DoNOF is a computational chemistry software program that stands for Donostia Natural Orbital Functional. The original code started on January 21, 2009 as PNOFID. It will run on essentially any machine with a FORTRAN 90 compiler for 64 bit processing.

DoNOF can perform computational chemistry calculations based on the Natural Orbital Functional Theory (NOFT), including PNOF5, PNOF6 and PNOF7. Correlation corrections after PNOF calculations can be estimated by second order perturbation theories. The total spin is conserved, not just the spin projection.

The solution is established optimizing the energy functional with respect to the occupation numbers (ONs) and to the natural orbitals (NOs), separately. The constrained nonlinear programming problem for the ONs is treated under pairing restrictions as an unconstrained minization, while the orbital optimization is carried out by a self-consistent procedure which yields the NOs automatically orthogonal. To achieve convergence, the direct inversion of the iterative subspace (DIIS) extrapolation technique is used, and a variable scale factor balances the symmetric matrix subject to the iterative diagonalizations.

The &INPRUN and &NOFINP namelists specify the input and output, and the fundamental job options. These options are controlled by the following keywords:

C-----

```
C
                  --- NAMELIST VARIABLES ---
(-----
C
C RUNTYP
                  specifies the run calculation
C
     = ENERGY 1) single-point energy calculation (Default)
      = GRAD 2) energy + gradients with respect to nuclear coord.
= OPTGEO 3) optimize the molecular geometry
C
      = GRAD
C
C
C MULT
                 Multiplicity of the electronic state
               singlet (Default)
C
       = 2,3,... doublet, triplet, and so on
C
C
C ICHARG
                 Molecular charge
C
      = 0
                Neutral Molecule (Default)
C
C IECP
                Effective Core Potentials (to be implemented)
              (Default) All electron calculation
Read ECP potentials in the $ECP group
       = 0
C
C
       = 1
C
               Electrostatic moments calculation skip calculation
C IEMOM
C
       = 0
                calculate monopole and dipole (Default)
C
       1
              also calculate quadrupole moments also calculate octopole moments
C
        2
C
C
C UNITS
                Distance units (any angles must be in degrees)
      = ANGS Angstroms (Default)
= BOHR Bohr atomic units
C
C
C
C EVEC
                  An array of the three x,y,z components of
                  the applied electric field, in a.u.
C
C
                  (1 a.u. = 1 Hartree/e*bohr = 5.1422082(15)d+11 V/m)
                  (Default)
C
       = 0.000
C
C DONTW
                  Do not write 2e- integrals on the disk (Unit=1)
C
       = T
                  (Default)
C
C-----
     NAMELIST/INPRUN/RUNTYP, MULT, ICHARG, IECP, IEMOM, UNITS, EVEC, DONTW
C
    Initial Values for the namelist variables
C-----
     RUNTYP = ENERGY
     MULT = 1
     ICHARG = 0
     IECP = 0
```

IEMOM = 1
UNITS = ANGS
EVEC = 0.0D0 ! EVEC(1,2,3)=0
DONTW = .TRUE.

C .	NAMEL	IST VARIABLES
C MAXIT C	= 1000	Maximum number of OCC-SCF iterations (DEFAULT)
C		
C Type of Calculation		
C		
C CICOEF C	= 0	Coefficient Optimization Optimize Energy only by the occupations
C	= 1	use the Iter. Diag. method (DEFAULT)
C	= 2	Optimize Energy only by the orbitals
C C C	= 3	Optimize Energy by all occupations and only core-fragment orbitals, the rest of fragment orbitals remain frozen
CIEINI C C	= 0	Calculate only the initial energy (DEFAULT)
C NO1 C	= -1	Max. index of NOs with Occupation = 1 Consider Core NOs (DEFAULT)
C C C	= 0 = Value	
C		
C Hartree-Fock		
C		
C HFID C C	= T	Use the Iterative Diagonalization Method to generate the HF Orbitals (DEFAULT)
C CNTHRESHEID	·	Convergence of the TOTAL ENERGY
C C C	= 8	THRESHEID=10.0**(-NTHRESHEID) (DEFAULT)
C MAXITID C C	= 30	Maximum number of external iterations (DEFAULT)
C		
C		
CIPNOF	= 5	Type of Natural Orbital Functional (NOF) PNOF5
C C C	= 6 = 7	PNOF6 PNOF7 (DEFAULT)
C Ista C C	= 0 = 1	Use Static version of PNOF7 PNOF7 (DEFAULT) PNOF7s
C C HighSpin C C	= F = T	Spin-uncompensated calculation type (DEFAULT) Multiplet state (Ms=0) High-spin uncompensated state (Ms=S)
C C NCWO	= 1	<pre>Number of coupled weakly occupied MOs per strongly occupied = Nc -&gt; PNOFi(Nc) NCWO = 1 (DEFAULT)</pre>
C C C	= 2,3, =-1	.  NCWO = NVIR/NDOC  NVIR: Number of HF virtual MOs (OCC=0)  NDOC: Number of strongly occupied MOs

NDOC: Number of strongly occupied MOs

C		
C	 in NOF c	
C		
C NTHRESHL		Convergence of the Lagrange Multipliers
C		THRESHL=10.0**(-NTHRESHL)
C	= 4	(DEFAULT)
C NTHRESHE		Convergence of the total energy
C		THRESHE=10.0**(-NTHRESHE)
C C	= 8	(DEFAULT)
CNTHRESHEC		Convergence of the total energy (ORBOPT)
C		THRESHEC=10.0**(-NTHRESHEC)
C	= 10	(DEFAULT)
C NTHRESHEN		Convergence of the total energy (OCCOPT)
C		THRESHEN=10.0**(-NTHRESHEN)
C C	= 10	(DEFAULT)
C		
-		ization Program (ID Method)
C		
C NOPTORB		Number of the optimized orbitals
C	= NBF	(DEFAULT)
C MAXLOOP		Maximum Iteration Number for the SCF-
C		iteration cycle in each ITCALLs
C	= 30	(DEFAULT)
C The straightforw	ard itera	tive scheme fails to converge very
C often due to the	values o	f some off-diagonal elements Fki. The
		ly small and of the same order of
		tor scales Fki. We establish an upper at when the absolute value of the
C matrix element F	ki is gre	ater than B, it is scaled by a factor
C Cki (F'ki = Cki*	Fki ), as	to satisfy ABS(Fki) <= B.
C SCALING		A variable factor scales Fki
C	= T	(DEFAULT)
C NZEROS		B = 10.0**(1-NZEROS).
C		Initial number of ZEROS in Fij. The
C		scaling factor varies until the number
C		of ZEROS (.000##) is equal for all elements Fij.
C	= 0	B = 10.0 (DEFAULT)
( N7EDOS m		B = 10.0**(1-NZEROSm)
C NZEROSm		Maximum number of zeros in Fij.
C	= 4	B = 10.0 (DEFAULT)
C NZEROSr		B = 10.0**(1-NZEROSr)
C NZERO31		Number of zeros in Fij to restart
C		automatically the calculation.
C	= 0	B = 10.0 (DEFAULT)
C ITZITER		Number of Iterations for constant scaling
C	= 10	(DEFAULT)
C DIIS		Direct Inversion in the Iterative
C		Subspace in the orbital optimization if
C	_	DUMEL < THDIIS every NDIIS loops
C	= T	(DEFAULT)
C NTHDIIS		Energy threshold to begin DIIS
C	= 3	THDIIS = 10.0**(-NTHDIIS) (DEFAULT)
C NDIIS		Number of considered loops to interpolate
C		the generalized Fock matrix in the DIIS
С	= 5	(DEFAULT)

C C PERDIIS C C C	= T = F	Periodic DIIS Apply DIIS every NDIIS (DEFAULT) DIIS is always applied after NDIIS		
C C Options for pertubat C	ive calcu			
C C OIMP2 C C	= F	NOF - Orbital Invariant MP2 (DEFAULT)		
C N01PT2 C C C C	= -1 = 0 = Value	Frozen MOs in perturbative calculations Maximum index of NOs with Occupation = 1 = NO1 (DEFAULT) All NOs are considered User specifies how many NOs are frozen		
•	= F	SC2-MCPT perturbation theory is used to correct the PNOF5 Energy. 2 outputs: PNOF5-SC2-MCPT and PNOF5-PT2 (DEFAULT)		
C	= 0	Number of excluded coupled orbitals in the PNOF5-PT2 calculation All NOs are included (DEFAULT)		
C C				
	Resta	rt Options		
C CRESTART C C	= F = T	Restart from GCF file (DEFAULT=F) INPUTGAMMA=0,INPUTC=0,INPUTFMIUG=0 INPUTGAMMA=1,INPUTC=1,INPUTFMIUG=1		
C CINPUTGAMMA C C	= 0 = 1	Guess for GAMMA variables (ONs) Close Fermi-Dirac Distribution (DEFAULT) Input from file GCF		
C C C C	= 0 = 1	Guess for Coefficient matrix (NOs) Use HCORE or HF Eigenvectors (DEFAULT) Input from file GCF		
C CINPUTFMIUG C C	= 0 = 1	Guess for Diagonal elements (FMIUG0) Use single diag. of Lagragian (DEFAULT) Input from file GCF		
C CINPUTCXYZ C C	= 0 = 1	Nuclear Coordinates (CXYZ) Input from input file (*.inp) (DEFAULT) Input from file GCF		
C C				
C CNPRINT C C C C	= 0 = 1 = 2	Output Option (DEFAULT VALUE: 0) Short Printing (Occ,Emom,Energies) Output at initial and final iterations including MOs,Pop,APSG,Lag,IPs,DMs,CJK Output everything in each iteration		
C CIWRITEC C C	= 0 = 1	Output Option for the Coefficient matrix No output (DEFAULT) Output the Coefficient Matrix (NOs)		
C CIMULPOP C C	= 0 = 1	Mulliken Population Analysis Not do it (DEFAULT) Do it		
C C PRINTLAG C	= F	Output Option for Lagrange Multipliers No Output (DEFAULT)		

C				
C DIAGLAG	Diagonalize Lagrange Multipliers			
C	Print new 1e- Energies, Canonical MOs, and new diagonal elements of the 1RDM			
C = F	Not do it (DEFAULT)			
C C IEKT	IPs by Ext. Koopmans' Theorem (EKT)			
C = 0	Not calculate the IPs			
C = 1 C	Calculate ionization potentials (IPs)			
C				
C C IAIMPAC	Write information into WFN file (UNIT 7)			
С	for THE AIMPAC PROGRAM			
C = 0 C = 1	DO NOT DO Write INTO WFN file (DEFAULT)			
C	WITCE INTO WIN TICE (DEFNOET)			
C NOUTRDM C = 0	Print OPTION for ATOMIC RDMs NO Output (DEFAULT)			
C = 1	Print ATOMIC RDMs IN 1DM and 2DM files			
C	TUDECUDM-10 011/ NTUDECUDM)			
C NTHRESHDM C = 6	THRESHDM=10.0**(-NTHRESHDM) (DEFAULT)			
C				
C NSQT C = 1	Print OPTION for 2DM file UNforMATTED (DEFAULT)			
C = 0	forMATTED (SEE SUBROUTINE OUTPUTRDMrc)			
C C NOUTCJK	Print OPTION for CJ12 and CK12			
C = 0	NO Output (DEFAULT)			
C = 1	Print CJ12 and CK12 in file 'CJK'			
CNTHRESHCJK	THRESHCJK=10.0**(-NTHRESHCJK)			
C = 6	(DEFAULT)			
C C NOUTTijab	Print OPTION for Tijab			
C = 0	NO Output (DEFAULT)			
C = 1 C	Print Tijab in file 'Tijab'			
C NTHRESHTijab	THRESHTijab=10.0**(-NTHRESHTijab)			
C = 6 C	(DEFAULT)			
C APSG	Open an APSG file for printing the			
C C	coefficient matrix (\$VEC-\$END) and the expansion coefficients of the APSG			
C	generating wavefunction.			
C = F C	Output (DEFAULT)			
C NTHAPSG	Threshold for APSG expansion coefficient			
C = 10	THAPSG = 10.0**(-NTHAPSG) (DEFAULT)			
C = 10 C	(DEFAULT)			
(				
C Optional Options C				
C	0.46			
C ORTHO C = F	Orthogonalize the initial orbitals No			
C = T	Yes (DEFAULT)			
C C CHKORTHO	Check the Orthonormality of the MOs			
C = F	No (DEFAULT)			
C = T	Yes			
C				
C Options related to Frozen coordinates in gradient computation C				
C C FROZEN	Is there any fixed coordinate			
C = F	(DEFAULT)			
C C IFROZEN	By pairs, what coordinate of which atom,			
C	e.g. 2,5,1,1 means "y" coordinate of			

```
C
                               MAXIMUM of frozen coordinates = 10
C
                      = 0
                               (DEFAULT)
C
C-
                         C
C..... ICGMETHOD
                               Define the Conjugate Gradient Method in
C
                               OCCOPTr, CALTijabIsym and OPTIMIZE
C
                      = 1
                               (DEFAULT)
C
                               SUMSL: CGOCUPSUMSLr, OPTSUMSL
C
                               SparseSymLinearSystem_CG
C
                      = 2
                               Use NAG subroutines:
C
                               E04DGF: OPTCGNAG, CGOCUPNAGr
C
                               F11JEF: SparseSymLinearSystem_NAG
C
                               LBFGS: OPTLBFGS, LBFGSOCUPr
C
C-----
     NAMELIST/NOFINP/MAXIT, ICOEF, IEINI, NO1, HFID, NTHRESHEID, MAXITID,
    &
                     IPNOF, Ista, HighSpin, NCWO, NTHRESHL, NTHRESHE,
    &
                     NTHRESHEC, NTHRESHEN, NOPTORB, MAXLOOP, SCALING,
    &
                     NZEROS, NZEROSm, NZEROSr, ITZITER, DIIS, NTHDIIS,
    &
                     NDIIS, PERDIIS, SC2MCPT, NO1PT2, NEX, OIMP2,
                     RESTART, INPUTGAMMA, INPUTC, INPUTFMIUG, INPUTCXYZ,
                     NPRINT, IWRITEC, IMULPOP, APSG, NTHAPSG, PRINTLAG,
    &
                     DIAGLAG, IAIMPAC, IEKT, NOUTRDM, NTHRESHDM, NSQT,
    &
                     NOUTCJK, NTHRESHCJK, NOUTTijab, NTHRESHTijab,
                     ORTHO, CHKORTHO, FROZEN, IFROZEN, ICGMETHOD
    Preset values to namelist variables
     MAXIT=1000
     Type of Calculation
     ICOEF=1
     IEINI=0
     N01 = -1
C
     Hartree-Fock
     HFID=.TRUE.
     NTHRESHEID=8
     MAXITID=30
C
     PNOF Selection
     IPNOF=7
     Ista=0
                                                 ! PN0F7n
     HighSpin=.FALSE.
                                                 ! Multiplet
     NCW0=1
                                                 ! Perfect Pairing
C
     Convergence Criteria in NOF calculation
     NTHRESHL=4
     NTHRESHE=8
     NTHRESHEC=10
     NTHRESHEN=10
     Options for the Orbital Optimization Program (ID Method)
C
     NOPTORB=-1
                                                ! NOPTORB=NBF
     MAXLOOP=30
     SCALING=.TRUE.
     NZEROS=0
     NZEROSr=0
     NZEROSm=NTHRESHL
                                                ! NTHRESHL
     ITZITER=10
     DIIS=.TRUE.
     NTHDIIS=3
     NDIIS=5
     PERDIIS=.TRUE.
C
     Options for pertubative calculations
     SC2MCPT=.FALSE.
     NEX=0
     N01PT2=-1
     OIMP2=.FALSE.
```

atom 5 and "x" coor of atom 1 to freeze.

```
Input Options for Gamma (Occ), C and Diagonal F
C
      RESTART=.FALSE.
      INPUTGAMMA=0
      INPUTC=0
      INPUTFMIUG=0
      INPUTCXYZ=0
      Output Options
C
      NPRINT=0
      for NPRINT>0
C
      IWRITEC=0
      IMULPOP=0
      APSG=.FALSE.
      NTHAPSG=10
      PRINTLAG=.FALSE.
      DIAGLAG=.FALSE. ! Only in the final Output
      IAIMPAC=1
      IEKT=0
                       ! Only in the final Output
C
      NOUTRDM=0
      NTHRESHDM=6
      NSQT=1
C
      NOUTCJK=0
      NTHRESHCJK=6
      NOUTTijab=0
      NTHRESHTijab=6
C
      Optional Options
      ORTHO=.TRUE.
      CHKORTHO=.FALSE.
C
      Frozen coordinates
      FROZEN=.FALSE.
      IFROZEN=0
C
      Options for the Conjugate Gradient Method
```

C-----

ICGMETHOD=1