second $CONTRL is not read, because it does not have a blank and a $ in the first two columns. Here a careful user has executed a CHECK job, and is now running the real calculation. The CHECK card is now just a comment line.

***

The final form of input is the fixed format group. These groups must be given IN CAPITAL LETTERS only! This includes the beginning $NAME and closing $END cards, as well as the group contents. The formatted groups are $VEC, $HESS, $GRAD, $DIPDR, and $VIB. Each of these is produced by some earlier GAMESS run, in exactly the correct format for reuse. Thus, the format by which they are read is not documented in section 2 of this manual.

***

Each group is described in the Input Description section. Fixed format groups are indicated as such, and the conditions for which each group is required and/or relevant are stated.

There are a number of examples of GAMESS input given in the Input Examples section of this manual.
Input Checking

Because some of the data in the input file may not be processed until well into a lengthy run, a facility to check the validity of the input has been provided. If EXETYP=CHECK is specified in the $CONTRL group, GAMESS will run without doing much real work so that all the input sections can be executed and the data checked for correct syntax and validity to the extent possible. The one-electron integrals are evaluated and the distinct row table is generated. Problems involving insufficient memory can be identified at this stage. To help avoid the inadvertent absence of data, which may result in the inappropriate use of default values, GAMESS will report the absence of any control group it tries to read in CHECK mode. This is of some value in determining which control groups are applicable to a particular problem.

The use of EXETYP=CHECK is HIGHLY recommended for the initial execution of a new problem.
Program Limitations

GAMESS can use an arbitrary Gaussian basis of spdfg type for computation of the energy or gradient. Some restrictions apply, for example, analytic hessians are limited to spd basis sets.

This program is limited to a total of 2,000 atoms. The total number of symmetry unique basis set shells cannot exceed 5,000, containing no more than 20,000 Gaussian primitives. Each contraction must contain no more than 30 Gaussians. The total number of contracted basis functions, or AOs, cannot exceed 8192. You may use up to 1050 effective fragments, of at most 5 types, containing no more than 2000 multipole/polarizability/other expansion points.

In practice, you will probably run out of CPU time or disk storage before you encounter any of these limitations. See Section 5 of this manual for information about changing any of these limits, or minimizing program memory use.

Except for these limits, the program is basically dimension limitation free. Memory allocations other than these limits are dynamic, from the storage requested by the input.
Restart Capability

The program checks for CPU time, and will stop if time is running short. Restart data are printed and punched out automatically, so the run can be restarted where it left off.

At present all SCF modules will place the current orbitals on the punch file if the maximum number of iterations is reached. These orbitals may be used in conjunction with the GUESS=MOREAD option to restart the iterations where they quit. Also, if the TIMLIM option is used to specify a time limit just slightly less than the job's batch time limit, GAMESS will halt if there is insufficient time to complete another full iteration, and the current orbitals will be punched.

When searching for equilibrium geometries or saddle points, if time runs short, or the maximum number of steps is exceeded, the updated hessian matrix is punched for restart. Optimization runs can also be restarted with the direct access file DICTNRY. See $STATPT for details.

Force constant matrix runs can be restarted from cards. See the $VIB group for details.

The two electron integrals may be reused. The Newton-Raphson formula tape for MCSCF runs can be saved and reused.

* * * *

The binary file restart options are rarely used, and so may not work well (or at all). Restarts which change the card input (adding a partially converged $VEC, or updating the coordinates in $DATA, etc.) are far more likely to be successful than restarts from the DAF file.