# TCCM homework 1: Computation of the MP2 energy

A. Ammar, Y. Damour, P. Reinhardt, A. Scemama November 22, 2024

You can work in groups of up to three people. We expect that you will use Git to work collaboratively on your project. We expect the following files and structure in your submission:

- A LICENSE file specifying the license for your code.
- An AUTHORS file listing the names of all contributors.
- A README.md file providing a brief description of the directory structure and the project.
- An INSTALL.md file with clear instructions on how to compile and run the program.
- A tests directory containing tests to ensure that the program behaves as expected.
- (Optional) A doc directory for additional documentation, if the project requires more detail than the README.md file can provide.
- A src directory containing all the source files of your program.

In this homework, you will read data (integrals, orbital energies) from a file and use it to compute the Hartree-Fock energy and the MP2 correlation energy for a closed-shell system.

## 1 Theory

#### 1.1 Hartree-Fock Energy Calculation

Using the one- and two-electron integrals in the molecular orbital (MO) basis, the closed-shell ( $N_{\uparrow} = N_{\downarrow}$ ) Hartree-Fock energy can be computed as:

$$E = E_{\rm NN} + 2\sum_{i=1}^{N_{\rm occ}} \langle i|h|i\rangle + \sum_{i=1}^{N_{\rm occ}} \sum_{j=1}^{N_{\rm occ}} \left[2\langle ij|ij\rangle - \langle ij|ji\rangle\right] \tag{1}$$

where:

- $E_{\rm NN}$  is the nuclear repulsion energy,
- 2  $\sum_i \langle i|h|i\rangle$  represents the kinetic energy and electron-nucleus potential term,
- $\langle pq|rs\rangle$  are the two-electron integrals,
- Indices i, j run over the occupied molecular orbitals.

Warning: The number of occupied orbitals,  $N_{\rm occ}$ , represents the number of up electrons  $N_{\uparrow}$  (or the number of down electrons  $N_{\downarrow}$ ). The formula (1) is given in terms of spatial orbitals, not spin orbitals.

#### 1.2 MP2 Energy Correction Calculation

If the MOs are eigenfunctions of the Fock matrix, the MP2 energy is given by:

$$E_{\text{MP2}} = \sum_{(i,j) \in \text{occupied } (a,b) \in \text{virtuals}} \frac{\langle ij|ab\rangle \left(2\langle ij|ab\rangle - \langle ij|ba\rangle\right)}{e_i + e_j - e_a - e_b} \tag{2}$$

where:

- $e_p$  is the energy of the p-th orbital (the p-th eigenvalue of the Fock matrix),
- i, j label occupied orbitals,
- $\bullet$  a, b label virtual orbitals.

# 2 Programming

#### 2.1 Installing TREXIO

You will need to read data from a TREXIO file, a format and library designed for exchanging wave function data between quantum chemistry programs. First, install the TREXIO library on your machine.

TREXIO depends on the HDF5 library, which can be installed using your package manager. On Ubuntu:

sudo apt install libhdf5-dev

On macOS:

brew install hdf5

Download TREXIO here: https://github.com/TREX-CoE/trexio/releases/download/v2.5.0/trexio-2.5.0.tar.gz and install it on your system:

tar -zxvf trexio-2.5.0.tar.gz
cd trexio-2.5.0
./configure
make
sudo make install

This installs the TREXIO library to /usr/local/lib.

Documentation for TREXIO is available here: https://trex-coe.github.io/trexio/.

### 2.2 Accessing Data from TREXIO Files

Include the TREXIO header in all source files that use the library:

#include <trexio.h>

A few sample files are provided in the data directory.

#### 2.2.1 Opening and Closing TREXIO Files

To open a TREXIO file:

```
trexio_exit_code rc;
trexio_t* trexio_file = trexio_open(filename, 'r', TREXIO_AUTO, &rc);
if (rc != TREXIO_SUCCESS) {
   printf("TREXIO Error: %s\n", trexio_string_of_error(rc));
   exit(1);
}
```

To close the TREXIO file:

```
rc = trexio_close(trexio_file);
if (rc != TREXIO_SUCCESS) {
  printf("TREXIO Error: %s\n", trexio_string_of_error(rc));
  exit(1);
}
trexio_file = NULL;
```

Each TREXIO function returns an exit code (trexio\_exit\_code), where TREXIO\_SUCCESS indicates success.

#### 2.2.2 Linking with TREXIO

Compile your program with the -ltrexio flag and specify the include and library paths:

```
gcc -I/usr/local/include -L/usr/local/lib -ltrexio my_code.c -o my_code
```

#### 2.2.3 Reading the Nuclear Repulsion Energy

Use this function to read the nuclear repulsion energy:

Be careful, this function takes as second argument a a pointer to a double variable. Similarly to scanf, you need to pass the address to modify a variable. You will need to use this function as

A correct implementation should return a nuclear repulsion energy of 9.19497 atomic units for the water molecule.

#### 2.2.4 Obtaining the Number of Occupied Orbitals

Recall that for a Hartree-Fock wave function the number of occupied orbitals is equal to the number of up-spin electrons. You can obtain the number of up-spin electrons using the following function:

#### 2.2.5 Reading One-Electron Integrals

Read the number of molecular orbitals using:

Read the one-electron integrals (core Hamiltonian) using:

The data array should be large enough to hold  ${\tt mo\_num} \times {\tt mo\_num}$  elements.

#### 2.2.6 Reading Two-Electron Integrals

Two-electron integrals are stored in a sparse format. First, get the number of non-zero integrals:

Allocate memory for indices and values:

```
int32_t* const index = malloc(4 * n_integrals * sizeof(int32_t));
if (index == NULL) {
    fprintf(stderr, "Malloc failed for index");
    exit(1);
}

double* const value = malloc(n_integrals * sizeof(double));
if (value == NULL) {
    fprintf(stderr, "Malloc failed for value");
    exit(1);
}
```

and read the integrals from the file using

where offset\_file=0 and buffer\_size = n\_integrals. Warning: you should pass the address of buffer\_size, because on output of this function buffer\_size contains the number of read integrals.

To access the *n*-th integral, you can get the 4 indices (i, j, k, l) and the corresponding value  $\langle ij|kl\rangle$  using

```
int i = index[4*n+0];
int j = index[4*n+1];
int k = index[4*n+2];
int l = index[4*n+3];
double integral = value[n];
```

Warning: Two-electron integrals obey 8-fold permutational symmetry:

$$\langle ij|kl\rangle = \iint \phi_i(\mathbf{r}_1)\phi_j(\mathbf{r}_2) \frac{1}{|\mathbf{r}_1 - \mathbf{r}_2|} \phi_k(\mathbf{r}_1)\phi_l(\mathbf{r}_2) d\mathbf{r}_1 d\mathbf{r}_2$$

- i and k can be swapped
- j and l can be swapped
- $\mathbf{r}_1$  and  $\mathbf{r}_2$  can be swapped

So  $\langle ij|kl\rangle = \langle il|kj\rangle = \langle kl|ij\rangle = \langle kj|il\rangle = \langle ji|lk\rangle = \langle li|jk\rangle = \langle lk|ji\rangle = \langle jk|li\rangle$ . To minimize the size of the file, for each given quartet of indices only one permutation is stored in the TREXIO file. In what follows, you must handle this symmetry during computations.

## 2.3 Computing the Hartree-Fock Energy

Use the data you have read to implement Eq.(1) and compute the Hartree-Fock energy. The expected result is -76.0267987 atomic units for the water molecule.

#### 2.4 Computing the MP2 Energy Correction

To compute the MP2 energy correction:

• Read orbital energies using:

• Implement Eq.(2).

The expected MP2 correction is -0.20395997 atomic units for the water molecule.