Remarks on Vconstr code interfaced with GAMESS-US

basic information, compilation and use

Last modified: May 13, 2013

1 General Information

 ${\scriptstyle 1}$ echo Files used on the master node master were: ${\scriptstyle 2}\; 1s\; \mbox{-1F}$

2 dbrain.f requires the following data

- hmatx core hamiltonian matrix
- tsmat kinetic energy operator integrals
- $\bullet\,$ vnmat electron-nuclear attraction operator integrals
- vmopao molecular orbitals in primitive ao basis
- $\bullet\,$ v
mo
ao moelcular 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 |
| осс | 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

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