# Introduction to running Gaussian

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# Contents

Gaussian Input Files	1
General Input File Format:	1
Molecule Specifications	2
Z-Matrix	
Z-Matrix Examples	3
Important Keywords (Route section)	
Output Level	
Model Chemistry	
Basis Set	4
Job Type	4
Notes about Keywords:	
Coordinate Driving in Internal Coordinates:	
	5
Important Link0 commands	5
Alternative Way to Specify Memory and CPUs	
Running Gaussian at Compute Canada	6
	6
Small Gaussian Job	-
Big Gaussian Job	
Specifying Resources with environment vars	
- • •	
Running Gaussian in parallel	
	8
Estimating memory usage with frequem	8
Links	9
Software	9
Online Databases/services	9
'	

# Gaussian Input Files

# General Input File Format:

1	%MEM=30GB	{LinkO Commands}
2	%CHK=mycoolmolecule.chk	
3	# B3LYP/6-31G(d) Opt Freq MaxDisk=400GB	{Keyword lines}
4		{EMPTY LINE}

```
This is a very cool molecule
                                                                 {Title section}
                                                                 {EMPTY LINE}
   0 1
                                                                 {Charge, Multiplicity}
   Η
         0.000000000
                          0.000000000
                                            0.000000000
                                                                 {Molecule spec}
   0
         1.0480000000
                          0.000000000
                                            0.000000000
   C
         1.5064000000
                          1.3312900000
                                            0.000000000
10
   Η
         2.5540780000
                          1.3053060000
                                           -0.000000000
11
   Η
         1.1371360000
                          1.7931990000
                                            0.8652100000
12
   Η
         1.1371360000
                          1.7931990000
                                           -0.8652100000
13
                                                                 {EMPTY LINE}
14
                                                                 {optional sections}
15
```

**Important**: Don't forget to include the following empty lines:

- ullet before and after the title section
- after the molecule specifications

## Molecule Specifications

Molecules (elements and coordinates) can be specified in a number of different ways, using:

- Cartesian (XYZ) coordinates
- Z-Matrix coordinates
- a combination of XYZ- and Z-Matrix

Further, it is possible to add more information such as:

- Atom-type and partial charge (for Molecular Mechanics calculations)
- non-default Isotopes (spectra/thermo-chemistry from frequency calculations)

MolSpec reference at: http://gaussian.com/molspec/

#### **Z-Matrix**

Z-Matrices are an alternative way to specify molecule geometries, which can be more intuitive to use for a chemist. They specify a number of distances (e.g. bond lengths), angles and dihedral angles between the atoms.

- 1. Atom 1 is placed at the origin.
- 2. Atom 2 is placed at a distance *R1* of atom 1 along the Z-axis (hence the name Z-Matrix).
- 3. Atom 3 is placed at a distance R2 from atom 1, so such atoms 3, 2 & 1 are at an angle A1.
- 4. Atom 4 is placed at
  - a distance R3 from atom 3
  - spanning an angle A2 between atoms 4, 3, 2
  - and a dihedral angle D1 between atoms 4, 3, 2, 1
- 5. All further atoms are defined by a distance, angle and dihedral with respect to any previously defined atoms.

Indeed atom 3 can also be defined distance 3-1 and angle 3-1-2 and atom 4 via any permutation of the already defined atoms 1, 2 & 3.

Z-Matrix reference http://gaussian.com/zmat/

## **Z-Matrix Examples**

Z-Matrix of H2O2 with values within the coordinates (compact):

```
1 H
2 O 1 0.9
3 O 2 1.4 1 105.0
4 H 3 0.9 2 105.0 1 120.0
```

Z-Matrix of H2O2 using symbolic values:

```
H
2 0 1 R1
3 0 2 R2 1 A
4 H 3 R1 2 A 1 D
5 Variables:
6 R1 0.9
7 R2 1.4
8 A 105.0
9 D 120.0
```

## Important Keywords (Route section)

## **Output Level**

The keyword line starts with a #-symbol either on it's own or followed by a letter T, N or P. This controls the amount of information that printed to the Gaussian output file:

- #N: "Normal" amount of output (same as #)
- #T: "Terse" (reduced) output
- #P: Print additional output

## **Model Chemistry**

In most cases the Model chemistry is defined by a combination of "method" and "basis set". Examples of commonly used methods are:

- HF (Hartree Fock),
- B3LYP (a common DFT method) and
- MP2 (second order Møller-Plesset expansion).

Most methods can be prefixed with R (closed-shell restricted wavefunctions), U (unrestricted open-shell wavefunctions) or RO (restricted open-shell wavefunctions).

#### Links:

- Model Chemistry Reference: http://gaussian.com/capabilities/
- Hartree Fock theory in "Computational Chemistry" course material

#### Basis Set

A basis set defines a set of basis functions, which are used to calculate the molecular orbitals. Common basis sets are:

- STO-3G (minimal basis using "Slater Type Orbitals"),
- 6-31G (a Pople-type split-valence basis set).
- 6-31G(d) (same but with d-type polarization functions)
- cc-pVDZ (correlation-consisted polarized Double-zeta basis set)

For some methods however, no basis sets can be specified, e.g.: PM7 (semi-empirical "Parametric Model number 7") or G3 ("Gaussian-3" composite method).

#### Links:

- Basis Set on Wikipedia
- Basis Sets in "Computational Chemistry" course material

## Job Type

The job type defines what kind of calculation should be performed:

- SP Single Point just calculate the energy for the given coordinates
- OPT Geometry Optimization typically an energy minimization
- Opt=TS Transition State search
- Scanning Potential Energy Surfaces

### Notes about Keywords:

- Capitalization does not matter. OPT is the same as Opt or opt.
- keywords and options may be shortened to their shortest unique abbreviation. The keyword is actually Optimization, but as no other keyword begins with Opt, it can be abbreviated.
- Some keywords can take options (see documentation) e.g. Opt=Z-Matrix or or Opt=TS. These options can also be combined e.g. Opt=(TS,Z-Matrix).

List of Gaussian Keywords: https://gaussian.com/keywords/

## Coordinate Driving in Internal Coordinates:

```
#N HF/6-31G(d) Opt=Z-Matrix NoSymm
   H202 rotational potential 0.0 - 180.0, HF/6-31G(d) level internal coordinates.
   0 1
   H1
   02 1 r1
   03 2 r2 1 a1
   H4 3 r1 2 a1 1 d1
     Variables:
10
   r1=1.0
   r2=1.3
12
   a1=110.0
   d1=0.0 S 18 +10.0
14
15
```

## Coordinate Driving in Redundant Internal Coordinates:

```
Opt=ModRed
    \#N HF/6-31G(d)
2
   H2O2 rotational potential 0.0 - 180.0, HF/6-31G(d) level
   redundant internal coordinates.
   0 1
   H1
       1 r1
   02
   03
       2 r2 1 a1
   H4 3 r1 2 a1 1 d1
10
     Variables:
11
   r1=1.0
12
   r2=1.3
   a1=110.0
14
   d1=0.0
15
16
   1 2 3 4 S 18 +10.0
17
18
```

## Important Link0 commands

- %MEM=30GB request 30GB of memory
- %NProcShared=4 (or %NProc=4) request a job for 4 CPUs
- %CHK=/scratch/username/mycoolmolecule.chk set checkpoint filename

Gaussian Link0 commands: http://gaussian.com/link0/

## Alternative Way to Specify Memory and CPUs

Order of precedence: Link0 > command line > environment variable

Link0 (%-lines)	command line	environment variable	Note
%Mem=30GB	-m=30GB	export GAUSS_MDEF="30GB"	*
%NProc=4		export GAUSS_PDEF="4"	†
%CPU=0-31	-c="0-31"	export GAUSS_CDEF="0-31"	‡

See Equivalency-tab on http://gaussian.com/options/ or http://gaussian.com/link0/.

<sup>\*:</sup> Unit is indicated as MB or GB (not M or G as for SLURM).

<sup>†:</sup> The command line option -p does not work for Gaussian 16.c01.

<sup>‡:</sup> The Gaussian manual suggests using %CPU instead of %NProc to make sure that Gaussian processes are "pinned" to the processors, which can avoid cache misses. On Compute Canada clusters, processes are pinned automatically by the scheduler and at the time of job-submission it cannot be determined which specific CPU cores will be assigned to the job. If at all, the %CPU notation (or its counterparts) should only be used when requesting whole nodes.

# Running Gaussian at Compute Canada

### Gaussian License restrictions

Gaussian is only available at Graham and Cedar.

In order to use Gaussian, the PI (supervisor) and each user needs to accept Gaussian's terms of use by sending an email with the following content to support@computecanada.ca:

I am not a member of a research group developing software competitive to Gaussian.

I will not copy the Gaussian software, nor make it available to anyone else.

I will properly acknowledge Gaussian Inc. and Compute Canada in publications.

I will notify Compute Canada of any change in the above acknowledgement.

## Small Gaussian Job

### small\_gaussian\_job.sh:

### h2o2\_b3lyp.com:

```
#N HF/6-31G(d) Opt
   H2O2 HF/6-31G(d) level
   0 1
   H 0.000000
               0.992020
                            0.835282
6
   0.000000
                0.650000
                           -0.104410
   0.000000
                -0.650000
                           -0.104410
   H 0.000000
               -0.992020
                            0.835282
10
```

## Big Gaussian Job

## big\_gaussian\_job.sh:

```
#!/bin/bash
#SBATCH --mem-per-cpu=4000M  # memory-per-cpu
#SBATCH --cpus-per-task=8  # cpus as defined by %NProcShared
#SBATCH --time=03-00:00  # expect run time (DD-HH:MM)
module load gaussian/g16.c01
# use localscatch:
export GAUSS_SCRDIR=$SLURM_TMPDIR
g16 < big_molecule.com > big_molecule.log
```

### big\_molecule.com:

```
%NProcShared=8
%MEM=30GB
%CHK=/scratch/<USERNAME>/big_molecule.chk
#N MP2/6-311++G(3df,2p) OPT FREQ MaxDisk=400GB

My Big Molecule with a large basis set
...
```

## Specifying Resources with environment vars

### big\_gaussian\_job.sh:

#### big\_molecule.com:

```
%CHK=/scratch/<USERNAME>/big_molecule.chk
%N MP2/6-311++G(3df,2p) OPT FREQ MaxDisk=400GB

My Big Molecule with a large basis set
...
```

Note that the unit for --mem is G, while for GAUSS\_MDEF it is GB!

Running Gaussian in parallel

- Gaussian recommends allocating 4GB or more per CPU-core for calculations with 50+ atoms and/or 500+ basis functions. http://gaussian.com/relnotes/?tabid=3
- Many, but not all stages of Gaussian calculations can be carried out utilizing multiple CPU cores and to varying degrees. Hartree-Fock and DFT energies, gradients and frequencies and MP2 energies and gradients seem to be well supported but comprehensive documentation is lacking.
- Number of CPUs that can be efficiently used depends on:
  - Method
  - number of Atoms
  - number of Basis functions
  - version of Gaussian

## Remember:

Your Mileage May Vary.

## Comparison: Large Molecule $(C_20O_4H_28)$ with 16, 32 CPUs

```
%MEM=120GB
%NProcShared=16
#N B3LYP/6-311++G(3df,2p) Opt FREQ
$ seff 26948781
Job ID: 26948781
Cluster: graham
User/Group: stuekero/stuekero
State: COMPLETED (exit code 0)
Cores per node: 16
                                                      <===
CPU Utilized: 13-06:23:40
CPU Efficiency: 96.40% of 13-18:17:52 core-walltime
Job Wall-clock time: 20:38:37
Memory Utilized: 37.02 GB
Memory Efficiency: 29.61% of 125.00 GB
                                                      <=== !!!
$ seff 26948862
Job ID: 26948862
Cluster: graham
User/Group: stuekero/stuekero
State: COMPLETED (exit code 0)
Cores per node: 32
                                                      <===
CPU Utilized: 13-07:42:35
CPU Efficiency: 99.85% of 13-08:11:12 core-walltime
Job Wall-clock time: 10:00:21
Memory Utilized: 73.00 GB
Memory Efficiency: 58.40% of 125.00 GB
                                                      <===
```

## Estimating memory usage with frequem

Gaussian frequent utility: http://gaussian.com/frequent/

The frequent utility helps to estimate the required memory for a frequency calculation.

It uses the following syntax:

## Links

- Gaussian Manual: http://gaussian.com/man/
- Gaussian on CC Docs Wiki: https://docs.computecanada.ca/wiki/Gaussian
- Gaussian error messages: https://docs.computecanada.ca/wiki/Gaussian\_error\_messages
- "Computational Chemistry" course material: www.schulz.chemie.uni-rostock.de/lehre/computerchemie/table-of-contents/
- Book "Exploring Chemistry with Electronic Structure Methods": http://gaussian.com/expchem3/
- Computational Chemistry List (CCL): http://ccl.net/

### Software

- Molecule Editors:
  - Open Source:
    - \* Avogardro: https://avogadro.cc/
    - \* Gabedit: http://gabedit.sourceforge.net/
    - \* Molden: http://cheminf.cmbi.ru.nl/molden/
    - \* Jmol: http://www.jmol.org/
  - commercial:
    - \* GaussView: http://gaussian.com/gaussview6
- Tools:
  - OpenBabel: http://openbabel.org/

## Online Databases/services

- NIH:
  - PubChem: https://pubchem.ncbi.nlm.nih.gov/
  - NIH CACTUS https://cactus.nci.nih.gov/
  - Online SMILES Translator and Structure File Generator https://cactus.nci.nih.gov/translate/
  - Chemical Structure Lookup Service (CSLS) https://cactus.nci.nih.gov/cgi-bin/lookup/search
- ChemSpider: http://www.chemspider.com/
- ChEMBL: https://www.ebi.ac.uk/chembl/
- Jmol Demo:
  - https://chemapps.stolaf.edu/jmol/jsmol/simple1.htm
  - http://ostueker.github.io/simple\_jmol\_demo/jmol\_surfaces.html