A practicalist guide to Self Consistent Field

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I. NOTATION

There exist many ways to label and describe quantum chemistry theories. In this section the quantities and indices used will be defined and outlined. Generalized multidimensional arrays will be referred to as "tensors" and two dimensional arrays will be referred to as "matrices".

A. Einstein summation notation

The generalized Einstein summation notation will be employed where repeated indices on the right hand side (RHS) that do not appear on left hand side (LHS) are assumed to be summed over. A matrix-matrix multiplication example:

Matrix convention:
$$C = AB$$
 (1)

Explicit Sum :
$$C_{ij} = \sum_{k} A_{ik} B_{kj}$$
 (2)

Einstein summation:
$$C_{ij} = A_{ik}B_{kj}$$
 (3)

an outer product example

$$C_{ikj} = A_{ik}B_{kj} \tag{4}$$

B. Indices

The following notation will be adhered to throughout this document. Occasionally indices will be redefined, but it should be clear in the context what the alternate meaning is.

• General atomic orbitals $\mu, \nu, \lambda, \sigma$

• Inactive (doubly occupied) molecular orbitals i, j, k, l

• Unoccupied (virtual) molecular orbitals a, b, c, d

• General index molecular orbitals p, q, r, s, m, n, o

C. Tensors

Common matrices and tensors utilized in this work.

• Core Hamiltonian matrix $H_{\mu\nu}$

• Molecular orbital coefficient matrix $C_{p\mu}$

• Overlap matrix $S_{\mu\nu}$

• SCF one-particle density matrix D_{pq}

• Atomic-orbital two-electron integral $g_{\mu\nu\lambda\sigma} = \int \mu(\mathbf{r}_1)\nu(\mathbf{r}_1)\frac{1}{r_{12}}\lambda(\mathbf{r}_2)\sigma(\mathbf{r}_2)d\mathbf{r}_1d\mathbf{r}_2$

II. SELF CONSISTENT FIELD

Here we will focus on a variant of Self Consistent Field (SCF) known as restricted Hartree-Fock (RHF) where the α and β orbitals are equal denoting a closed-shell molecule with all paired electrons. To begin, let us start with the AO Fock matrix, which is represented as^{1,2}

$$F_{\mu\nu} = H_{\mu\nu} + 2g_{\mu\nu\lambda\sigma}D_{\lambda\sigma} - g_{\mu\lambda\nu\sigma}D_{\lambda\sigma} \tag{5}$$

and the one-particle RHF density matrix D is computed from the orbitals C (assumed real).

$$D_{\lambda\sigma} = C_{i\sigma}C_{i\lambda} \tag{6}$$

It should be noted that if D is converted to the MO basis, we have a diagonal matrix of ones up to the number of occupied indices and zero thereafter

$$D_{pq} = \begin{pmatrix} 1_{ii} & 0 \\ 0 & 0 \end{pmatrix} \tag{7}$$

The total RHF energy can be expressed as a sum of the electronic and Born-Oppenheimer (BO) nuclear energies

$$E^{RHF} = E_{electronic}^{RHF} + E_{nuclear}^{BO} \tag{8}$$

$$E_{electronic}^{RHF} = (F_{\mu\nu} + H_{\mu\nu})D_{\mu\nu} \tag{9}$$

$$E_{nuclear}^{BO} = \sum_{i>j} \frac{Z_i Z_j}{r_{ij}} \tag{10}$$

where Z_i is the nuclear charge of atom i.

Examining these equations, it is clear that the most computationally demanding portions are the convolution of the density matrix with the two-electron integral tensor. Therefore, it is often convenient to define the following quantities

$$J[D_{\lambda\sigma}]_{\mu\nu} = g_{\mu\nu\lambda\sigma}D_{\lambda\sigma} \tag{11}$$

$$K[D_{\lambda\sigma}]_{\mu\nu} = g_{\mu\lambda\nu\sigma}D_{\lambda\sigma} \tag{12}$$

The resulting J and K matrices are often called the Coulomb and Exchange matrices, respectively. Often quantum chemistry programs have very efficient routines to compute these equations, thus utilizing these routines is of the utmost importance.

A. Roothaan Equations

The SCF equations are often solved through the Roothan equations², shown in matrix formalism

$$FC = SC\epsilon \tag{13}$$

This is a generalized pseudo-eigenvalue problem, where **S** represents our orthogonality constraints. At every iteration we need to solve for the coefficients **C** that diagonalize the Fock matrix. An ideal world would produce AO orbitals that are orthogonal, unfortunately, as demonstrated by non-diagonal overlap matrices, this is not the case. To overcome this problem, a orthonormalized Fock matrix is diagonalized instead, shown in matrix formalism

$$\mathbf{A} = \mathbf{S}^{-1/2} \tag{14}$$

$$\mathbf{F}' = \mathbf{A}^T \mathbf{F} \mathbf{A} \tag{15}$$

$$\mathbf{F}'\mathbf{C}' = \mathbf{C}'\epsilon \tag{16}$$

$$\mathbf{C} = \mathbf{AC'} \tag{17}$$

At every iteration we construct our Fock matrix from the previous orbitals (\mathbf{C}_{n-1}) and compute new orbitals (\mathbf{C}_n) until convergence is reached.

For the rest of this section, the "physicist's water molecule" (O-H = 1.1Å, \angle HOH = 104°) will be utilized in the cc-pVDZ basis set for illustrating convergence patterns. A pure iterative diagonalization approach is shown in Fig. 1. As can be seen, the convergence for this simple molecule is quite slow and the next sections will detail convergence acceleration.

FIG. 1. A RHF computation of "physicist's water molecule" starting with a core Hamiltonian guess in the cc-pVDZ basis set. dE is the energy difference between iterations and dRMS is the root mean square of the orbital gradient.

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1: Energy = -68.98003273414295
RHF Iteration
                                               dE = -6.898E+01
                                                                 dRMS = 1.165E-01
RHF Iteration
               2: Energy = -69.64725442845806
                                               dE = -6.672E-01
                                                                 dRMS = 1.074E-01
               3: Energy = -72.84030309363035
RHF Iteration
                                               dE = -3.193E+00
                                                                 dRMS = 1.039E-01
RHF Iteration
               4: Energy = -72.89488390650019
                                               dE = -5.458E-02
                                                                 dRMS = 8.660E-02
RHF Iteration
               5: Energy = -74.12078064688371
                                               dE = -1.225E+00
                                                                 dRMS = 8.646E-02
RHF Iteration
               6: Energy = -74.86718194576882
                                               dE = -7.464E-01
                                                                 dRMS = 6.528E-02
RHF Iteration 7: Energy = -75.41490878039029
                                               dE = -5.477E-01
                                                                 dRMS = 5.216E-02
RHF Iteration 22: Energy = -75.98979285429830
                                               dE = -3.768E-06
                                                                 dRMS = 1.188E-04
RHF Iteration 23: Energy = -75.98979450314014
                                                                 dRMS = 7.860E-05
                                               dE = -1.648E-06
RHF Iteration 24: Energy = -75.98979522446778
                                               dE = -7.213E-07
                                                                 dRMS = 5.198E-05
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REFERENCES

¹A. Szabo and N. S. Ostlund, *Modern Quantum Chemistry*, Introduction to Advanced Electronic Structure Theory. Courier Corporation, 1996.

²I. N. Levine, *Quantum Chemistry*. Prentice-Hall, New Jersey, 5th edition, 2000.