

Remarks on Vconstr code interfaced with GAMESS-US

basic information, compilation and use

Last modified: May 13, 2013

1 General Information

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1 echo Files used on the master node master were:
2 ls -lF
```

2 dbrain.f requires the following data

- **hmatx** core hamiltonian matrix
- **tsmat** kinetic energy operator integrals
- **vnmat** electron-nuclear attraction operator integrals
- **vmopao** molecular orbitals in primitive ao basis
- **vmoao** molecular orbitals in mo basis
- **pmo** mo density matrix in mo basis
- **pnomo** ci density matrix in mo basis

3 dsfun.x input

Table 1: dsfun input parameters

name	type	comment
title	character(len=8)	
nmos	integer	number of molecular orbitals
occ	real(1:nmos)	occupation numbers
df	real	shift
nppr	integer	pprint, intermediate storage of Vxc at every iteration
scfdmp	real	damping
lrfun	logical	linear response
dvdmp	real	linear response procedure damping
lfield	logical	electric field
fzyx	real(1:3)	components of the electric field
nvpr	integer	vprint
lsym	logical	symmetrize the orbitals
lintsm	logical	determine symmetry using orbitals(if false h-matrix)
ismo	integer	mo section in the atmol dumpfile
isno	integer	no section in the atmol dumpfile
isao	integer	adapt section in the atmol dumpfile from which the Vmopao matrix is res

References