

Linux commands

cd	Change directory
mkdir	Create directory
rmdir	Remove directory
rm	Remove file
cp	Copy file
ls	List of content of current directory

Exemple: `cp he.inp he3216.inp`

This will copy the file he.inp and rename the copy as he3216.inp

The studied atom/molecule

Multiplicity (n number of unpaired electron)	n+1(n number of unpaired electron)
Charge	+/-/0

Run gameSS

he.inp	name of the input script
he.out	name of the output script
rungms	run gameSS

Exemple : `rungms he> he.out`

Will run gameSS with the script he.inp, and give you the output in the file he.out

Output file

How to see if my calculation worked?

Look density converged in the output file

Where to see the final energy (HF) ?

Right after the message density converged the Hartree Fock energy calculation is provided
`FINAL RHF ENERGY I S=`

Where to see the final energy (MP2/CCSD) ?

Look for a message `E (MP2/ CCSD) =`

How to see the number of basis fonction?

Beginning of the output script there is a recap of the input script, and search for Nb of cartesian basis fonction=

⚠ RHF final energy is always provided even if you don't do HF calculation.

Input in \$BASIS

STO-nG	GBASIS=STO	NGA
n-21G	GBASIS=N21	NGA
n-31G	GBASIS=N31	NGA
n-311G	GBASIS =N311	NGA
cc-PVnZ	GBASIS=CCn	! I =1 i TRC

x shell NxFUNC=number of set x=p
 polari-
 sation
 function

Semi empirical methods
 GBASIS =MNDO

GBASIS=AM1

GBASIS=PM3

Exemple:

- Calculation with 6-31G* Basis set for He
`GBASIS=N31 NGAUSS=6 NPFUNC=1`
 - Calculation with 6-31G** Basis set for He
`GBASIS=N31 NGAUSS=6 NPFUNC=2`
- Use P because He is full on s, not on p



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Input in \$DATA

1. title/details
2. symmetry point group, usually C1
3. empty line, unless symmetry different than C1
4. Atomic coordinates
 - ▶ COORDS =CART

atome name / nuclear charge / X / Y / Z

```
H 1 0.0 0.0 0.0
C 6 1.1 0.0 0.0
```

▶ COORDS=ZMT

atom / l / distance / J / angle / K / torsion

```
H 1 1.1 2 125.0 3 180.0
```

▶ COORDS =ZMTMPC

atom / distance / l / angle / l / torsion / l / J / K

```
H 1.1 1 125.0 1 180.0 1 2 3
```

⚠ For COORDS=ZMT and COORDS =ZMTMPC, only put the required data !

Atom 1 : only name
 Atome 2 : name + distance
 Atome 3 : name + distance + angle
 Atome 4 and + : all data

Input in \$CONTRL

Method/Wave function choice	Close Shell System	SCFTYP=RHF default
	Open Shell System	SCFTYP= UHF
		SCFTYP= ROHF
		SCFTYP=GVB
		SCFTYP= MCSCF
Electron correlation method	Mollera-Plesseta MPn	MPLEVL=n
	DFT	DFTTYP= word
	coupled-cluster CC	CCTYP= word
	configuration interaction CI	CITYP= word
	valence bond VB	VBTP= word
	TDDFT	TDDFT= word

Input in \$CONTRL (cont)

Other	Charge <i>n</i> of molecule	ICHARG= <i>n</i>
	Multiplicity <i>n</i> of molecule	MULT= <i>n</i>
	Format of geometry in \$D	COORDS= word ATA

DFT method's words: VWN/BP 86/ BLYP . . .

CC method's words: CCD/CC SD/ CCS D (T) . . .

Words for coordinates: CART/Z MT/ ZMTM PC



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