FUKUI v231003

A Program to Compute Fukui Indices, Dipole Moments and Atomic Overlap Matrices

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1 Fukui: scheme and software

This document is a short guide to use Fukui, a program developed as a multipurpose tool to perform:

- chemical reactivity indices called "Fukui indices",
- σ/π separation of atomic populations (reading an input file with the molecular orbital symmetry),
- the calculation of dipole moment components,
- the calculation of atomic overlap matrices as a previous step to compute the N-electron delocalization indices (employed for aromaticity studies, for example) and condensed 2-center Fukui indices, used for studies of reactivity and resonance effects.

The program reads grid points from several sources and, therefore, it is independent of the type of real-space atomic electron density partitioning employed.

1.1 Scheme of the program [Go To Table Of Contents]

Fukui is a tool designed to perform several tasks in a parallelized and efficient way using a grid of points:

- Compute the total population from the aforementioned grid.
- Real space molecular partitioning schemes, such as QTAIM¹ and Hirshfeld-based^{2–4} methodology, can be employed for the calculation of condensed Fukui indices using the fragment of molecular response (FMR) approach.^{5,6}
- Real space atomic overlap matrix (AOM) can be computed within the grid domains as a previous step to obtain bond orders and N-electron delocalization indices through a grid of points following Mandado's implementation.^{7,8}
- σ/π molecular orbital (MO) contributions can be separated from the total population.
- Compute the dipole moment components from the grid. If an atom number is specified as third argument, the dipole moment is divided into two components: the interatomic charge transfer (μ_c) and the polarizations of the individual atomic distributions $(\mu_a)^{1}$

1.2 Program details [Go To Table Of Contents]

- Programming language: Fortran 2008
- Operating systems: Any with Fortran (compilers), CMake and, optionally, OpenMP (for parallelization support).
- List of source files: fukui.f90 goon.f90 help.f90 onlygrid.f90 modules/computation.f90 modules/grid.f90 modules/main.f90 modules/output.f90 modules/pi.f90 modules/wfn.f90 openm-p/init_par.f90
- List of CMake files: CMakeLists.txt modules/CMakeLists.txt openmp/CMakeLists.txt
- List of utilities: bstk_astk.f90 cube2stk.f90 fukui.sh mwfn2piorbs.sh som2sg.sh

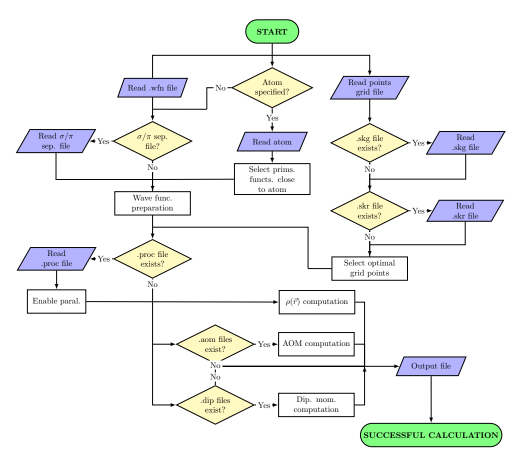


Figure 1.1: Simplified flowchart of the Fukui 231003 program execution. Terminal blocks are represented by a green stadium shape, while the input/output and decision blocks correspond to blue rhomboid and yellow rhombus shapes, respectively. Last, the rectangle is used to stand for process blocks in the program. The acronyms of atomic overlap matrix (AOM), "Dip. mom." (dipole moment) and the symbol $\rho(\vec{r})$ (electron density) are also employed in this chart.

1.3 Units [Go To Table Of Contents]

If not stated otherwise, the Fukui program outputs its results in atomic units (electrons) for populations.

1.4 Structure of the program [Go To Table Of Contents]

The structure and steps of the program are summarized in Figure 1.1.

1.5 Terms of use, dependencies and license [Go To Table Of Contents]

FUKUI is free software under GPLv3 license. We refer to the GitHub web page for more details about the license:

https://github.com/nom05/fukui

This manual is open source under a MIT license. We refer to the GitHub web page for more details about the license https://github.com/nom05/fukui_manual

1.6 How to cite Fukui 231003 [Go To Table Of Contents]

• N. Otero, M. Mandado and R. Mosquera, Fukui 231003 (2023), Universidade de Vigo.

2 Program installation

NOTE: The preferred platform or, more specifically, the platform the developers use is GNU/Linux. MS Windows is an untested alternative. Through WSL or WSL2 the compilation would be equivalent to the following instructions. It is beyond the scope of this manual to explain how to activate these options in MS Windows.

The program source code will be available in https://github.com/nom05/fukui. To install the program, follow the instructions:

1. Clone the repository (if not previously downloaded/included in a physical medium):

```
\triangleright git clone github.com/nom05/fukui
```

2. Enter in the directory:

```
⊳ cd fukui
```

3. Verify your current CMake version* is equal or greater than 2.8.12.

```
cmake --version

cmake version 3.22.3

CMake suite maintained and supported by Kitware (kitware.com/cmake).
```

4. Choose the Fortran compiler. By default, GNU Fortran (gfortran) will be employed. To use an alternative, edit the file called "CMakeLists.txt", in the parent directory of the source code, with your favorite editor (nano, vi, vim, Visual Studio Code, etc.). Comment (adding the symbol '#') or uncomment (remove '#') with the purpose of disabling or enabling, respectively, your personal options:

```
| The content of the content o
```

5. Create a new directory called, for example, "build" and enter inside:

```
⊳ mkdir build; cd build
```

6. Now, create the compilation environment:

^{*}From Wikipedia: In software development, CMake is cross-platform free and open-source software for build automation, testing, packaging and installation of software by using a compiler-independent method. CMake is not a build system but rather it generates another system's build files. It supports directory hierarchies and applications that depend on multiple libraries. It is used in conjunction with native build environments such as Make, Qt Creator, Ninja, Android Studio, Apple's Xcode, and Microsoft Visual Studio. It has minimal dependencies, requiring only a C++ compiler on its own build system.

```
⊳ cmake ..
                 The C compiler identification is GNU 11.2.0
The CXX compiler identification is GNU 11.2.0
Detecting C compiler ABI info
Detecting C compiler ABI info - done
  4
                Detecting C compiler ABI info - done

Check for working C compiler: /usr/bin/cc - skipped

Detecting C compile features

Detecting C compile features - done

Detecting CXX compiler ABI info

Detecting CXX compiler ABI info - done

Check for working CXX compiler: /usr/bin/c++ - skipped
  6
7
8
  9
10
                  Detecting CXX compile features
11
12
                  Detecting CXX compile features - done
                 The Fortran compiler identification is GNU 11.2.0 Detecting Fortran compiler ABI info Detecting Fortran compiler ABI info - done
13
14
15
                 Check for working Fortran compiler: /usr/bin/gfortran - skipped Found OpenMP_C: -fopenmp (found version "4.5")
Found OpenMP_CXX: -fopenmp (found version "4.5")
Found OpenMP_Fortran: -fopenmp (found version "4.5")
Found OpenMP: TRUE (found version "4.5")
16
18
19
20
         OPENMP FOUND
-- Configuring done
21
23
                  Generating done
                  Build files have been written to: [...]/fukui/src/build
```

7. And finally compile the code:

```
⊳ make
             Scanning dependencies of target fukui.x
[ 9%] Building Fortran object CMakeFiles/fukui.x.dir/modules/main.f90.o
/home/nicux/Documentos/research/code/fortran/fukui/src/modules/main.f90:206:45:
   3
   4
                                                 do i = 1,imos
   5
   \frac{6}{7}
             206 |
                                                                                       write(charintd,'(\overline{17})') moc(\underline{i-1})
   9
             Warning: Array reference at (1) out of bounds (0 < 1) in loop beginning at (2) [-Wdo-subscript] [ 18%] Building Fortran object CMakeFiles/fukui.x.dir/modules/grid.f90.o [ 27%] Building Fortran object CMakeFiles/fukui.x.dir/modules/wfn.f90.o
 10
 11
 12
                 2/k] Building Fortran object CMakeFiles/fukui.x.dir/modules/computation.f90.o
45%] Building Fortran object CMakeFiles/fukui.x.dir/modules/computation.f90.o
45%] Building Fortran object CMakeFiles/fukui.x.dir/modules/pi.f90.o
54%] Building Fortran object CMakeFiles/fukui.x.dir/modules/output.f90.o
63%] Building Fortran object CMakeFiles/fukui.x.dir/fukui.f90.o
72%] Building Fortran object CMakeFiles/fukui.x.dir/help.f90.o
81%] Building Fortran object CMakeFiles/fukui.x.dir/goon.f90.o
90%] Building Fortran object CMakeFiles/fukui.x.dir/openmp/init_par.f90.o
 13
 14
 15
 16
 17
 19
             [100%] Linking Fortran executable fukui.x
[100%] Built target fukui.x
20
```

8. You will find the executable in the current compilation directory:

```
□ Ls -tr

CMakeCache.txt openmp modules Makefile cmake_install.cmake main.mod grid.mod wfn.mod computation.mod pi.mod output.mod fukui.x CMakeFiles
```

fukui.x

TIP: To compile faster use several processor threads (increase the number of threads, #threads) through the option "-j#threads". For example, "make -j3" will set three (3) threads.

3 How to run Fukui

FUKUI has a special way to specify the options of the calculation. Without arguments, the executable prints a short help:

```
⊳ ./fukui.x
            PROGRAM FUKUI 231003
            ** WFN file is not included as argument **
  3
  4
5
            Type: ./fukui.x wfnfile stk_file/cube_file [# atom]
  6
                    Extension for stk and wfn files are optional, extension for cube file is mandatory.
                   Extension for stk and wfn files are optional, extension for cube file is m stk_file can be either a Fortran binary file or an ordinary text file. They have to include, following this order: x, y, z, Gauss quadrature weight and electron density values.

If the ordinary text format is considered, then the format to read must be specified, the label free can be used to use free format specification.

[# atom] is optional and enables gauss, func. skipping.

If wfnfile.skd exists, it reads cut-off1 inside.
  8
10
11
12
13
                   Sigma/pi separation: create .sg or .som (if both files exist, the first file sought is the .sg file) Only pi MOs: .sg/.som + .pi (command> touch wfnfile.pi).

Skip points with gauss. quad. weight < cut-off2: touch wfnfile.skg

Skip points with ref. dens < cut-off3: touch wfnfile.skr.

cut-off2 and cut-off3 can be changed by editing the values in the
14
15
16
17
18
19
                     previously created files.
                   Enable parallelization: Max ->touch wfnfile.proc

Set # procs->edit wfnfile.proc

Atomic Overlap Matrix (AOM) calculation: touch wfnfile.aom

Atomic Dipole moment calculation: touch wfnfile.dip

Specifying the number of atom [# atom] a charge transfer and intrinsic contributions are separated.
20
21
22
23
24
                    Specifying the number of atom [# atom] a charge transfer and intrinsic contributions are separated. Including the text OnlyGrid inside, the program will not compute the density, only grid will be used Change max dim. of array allocation: edit wfnfile.mxal Include in this file (all lines are mandatory):

o Line 1: Max # atoms.

o Line 2: Max # mol. orbs.

o Line 3: Max # points.

o Line 4: Max # prim. funcs.
25
26
27
28
29
30
```

As you can observe in the previous execution output without arguments, the program needs a file containing wave function information (see Section A.1 and Section A.2 in the Appendix for instructions on how to create it and convert among wave function files, respectively) and another file including the grid points.

3.1 File including grid of points [Go To Table Of Contents]

FUKUI supports two types of grid points files:

• Cube files, following the file format used by Gaussian utilities reference (https://gaussian.com/cubegen/). This grid is based on points set using a step size in the three directions of space, forming a cube (strictly speaking, a parallelepiped) containing the molecule. These cube files are built from the following scheme:

Line 1 and 2	Computational details and a comment line.		
Line 3	Number of atoms (NA), the grid origin in Cartesian coordinates and the number of values per point. In the case of electron density, the value is equal to 1.		
Line 4	Number of steps in the slowest running direction and the vector whose modulus corresponds to the distance between two points in this direction.		
Line 5	Number of steps in the intermediate running direction and the vector whose modulus corresponds to the distance between two points in this direction.		

Line 6

Number of steps in the fastest running direction and the vector whose modulus corresponds to the distance between two points in this direction.

Line 7 and the NA following Atomic number, charge, and coordinates of the atoms.

Line 7+NA+1 until the end Values of the electron density at each point in the grid. The total number of values is obtained as the product of the number of the steps in the three directions.

```
Cube file structure:
     tetracene B3LYP/6-31G** Density
2
     Electron density from Total SCF Density
                        -15.750103
3
       30
             -6.512752
                                      -9.176246
4
5
       53
             0.248108
                          0.000000
                                       0.000000
             0.000000
                          0.248108
      128
                                       0.00000
6
             0.000000
                                       0.248108
       75
                          0.000000
7
             6.000000
                          0.000000
                                       9.237351
                                                    1.351534
8
        6
             6.000000
                          0.00000
                                       7.012694
                                                    2.663494
       .]
10
              1,000000
                          0.000000
                                     -11.027453
                                                   -2.355229
11
      1.41965E-18 4.15287E-18 1.16752E-17
                                               3.15458E-17
                                                             8.19189E-17
                                                                          2.04457E-16
```

- A grid of points obtained by means of a gaussian quadrature, containing x, y, z, numerical integration weights and electron density values, strictly in this order. The program is able to
 - Fortran binary files, saved in unformatted files. They are the fastest way to read the set of points.
 - Text files with a Fortran specific format. The format to read must be specified. The label "free" can be used for Fortran's free format specification. In contrast, the latter is the slowest way to read a file in Fortran.

```
Example 1: text files including a set of points with a specific Fortran
   format:
   (3(F16.8),2(E18.8))
2
         0.0000000
                         2.62795893
                                         0.00031152
                                                       0.12858975E-09
                                                                         0.11765072E+03
3
         0.0000000
                         2.62795893
                                        -0.00031152
                                                       0.12858975E-09
                                                                         0.11765072E+03
4
         0.00000000
                         2.62827045
                                         0.00000000
                                                       0.12858975E-09
                                                                         0.11765075E+03
5
         0.00000000
                         2.62764741
                                         0.00000000
                                                       0.12858974E-09
                                                                         0.11765069E+03
6
         0.00031152
                         2.62795893
                                         0.0000000
                                                       0.12858975E-09
                                                                         0.11765072E+03
        -0.00031152
                         2.62795893
                                         0.0000000
                                                       0.12858975E-09
                                                                         0.11765072E+03
```

```
Example 2: text files including a set of points with free Fortran format:
   Free
                                                        0.12858975E-09
2
         0.0000000
                          2.62795893
                                          0.00031152
                                                                          0.11765072E+03
3
         0.00000000
                          2.62795893
                                         -0.00031152
                                                        0.12858975E-09
                                                                          0.11765072E+03
4
5
         0.00000000
                          2,62827045
                                          0.00000000
                                                        0.12858975E-09
                                                                          0.11765075E+03
         0.00000000
                          2.62764741
                                          0.0000000
                                                        0.12858974E-09
                                                                          0.11765069E+03
6
         0.00031152
                          2.62795893
                                          0.0000000
                                                        0.12858975E-09
                                                                          0.11765072E+03
         -0.00031152
                          2.62795893
                                          0.0000000
                                                        0.12858975E-09
                                                                          0.11765072E+03
```

3.2 How to perform a calculation [Go To Table Of Contents]

The mandatory files needed for the FUKUI program are the aforementioned wave function and grid files. For example, if the files are called "wavefunction.wfn" and "grid_file_name.stk", respectively, the way to run the calculation is:

```
⊳ executable path/fukui.x wavefunction.wfn grid file name.stk
```

The "executable path" is the directory containing the executable, for example "./" if you are in the current compilation directory.

The extensions are optional with the exception of the cube grid file.

3.3 Output file name [Go To Table Of Contents]

In the case the user had the following file names: wavefunction.wfn, grid_file_name.stk, Fukui will use the following output file name: "wavefunction grid file name.fuk".

3.4 Optional: specifying the corresponding atom where the grid points is centered on [Go To Table Of Contents]

In the case the set of points is centered on an atom, it is recommended to activate the "primitive gaussian functions skipping". This option avoids to include primitive functions far from the specified atom. The default distance is 18 au (a very safe default, according to our tests) but it can be modified creating a file "file.skd" employing the wave function file name and replacing the extension by "skd". There, a new value can be set. For example:

ho echo 4.000 > wavefunction.skd

As can be seen in Section 3.5, the specification of an atom opens the possibility to separate the dipole moment into two terms: charge transfer and the intrinsic atomic dipole moment.

3.5 Other options [Go To Table Of Contents]

In general, the Fukui program is employed massively in a large set of atoms. Instead of creating a unique input and performing one parallelized calculation, each grid can be processed in an independent way and, therefore, the user can submit several parallelized calculations in different nodes of a high-performance computing cluster, obtaining faster results. In this way, in order to facilitate the common options for the different files, the program reads or detects the existence of files to activate its options.

Increasing the maximum default employed resources [Go To Table Of Contents]

In the opinion of the developers, it is recommended to define consciously the maximum array dimensions instead of allocating the necessary memory without human control. The user must know if the available computational resources are enough in the case of (very) large systems instead of observing how the system kernel stops the program execution or the computer crashes. For this reason, the Fukui program includes these default cut-offs, modifiable through a configuration file:

# atoms	# MOs	# points	# primitives
50	250	1000000	1000

To increase these modest cut-offs (50 atoms are not a lot), the user will create a file including the following lines: Line 1: Max number of atoms. Line 2: Max number of molecular orbitals. Line 3: Max number of points. Line 4: Max number of primitive functions.

Enabling parallelization [Go To Table Of Contents]

FUKUI was developed with OpenMP parallelization (a calculation cannot be shared with other machines). To activate the full parallelization, employing all the threads of the computer, we only need to create an empty file with the wave function file name and replacing the current extension by the text "proc":

b touch wavefunction.proc

In the case a specific number of threads is required, the user can set it in the aforementioned file:

\triangleright echo 8 > wavefunction.proc

Modifying the distance in the primitive gaussian functions skipping [Go To Table Of Contents] See Section 3.4.

Skipping points with negligible Gauss quadrature weights [Go To Table Of Contents]

If the grid file was not processed previously removing negligible Gauss quadrature weights, the user can set the option to skip them. Typing the wave function file name and replacing the extension by "skg", this option will be enabled:

b touch grid file name.skg

The default value is 1.0×10^{-10} , a very reasonable cut-off according to our tests, but it is modifiable adding a new value in the file:

```
⊳ echo 1.e-9 > grid file name.skg
```

The program will print on screen and in the output file the number of points skipped and the total electron density with and without these negligible points.

Skipping points with negligible reference electron densities [Go To Table Of Contents]

If reference electron densities of the grid file were not previously processed, the user could be interested in skipping them. Typing the wave function file name and replacing the extension by "skr", this option will be set:

\triangleright touch grid_file_name.skr

The default value is 5.0×10^{-9} , a very reasonable cut-off according to our tests, but it is modifiable adding a new value in the file:

```
\triangleright echo 1.e-9 > grid_file_name.skr
```

Setting σ/π separation or only including π spin orbitals in the calculation [Go To Table Of Contents]

For many years in our research group, we have used a file that includes the identification of the σ/π symmetry of the spin orbitals obtained by a program we called "simon.x", and the outputs use the ".som" extension. It includes some text including the identification and it is not the easiest way to save a list of π spin orbitals for an user from out of our group. For this reason, Fukui is also compatible with a more simplified format:

```
Structure and format of a .sg file

#MO #piMO #sigmaMO #no_identifiedMO

\pi spin orbitals specification

\sigma spin orbitals specification

Specification of unidentified spin orbitals
```

Here is an example for tetracene:

Therefore, the tetracene has 60 MOs, 9 of which have π symmetry, and 51 are σ ones. The contents of the previous example can be saved in a file with the wave function file name and ".sg" extension.

When the user needs to obtain populations or AOMs defined by π MOs only, we can create an empty file with the extension ".pi" and the wave function file name:

⊳ touch wfn file name.pi

The calculation will be notably faster in addition.

Enabling the calculation of atomic overlap matrices (AOMs) [Go To Table Of Contents]

The atomic overlap matrices (AOMs) are needed as a previous step to calculate bond orders, exchange-correlation Fukui functions and/or N-delocalization indices. This program allows computing it for any region of the molecular space and/or atomic partitioning while a grid of points is provided.

Similarly to other options, the user will create an empty file to activate this computation using the wave function file name and the ".aom" extension.

⊳ touch wfn file name.aom

Enabling the calculation of the dipole moment [Go To Table Of Contents]

Dipole moment is locally defined as:

$$\mu_{\alpha} = -\int r_{\alpha} \rho(\vec{r}) \, d\vec{r} \tag{3.1}$$

where α is the corresponding Cartesian component (x, y or z) of the position vector and the dipole moment. Equation 3.1 is easily partitioned by including a real space weight function $w_A(\vec{r})$ for an hypothetical atom A with respect to the position vector \vec{r} :

$$\mu_{\alpha}(\mathbf{A}) = -\int w_{A}(\vec{r}) r_{\alpha} \rho(\vec{r}) \, d\vec{r}$$
(3.2)

One can always add a null term to the expression, keeping it unmodified:

$$\mu_{\alpha}(\mathbf{A}) = \overbrace{(R_{\mathbf{A}}^{\alpha} - R_{\mathbf{A}}^{\alpha})N_{\mathbf{A}}}^{=0} - \int w_{A}(\vec{r})r_{\alpha}\rho(\vec{r})\,\mathrm{d}\vec{r}$$
(3.3)

 $N_{\rm A}$ is the population of A according to the real space partitioning and $R_{\rm A}^{\alpha}$ corresponds to the α Cartesian component of the nucleus A. Therefore, keep in mind that you will need to specify the atom as argument to divide the dipole moment in the terms obtained below. Expanding $N_{\rm A}$ and restructuring the expression, we obtain:

$$\mu_{\alpha}(\mathbf{A}) = \overbrace{-\int (-R_{\mathbf{A}}^{\alpha}) w_{A}(\vec{r}) \rho(\vec{r}) \, \mathrm{d}\vec{r} - R_{\mathbf{A}}^{\alpha} N_{\mathbf{A}} - \int w_{A}(\vec{r}) r_{\alpha} \rho(\vec{r}) \, \mathrm{d}\vec{r}}^{R_{\mathbf{A}}^{\alpha} N_{\mathbf{A}}} - \int w_{A}(\vec{r}) r_{\alpha} \rho(\vec{r}) \, \mathrm{d}\vec{r}}$$
(3.4)

$$= \underbrace{-R_{\rm A}^{\alpha} N_{\rm A}}_{\mu_{\alpha}^{\rm intr}(A)} - \int (r_{\alpha} - R_{\rm A}^{\alpha}) w_{A}(\vec{r}) \rho(\vec{r}) \, d\vec{r}$$

$$\underbrace{-\frac{\mu_{\alpha}^{\rm CT}(A)}{\mu_{\alpha}^{\rm intr}(A)}}_{\mu_{\alpha}^{\rm intr}(A)}$$
(3.5)

This decomposition allows us to define two terms. On the one hand, $\mu_{\alpha}^{CT}(A)$ stands for a component dependent on the absolute position of the atom A and called charge transfer. On the other hand, $\mu_{\alpha}^{intr}(A)$ corresponds to a term uniquely centered on A, i.e., an intrinsic component.

Similarly to other options, the user will create an empty file to activate this computation using the wave function file name and the ".dip" extension.

> touch wfn file name.dip

Including the word "OnlyGrid" (case insensitive) inside, the program only compute the dipole moment from the density in the file and stops, omitting the rest of the computation:

 \triangleright echo "OutGrid" > wfn file name.dip

4 Output of a successful calculation

In this section we will discuss the program output on screen (Section 4.1) and the differences with the saved file (Section 4.2).

4.1 Output of a successful calculation on screen [Go To Table Of Contents]

The structure of the output on screen will be explained below. We will use all the options enabled to obtain the population, σ/π separation, the dipole moment computation and the atomic overlap matrices of a .stk file for the optimized tetracene molecule obtained at B3LYP/6-31G** level. The first text we will observe is the version, in this case 231003. After that, the wave function and grid files we are using, that is, "tetracene.wfn" and "tetracene_C001.stk", respectively. To enable the parallelization, a .proc file is needed. If you will not use the full available computer threads, specify the number in the file. Next, we will see simple information about the parallelization and a "Hello" message coming from each thread. The command we use is:

```
▷ ./fukui.x tetracene.wfn tetracene C001.stk 1 (part 1 of 6)
    PROGRAM FUKUI 231003
3
    >> Wave function file found: tetracene.wfn
      Grid file using .stk format: tetracene_C001.stk
       tetracene.proc was found. Parallelization enabled!!
    ********** THREAD INFORMATION ***********
9
      Job running using OpenMP.
10
      The number of processors is The number of threads is \dots
    Hello from process
12
13
   Hello from
Hello from
14
               process
15
    Hello from
16
    Hello from
               process
17
         from
          from
              process
19
20
                    clock time
21
22
```

Next, the default maximum resources will be printed. If you need to change them, create the .mxal file following the instruction in Section 3.5.

```
./fukui.x tetracene.wfn tetracene_C001.stk 1 (part 2 of 6)

// Using default max array allocations:
natomx nommx npmx nprimx
50 250 1000000 1000
```

The program detects a .sg file. It will read the orbital symmetry specification. Due to the fact that we add the third argument to specify the atom in which the grid is centered on, Fukui will skip gaussian functions far from this atom (with distance greater than 18.0 au by default). Next, we have created the .skg and .skr files and, therefore, the program will skip the Gauss quadrature weight functions and electron density values lower than the printed cut-off. In addition, we set the calculation of the AOM creating the "tetracene.aom" file. Finally, the computation of the dipole moment will be carried out due to the existence of the "tetracene.dip" file.

The program will detect if the .stk is a Fortran unformatted file. Next, it will read both wave function and grid files, verifying all the components are correct and printing how many points will be skipped with our current settings (defined in the .skg and .skr files or using the default cut-off values). The following step, if a .sg or .som file is available, is to identify the π MOs. Before the real calculation starts, the program will print the percentage of gaussian functions considered. This option is very interesting to reduce drastically the dependency with the molecule size. The last line before the computation corresponds to the calculation type: e dens (electron density), dip mom (dipole moment) and/or atomic overlap matrix (AOM). In this example, the program will compute all the available options (e dens + dip mom + AOM).

```
▷ ./fukui.x tetracene.wfn tetracene C001.stk 1 (part 4 of 6)
   >> Opening grid file: tetracene_C001.stk
   * unformatted grid file detected after 23 iterations
3
   [TT] Settings prepared in ...... 0.00 seconds
5
6
7
   >> Reading wave function file:
   * 60 MOs, 588 gaussian functions, 30 atoms.
8
     Coordinates and nuclear charges ..... OK
9
     Primitive centers .....
                                            OK
10
     Primitive types .....
11
     Primitive exponents
12
       coefficients .....
13
14
   [TT] Wave function read in ..... 0.02 seconds
15
16
   >> Reading grid file with a set of points:
     82466 points (total)
Binary stk file
17
18
19
     29154 of 82466 points were considered -> 64.65% skipped
20
^{21}
   [TT] Grid file read in ...... 0.03 seconds
22
23
   >> Opening sigma/pi separation file:
24
   * PI orbitals read .......
                                      ..... 46,48,51,55-60 PI(9)
25
   >> Preparing calculation:
26
   * Percentage of gaussian functions used ... 88.10\%
27
   [TT] Intermediate steps in ...... 0.00 seconds
```

We will see how the program splits the grid points among the threads of the computer we are requested:

```
▷ ./fukui.x tetracene.wfn tetracene C001.stk 1 (part 5 of 6)
     >> Computing grid points:
2
       CPU
            2: from 10933 to 14577 of 29154
            0: from 1 to 3644 of 29154
5: from 7289 to 10932 of 29154
1: from 14578 to 18221 of 29154
6: from 3645 to 7288 of 29154
3
       CPU 0:
4
5
       CPII
       CPU
6
       CPU
            3: from
                      25510 to 29154 of 29154
       CPU
       CPU
                      21866
                                 25509 of
                from
            4: from
9
       CPU
                      18222
                             to 21865 of 29154
10
     [TT] Grid points computed in ...... 0.40 seconds
```

The only results the program prints on screen are the total populations and dipole moment separation, removing the excessive data from the MO populations and the AOMs.

```
RESULTS OF THE INTEGRATION (ATOM 1)
POP = 6.085619749597D+00 au
 2
 3
    REF POP
                              6.085620961232D+00 au
 4
    FUKUI = -0.000 au
8
9
    DTP MOM
      INTRINSIC
                            2.80764995D-15
                                              -1.44706705D-03
                                                                 -4.48692077D-02
10
      CHARGE TRANSFER
                            -0.0000000D+00
                                               -5.62150045D+01
                                                                 -8.22492103D+00
                                                                                   au
                                                                 -8.26979024D+00
11
      TOTAL
                            2.80764995D-15
                                               -5.62164516D+01
                                                                                   au
12
    REF DIP MOM
                                              -1.44598784D-03
13
      INTRINSIC
                           -2.16778707D-15
                                                                 -4.48687786D-02
                                                                 -8.22492267D+00
-8.26979145D+00
14
      CHARGE TRANSFER
                           -0.0000000D+00
                                              -5.62150157D+01
                                                                                   au
      TOTAL
                           -2.16778707D-15
                                              -5.62164617D+01
15
16
17
    Considering SKIPPING:
18
    REF POP = 6.08562062
DIF REF POP = 3.407D-07 au
FUKUI = -0.000 au
19
                     6.08562062048363D+00 au
\frac{20}{21}
22
23
    REF DIP MOM
24
      INTRINSIC
                            -2.16599752D-15
                                               -1.44680540D-03
                                                                 -4.48691350D-02
25
      DIF INTRINSIC
                           -1.78955799D-18
                                               8.17557978D-07
                                                                  3.56385705D-07
26
27
      CHARGE TRANSFER
                                -0.0000000D+00
                                                   -5.62150126D+01
                                                                      -8.22492221D+00
      DIF CHARGE TRANSFER =
                                                                     -4.60533116D-07
                                0.0000000D+00
                                                  -3.14761335D-06
28
                                                                 -8.26979134D+00
                                              -5.62164594D+01
                           -2.16599752D-15
                                                                                  au
      DIF TOTAL
                            -1.78955799D-18
                                               -2.33005537D-06
                                                                 -1.04147410D-07
30
31
    SIGMA/PI CONTRIBUTIONS:
                                   molecular orbitals with PI symmetry
32
    SIGMA
            5.14519
                              0.94043
33
    0.54 seconds
34
```

Each value represents:

POP

is the total population computed by the program from the wave function

REF POP

is the population from the .stk file (not computed using the wave function file)

FUKUI

corresponds to the difference between REF POP and POP. Depending on the kind of calculation we are performing could be the Fukui index.

DIP MOM

starts the section for the corresponding dipole moment results, following Equation 3.5 formulated in Section 3.5 and computing the electron density.

- INTRINSIC: $\mu_{\alpha}^{intr}(A)$.
- CHARGE TRANSFER: $\mu_{\alpha}^{\rm CT}({\bf A})$.
- TOTAL: sum of the two previous components.

REF DIP MOM

corresponds to the dipole moment computed from the electron densities included in the .stk or .cube file, following Equation 3.5 formulated in Section 3.5.

- INTRINSIC: $\mu_{\alpha}^{intr}(A)$.
- CHARGE TRANSFER: $\mu_{\alpha}^{\text{CT}}(A)$.
- TOTAL: sum of the two previous components.

Considering SKIPPING: values obtained from the sum over the smaller grid, skipping points with negligible Gauss quadrature weights and electron density values.

REF POP this value represents the overall sum from the densities in the .stk file.

DIF REF POP

is the subtraction between both REF POP values, with and without skipping. As you can observe, the condensed populations are affected in the 7th decimal position. Therefore, our settings are a very good approach.

FUKUI

is defined identically as the another FUKUI value but including only the smaller grid of points.

REF DIP MOM corresponds to the dipole moment computed from the electron densities included in the .stk or .cube file but with a smaller set of points.

- INTRINSIC: $\mu_{\alpha}^{intr}(A)$.
- DIF INTRINSIC: difference between both INTRINSIC components.
- CHARGE TRANSFER: $\mu_{\alpha}^{CT}(A)$.
- DIF CHARGE TRANSFER: difference between both CHARGE TRANSFER components.
- TOTAL: sum of the two previous components.
- DIF TOTAL: difference between both TOTAL dipole moment values.

Finally, the results of the σ/π separation, the time spent on the execution and the performance are printed.

4.2 File output of a successful calculation [Go To Table Of Contents]

Part of the information on screen, discussed in Section 4.1, is also written in a file called as "wavefunction grid file.fuk". We will discuss the new parts with respect to the output on screen (Section 4.1). Most of the additions are after printing the populations and dipole moment (if requested):

```
    ▷ cat wavefunction grid file.fuk (part 1 of 2)

             ORBITAL CONTRIBUTIONS:
               N(1)
                                   0.000000 au
                   2)
 3
                                   0.00000

    \begin{array}{r}
      4 \\
      5 \\
      6 \\
      7 \\
      8 \\
      9 \\
      10
    \end{array}

                                   0.000035
                   4)
5)
                                   0.000035
                                   0.000036
                   6)
7)
8)
                                   0.000036
                                   0.424963
                    9)
11
               N(10)
                                   0.337798 au
12
13
14
               N(57)
N(58)
15
16
17
                                   0.140636
               N(59)
                                       101862
               N(60)
                                   0.081479 au
```

As you can see, Fukui prints the orbital contributions for the total populations. Next, if the .sg or .som file is available, the σ/π separation and, finally, the AOM using an 8-column format of real values in scientific notation:

```
    ▷ cat wavefunction grid file.fuk (part 2 of 2)

              The Atomic Overlap Matrix
2
3
4
5
6
       0.525717804208E-08
                            0.176756945118E-07
      0.735609271729E-08
```

Lastly the elapsed time and performance.

5 Utilities

Some small programs or utilities have been included. They are independent from the main program, and some of them (with .f90 extension) are compiled in the usual way (compiler source_code.f90 -o executable_name.x).

5.1 bstk astk.f90 [Go To Table Of Contents]

This small program transforms Fortran formatted .stk files (ASCII text) into unformatted ones (binary data) and vice versa. The program needs the reference .stk file (only include the file name without extension) and the type of file we are using. If no arguments, the program will ask for them.

5.2 cube2stk.f90 [Go To Table Of Contents]

This utility transforms a cube file into an unformatted .stk file. It includes a short help and its use is "cube2stk.x cube_file atom_number". The last argument is optional.

5.3 fukui.sh [Go To Table Of Contents]

This BASH script is useful to systematize the execution of a set of .stk files using the following file name scheme: "stkfilename_X001.stk", being "stkfilename" the root file name of the .stk file, "X" the element symbol and "001" the corresponding atom number in the molecule. The script will take all the file names and arguments automatically. For example, "./fukui.sh tetracene_C001.stk" will call the Fukui program using the wave function called "tetracene.wfn", and it will include the third argument "1" to enable the primitive skipping. The program will also create the .mxal with the resources needed.

5.4 mwfn2piorbs.sh [Go To Table Of Contents]

This BASH script will obtain the .sg file to carry out the σ/π separation using the program called Multiwfn. Edit the script to comply with the corresponding executable paths.

5.5 som2sg.sh [Go To Table Of Contents]

This BASH script is for internal use. It transforms .som into .sg files to perform the σ/π separation.

Bibliography

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A Appendices

A.1 Obtaining PROAIMS wave function file [Go To Table Of Contents]

We will see how to obtain a PROAIMS wave function file with an example for the Gaussian suite of programs. More concretely, we have considered the benzene molecule:

```
⊳ cat C6H6.gjf
    %nproc=4
    %mem=1gb
%chk=C6H6.chk
3
    # b3lyp 6-31g(d,p) 6d integral=ultrafinegrid out=wfn
5
6
7
8
9
    Title Card Required
10
                          0.0000000
                                           1.39499067
                                                           0.0000000
11
                          -1.20809735
                                           0.69749533
                                                           0.0000000
12
13
14
15
16
17
18
19
20
21
                          -1.20809735
                                          -0.69749533
                                                           0.0000000
                          0.0000000
                                          -1.39499067
                                                           0.0000000
                                          -0.69749533
                           1.20809735
                                                           0.00000000
                           1.20809735
                                           0.69749533
                                                           0.0000000
                           0.0000000
                                           2.49460097
                                                           0.0000000
                          2.16038781
                                           1.24730049
                                                           0.0000000
                          -2.16038781
                                            .24730049
                                                           0.0000000
                          0.0000000
                                          -2.49460097
                                                           0.0000000
                          2.16038781
                                             24730049
                                                           0.0000000
                          2.16038781
                                           1.24730049
                                                           0.0000000
    C6H6.wfn
```

In order to print a .wfn file, the label out=wfn must be included. There exist equivalent forms of this label, such as output=wfn, out=psi and output=psi.

A.2 Converting from another file format to PROAIMS wave function [Go To Table Of Contents]

One of the most updated software with an active development to convert among formats is the Multiwfn program (http://sobereva.com/multiwfn/). In the version 3.7 the steps to obtain the .wfn file are shown below:

```
Multiwfn -- A Multifunctional Wavefunction Analyzer
Version 3.7, release date: 2020-Aug-14
Project leader: Tian Lu (Beijing Kein Research Center for Natural Sciences)
Below paper ***MUST BE CITED*** if Multiwfn is utilized in your work:
Tian Lu, Feiwu Chen, J. Comput. Chem., 33, 580-592 (2012)
 2
 6
         Total/Alpha/Beta electrons:
                                                                     42.0000
                                                                                            21.0000
                                                                                                                   21.0000
       Net charge: 0.00000 Expected multiplicity: 1

Atoms: 12, Basis functions: 120, GTFs: 210

Total energy: -232.113832873650 Hartree, Virial ratio: 2.00945783

This is a restricted single-determinant wavefunction
 8
 9
10
11
       Orbitals from 1 to 21 are occupied
Title line of this file: Title Card Required
12
13
14
        Loaded C6H6.fchk successfully!
15
16
        Formula: H6 C6
        Molecule weight:
17
       Point group: D6h
18
19
       "q": Exit program gracefully "r": L
******** Main function menu **********
0 Show molecular structure and view orbitals
                                                                               "r": Load a new file
20
21
23
           Output all properties at a point
^{24}
           Topology analysis
       2 Topology analysis
3 Output and plot specific property in a line
4 Output and plot specific property in a plane
5 Output and plot specific property within a spatial region (calc. grid data)
6 Check & modify wavefunction
7 Population analysis and atomic charges
25
26
27
28
29
30
       8 Orbital composition analysis
       9 Bond order analysis
10 Plot total DOS, partial DOS, OPDOS, local DOS and photoelectron spectrum
11 Plot IR/Raman/UV-Vis/ECD/VCD/ROA/NMR spectrum
31
32
33
       12 Quantitative analysis of molecular surface
34
       13 Process grid data (No grid data is presented currently)
14 Adaptive natural density partitioning (AdNDP) analysis
35
36
37
        15 Fuzzy atomic space analysis
38
        16 Charge decomposition analysis (CDA) and plot orbital interaction diagram
       17 Basin analysis
19 Orbital localization analysis
21 Energy decomposition analysis
100 Other functions (Part 1)
300 Other functions (Part 3)
39
                                                                              18 Electron excitation analysis 20 Visual study of weak interaction
40
42
                                                                              200 Other functions (Part 2)
```

\triangleright Type "100" (without inverted commas) and the program will print a text similar to this:

```
======== Other functions (Part 1) =========
 2
        0 Return
        1 Draw scatter graph between two functions and generate their cube files
2 Export various files (mwfn/pdb/xyz/wfn/wfx/molden/fch/47/mkl...) or generate input file of quantum che
 3
       2 Export various files (mwin/pub/xyz/win/wix/morden/ith/4//mxi...) of generate 3 Calculate molecular van der Waals Volume
4 Integrate a function in whole space
5 Show overlap integral between alpha and beta orbitals
6 Monitor SCF convergence process of Gaussian
8 Generate Gaussian input file with initial guess from fragment wavefunctions
 5
 6
 8
 9
           Evaluate interatomic connectivity and atomic coordination number 1 Calculate overlap and centroid distance between two orbitals
10
        12 Perform biorthogonalization between alpha and beta orbitals
13 Calculate HOMA and Bird aromaticity index
14 Calculate LOLIPOP (LOL Integrated Pi Over Plane)
12
13
14
        15 Calculate intermolecular orbital overlap
16 Calculate various quantities in conceptual density functional theory (CDFT)
15
16
        18 Yoshizawa's electron transport route analysis
        19 Generate promolecular .wfn file from fragment wavefunctions
20 Calculate Hellmann-Feynman forces
21 Calculate properties based on geometry information for specific atoms
22 Detect pi orbitals, set occupation numbers and calculate pi composition
23 Fit function distribution to atomic value
18
19
20
21
        24 Obtain NICS_ZZ value for non-planar or tilted system
```

```
> Type "2" (without inverted commas) and the program will print a text similar to this:
    0 Return
    Export system to various formats of files:  \\
      Output current structure to .pdb file Output current structure to .xyz file
3
 5
       Output current structure and atomic charges to .chg file
      Output current wavefunction as .wfx file Output current wavefunction as .wfn file
 6
      Output current wavefunction as Molden input file (.molden) Output current wavefunction as .fch file
       Output current wavefunction as .47 file
11
      Output current wavefunction as old Molekel input file (.mkl)
12
    31 Output current structure to .cml file
    32 Output current wavefunction as .mwfn file
13
    Generate input file of quantum chemistry codes:
14
15
    10 Gaussian
16
    11 GAMESS-US
17
       ORCA
                               13 NWChem
    14 MOPAC
16 MRCC
                               15 PSI4
17 CFOUR
18
19
20
    18
20
       Molpro
Molcas
                               19
                                  Dalton
                               21 Q-Chem
    ▷ Type "5" (without inverted commas) and the program will print a text similar to this:
    Input path for exporting file, e.g. C:\ltwd.wfn
```

▷ Type the file name: C6H6.wfn

> Exit following the menu codes.

You will find the PROAIMS wave function file called "C6H6.wfn" in the current directory together with the reference file.

Another way to convert from .fchk to .wfn format is using the AIMAll program.