

Week 47, Coupled-Cluster theory

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Week 47, November 17-21, 2025

Thursday:

1. Coupled-cluster theory and basic equations, see also notes from last week at <https://github.com/ManyBodyPhysics/FYS4480/blob/master/doc/pub/week46/pdf/week46.pdf>

Friday:

1. Derivation of doubles excitation equation
2. Applications to the pairing model from the second midterm and discussions of second midterm
3. Lecture material: Lecture notes (these notes) and chapter 9 of Shavitt and Bartlett, in particular sections 9-1-9.3, see also slides from last week at <https://github.com/ManyBodyPhysics/FYS4480/blob/master/doc/pub/week46/pdf/week46.pdf>

Introduction

The coupled-cluster method is an efficient tool to compute properties of many-body systems with an effort that grows polynomial with system size. While this might still be expensive, it is now possible to compute for example atomic nuclei with mass numbers about $A \approx 208$ (which corresponds to lead) with this method. Recall that full configuration interaction (FCI) exhibits an exponential cost and is therefore limited to systems with few degrees of freedom.

For some recent applications see

1. **Nuclear Physics:** see

<https://www.nature.com/articles/s41567-022-01715-8>

2. **Quantum chemistry:** see <https://journals.aps.org/rmp/abstract/10.1103/RevModPhys.79.291>

The normal-ordered Hamiltonian

We start from the reference state

$$|\Phi_0\rangle = \prod_{i=1}^N a_i^\dagger |0\rangle \quad (1)$$

for the description of a system with N particles. Usually, this reference is the Hartree-Fock state, but that is not necessary as discussed throughout this course.

Notations again

After we introduced the particle-hole formalism, we have opted for a convention where the indices i, j, k, \dots run over hole states, i.e. orbitals occupied in the reference state, while a, b, c, \dots run over particle states, i.e. unoccupied orbitals. Indices p, q, r, s can identify any orbital. Let n_u be the number of unoccupied states, and N is of course the number of particles, which is also the number of occupied states used to define the ansatz for the ground state.

Hamiltonian

We consider the Hamiltonian

$$H = \sum_{pq} \langle p | h_0 | q \rangle a_p^\dagger a_q + \frac{1}{4} \sum_{pqrs} \langle pq | v | rs \rangle a_p^\dagger a_q^\dagger a_s a_r \quad (2)$$

The reference state

The reference state is a non-trivial vacuum of our theory. We normal order this Hamiltonian with respect to the nontrivial vacuum state given by the Hartree-Fock reference and obtain the normal-ordered Hamiltonian

$$H_N = \sum_{pq} \langle p|f|q\rangle \left\{ a_p^\dagger a_q \right\} + \frac{1}{4} \sum_{pqrs} \langle pq|V|rs\rangle \left\{ a_p^\dagger a_q^\dagger a_s a_r \right\}. \quad (3)$$

Hartree-Fock basis

Here,

$$\langle p|f|q\rangle = \langle p|h_0|q\rangle + \sum_i \langle pi|V|qi\rangle \quad (4)$$

is the Fock matrix. We note that the Fock matrix is diagonal in the Hartree-Fock basis. The brackets $\{\dots\}$ denote normal ordering, i.e. all operators that annihilate the nontrivial vacuum are to the right of those operators that create with respect to that vacuum. Normal ordering implies that $\langle \Phi_0 | H_N | \Phi_0 \rangle = 0$.

What does “good” mean?

The coupled-cluster method is a very efficient tool to compute properties of many-particle systems when a “good” reference state is available. Let us assume that the reference state results from a Hartree-Fock calculation.

How do you know whether a Hartree-Fock state is a “good” reference? Which results of the Hartree-Fock computation will inform you?

Answer

Once the Hartree-Fock equations are solved, the Fock matrix becomes diagonal, and its diagonal elements can be viewed as single-particle energies. Hopefully, there is a clear gap in the single-particle spectrum at the Fermi surface, i.e. after N orbitals are filled.

To nuclear physics aficionados: How many nuclei are accessible with the coupled cluster method based on spherical mean fields?

If symmetry-restricted Hartree-Fock is used, one is limited to compute systems with closed subshells for neutrons and for protons. On a first view, this might seem as a severe limitation. But is it? If one limits oneself to nuclei with mass number up to mass number $A = 60$, how many nuclei can potentially be described with the coupled-cluster method? Which of these nuclei are potentially interesting? Why?

Answer

Nuclear shell closures are at $N, Z = 2, 8, 20, 28, 50, 82, 126$, and subshell closures at $N, Z = 2, 6, 8, 14, 16, 20, 28, 32, 34, 40, 50, \dots$. In the physics of nuclei, the evolution of nuclear structure as neutrons are added (or removed) from an isotope is a key interest. Examples are the rare isotopes of helium (He-8,10) oxygen (O-22,24,28), calcium (Ca-52,54,60), nickel (Ni-78) and tin (Sn-100,132). The coupled-cluster method has the potential to address questions regarding these nuclei, and in several cases was used to make predictions before experimental data was available. In addition, the method can be used to compute neighbors of nuclei with closed subshells.

An FCI reminder to link with coupled-cluster theory and MBPT

We had

$$(\hat{H} - E)|\Psi_0\rangle = (\hat{H} - E) \sum_{P'H'} C_{H'}^{P'} |\Phi_{H'}^{P'}\rangle = 0.$$

As stated previously, one solves this equation normally by diagonalization. If we are able to solve this equation exactly (that is numerically exactly) in a large Hilbert space (it will be truncated in terms of the number of single-particle states included in the definition of Slater determinants), it can then serve as a benchmark for other many-body methods which approximate the correlation operator \hat{C} .

Rewriting the equations

For reasons to come (link with Coupled-Cluster theory and Many-Body perturbation theory), we will rewrite the full FCI equation as a set of coupled non-linear equations in terms of the unknown coefficients C_H^P .

To see this, we look at $\langle \Phi_H^P | = \langle \Phi_0 |$, that is we multiply with $\langle \Phi_0 |$ from the left in

$$(\hat{H} - E) \sum_{P'H'} C_{H'}^{P'} |\Phi_{H'}^{P'}\rangle = 0,$$

and we assume that we have a two-body operator at most. Using the Condon-Slater rules gives then an equation for the correlation energy in terms of C_i^a and C_{ij}^{ab} . We get then

$$\langle \Phi_0 | \hat{H} - E | \Phi_0 \rangle + \sum_{ai} \langle \Phi_0 | \hat{H} - E | \Phi_i^a \rangle C_i^a + \sum_{abij} \langle \Phi_0 | \hat{H} - E | \Phi_{ij}^{ab} \rangle C_{ij}^{ab} = 0,$$

or

$$E - E_0 = \Delta E = \sum_{ai} \langle \Phi_0 | \hat{H} | \Phi_i^a \rangle C_i^a + \sum_{abij} \langle \Phi_0 | \hat{H} | \Phi_{ij}^{ab} \rangle C_{ij}^{ab},$$

where the E_0 is the reference energy and ΔE becomes the

Slight rewrite

We can rewrite

$$E - E_0 = \Delta E = \sum_{ai} \langle \Phi_0 | \hat{H} | \Phi_i^a \rangle C_i^a + \sum_{abij} \langle \Phi_0 | \hat{H} | \Phi_{ij}^{ab} \rangle C_{ij}^{ab},$$

as

$$\Delta E = \sum_{ai} \langle i | \hat{f} | a \rangle C_i^a + \sum_{abij} \langle ij | \hat{v} | ab \rangle C_{ij}^{ab}.$$

Equations for the coefficients

This equation determines the correlation energy but not the coefficients C . We need more equations. Our next step is to multiply from the left with Φ_i^a

$$\langle \Phi_i^a | \hat{H} - E | \Phi_0 \rangle + \sum_{bj} \langle \Phi_i^a | \hat{H} - E | \Phi_j^b \rangle C_j^b + \sum_{bcjk} \langle \Phi_i^a | \hat{H} - E | \Phi_{jk}^{bc} \rangle C_{jk}^{bc} + \sum_{bcdjkl} \langle \Phi_i^a | \hat{H}$$

as this equation will allow us to find an expression for the coefficients C_i^a since we can rewrite this equation as

$$\langle i | \hat{f} | a \rangle + \langle \Phi_i^a | \hat{H} - E | \Phi_i^a \rangle C_i^a + \sum_{bj \neq ai} \langle \Phi_i^a | \hat{H} | \Phi_j^b \rangle C_j^b + \sum_{bcjk} \langle \Phi_i^a | \hat{H} | \Phi_{jk}^{bc} \rangle C_{jk}^{bc} + \sum_{bcdjkl} \langle \Phi_i^a | \hat{H}$$

Iterative solutions

We rewrite this equation as

$$C_i^a = -(\langle \Phi_i^a | \hat{H} - E | \Phi_i^a \rangle)^{-1} \left(\langle i | \hat{f} | a \rangle + \sum_{bj \neq ai} \langle \Phi_i^a | \hat{H} | \Phi_j^b \rangle C_j^b + \sum_{bcjk} \langle \Phi_i^a | \hat{H} | \Phi_{jk}^{bc} \rangle \right)$$

Coefficients for single excitations

Since these equations are solved iteratively (that is we can start with a guess for the coefficients C_i^a), it is common to start the iteration by setting

$$C_i^a = -\frac{\langle i|\hat{f}|a\rangle}{\langle \Phi_i^a|\hat{H} - E|\Phi_i^a\rangle},$$

and the denominator can be written as

$$C_i^a = \frac{\langle i|\hat{f}|a\rangle}{\langle i|\hat{f}|i\rangle - \langle a|\hat{f}|a\rangle + \langle ai|\hat{v}|ai\rangle - E}.$$

The observant reader will however see that we need an equation for C_{jk}^{bc} and C_{jkl}^{bcd} as well. To find equations for these coefficients we need then to continue our multiplications from the left with the various Φ_H^P terms.

Double excitations

For C_{jk}^{bc} we need then

$$\langle \Phi_{ij}^{ab} | \hat{H} - E | \Phi_0 \rangle + \sum_{kc} \langle \Phi_{ij}^{ab} | \hat{H} - E | \Phi_k^c \rangle C_k^c + \sum_{cdkl} \langle \Phi_{ij}^{ab} | \hat{H} - E | \Phi_{kl}^{cd} \rangle C_{kl}^{cd} +$$

$$\sum_{cdeklm} \langle \Phi_{ij}^{ab} | \hat{H} - E | \Phi_{klm}^{cde} \rangle C_{klm}^{cde} + \sum_{cdefklmn} \langle \Phi_{ij}^{ab} | \hat{H} - E | \Phi_{klmn}^{cdef} \rangle C_{klmn}^{cdef} = 0,$$

and we can isolate the coefficients C_{kl}^{cd} in a similar way as we did for the coefficients C_i^a .

n-coupled excitations

At the end we can rewrite our solution of the Schrödinger equation in terms of n coupled equations for the coefficients C_H^P . This is a very cumbersome way of solving the equation. However, by using this iterative scheme we can illustrate how we can compute the various terms in the wave operator or correlation operator \hat{C} . We will later identify the calculation of the various terms C_H^P as parts of different many-body approximations to full CI. In particular, we will relate this non-linear scheme with Coupled Cluster theory and many-body perturbation theory.

If we use a Hartree-Fock basis, how can one simplify the equation

$$\Delta E = \sum_{ai} \langle i | \hat{f} | a \rangle C_i^a + \sum_{abij} \langle ij | \hat{v} | ab \rangle C_{ij}^{ab}?$$

The similarity transformed Hamiltonian

There are several ways to view and understand the coupled-cluster method. A first simple view of coupled-cluster theory is that the method induces correlations into the reference state by expressing a correlated state as

$$|\Psi\rangle = e^T |\Phi_0\rangle, \quad (5)$$

Clusters of excited states

Here, T is an operator that induces correlations. We can now demand that the correlate state (5) becomes an eigenstate of the Hamiltonian H_N , i.e. $H|\Psi\rangle = E|\Psi\rangle$. This view, while correct, is not the most productive one. Instead, we left-multiply the Schrödinger equation with e^{-T} and find

$$\overline{H_N}|\Phi_0\rangle = \Delta E|\Phi_0\rangle. \quad (6)$$

Here, ΔE is the correlation energy, and the total energy is $E = \Delta E + E_0^{\text{ref}}$.

Similarity transformation

The similarity-transformed Hamiltonian is defined as

$$\overline{H_N} \equiv e^{-T} H_N e^T. \quad (7)$$

A more productive view on coupled-cluster theory thus emerges:
This method seeks a similarity transformation such that the
uncorrelated reference state becomes an exact eigenstate of the
similarity-transformed Hamiltonian (7).

What T leads to Hermitian $\overline{H_N}$?

What are the conditions on T such that $\overline{H_N}$ is Hermitian?

Answer

For a Hermitian $\overline{H_N}$, we need a unitary e^T , i.e. an anti-Hermitian T with $T = -T^\dagger$

As we will see below, coupled-cluster theory employs a non-Hermitian Hamiltonian.

Understanding (non-unitary) similarity transformations

Show that $\overline{H_N}$ has the same eigenvalues as H_N for arbitrary T .
What is the spectral decomposition of a non-Hermitian $\overline{H_N}$?

Answer

Let $H_N|E\rangle = E|E\rangle$. Thus

$$\begin{aligned} H_N e^T e^{-T} |E\rangle &= E|E\rangle, \\ \left(e^{-T} H_N e^T \right) e^{-T} |E\rangle &= E e^{-T} |E\rangle, \\ \overline{H_N} e^{-T} |E\rangle &= E e^{-T} |E\rangle. \end{aligned}$$

Thus, if $|E\rangle$ is an eigenstate of H_N with eigenvalue E , then $e^{-T}|E\rangle$ is eigenstate of $\overline{H_N}$ with the same eigenvalue.

Non-hermitian operator

A non-Hermitian $\overline{H_N}$ has eigenvalues E_α corresponding to left $\langle L_\alpha |$ and right $|R_\alpha \rangle$ eigenstates. Thus

$$\overline{H_N} = \sum_{\alpha} |R_\alpha\rangle E_\alpha \langle L_\alpha| \quad (8)$$

with bi-orthonormal $\langle L_\alpha | R_\beta \rangle = \delta_\alpha^\beta$.

More formalism

To make progress, we have to specify the cluster operator T . In coupled cluster theory, this operator is

$$T \equiv \sum_{ia} t_i^a a_a^\dagger a_i + \frac{1}{4} \sum_{ijab} t_{ij}^{ab} a_a^\dagger a_b^\dagger a_j a_i + \dots + \frac{1}{(N!)^2} \sum_{i_1 \dots i_N a_1 \dots a_N} t_{i_1 \dots i_N}^{a_1 \dots a_N} a_{a_1}^\dagger \dots a_{a_N}^\dagger \quad (9)$$

Particle-hole excitations

Thus, the operator (9) induces particle-hole (p-h) excitations with respect to the reference. In general, T generates up to $A_p - A_h$ excitations, and the unknown parameters are the cluster amplitudes t_i^a , t_{ij}^{ab} , ..., $t_{i_1, \dots, i_A}^{a_1, \dots, a_A}$.

Full cluster operator

Thus, the coupled-cluster method with the full cluster operator (9) is exponentially expensive, just as FCI. To make progress, we need to make an approximation by truncating the operator. Here, we will use the CCSD (coupled clusters singles doubles) approximation, where

$$T \equiv \sum_{ia} t_i^a a_a^\dagger a_i + \frac{1}{4} \sum_{ijab} t_{ij}^{ab} a_a^\dagger a_b^\dagger a_j a_i. \quad (10)$$

Unknown amplitudes

We need to determine the unknown cluster amplitudes that enter in CCSD. Let

$$|\Phi_i^a\rangle = a_a^\dagger a_i |\Phi_0\rangle, \quad (11)$$

$$|\Phi_{ij}^{ab}\rangle = a_a^\dagger a_b^\dagger a_j a_i |\Phi_0\rangle \quad (12)$$

be 1p-1h and 2p-2h excitations of the reference. Computing matrix elements of the Schrödinger Equation (6) yields

$$\langle \Phi_0 | \overline{H_N} | \Phi_0 \rangle = E_c, \quad (13)$$

$$\langle \Phi_i^a | \overline{H_N} | \Phi_0 \rangle = 0, \quad (14)$$

$$\langle \Phi_{ij}^{ab} | \overline{H_N} | \Phi_0 \rangle = 0. \quad (15)$$

Correlation energy

The first equation states that the coupled-cluster correlation energy is an expectation value of the similarity-transformed Hamiltonian.

The second and third equations state that the similarity-transformed Hamiltonian exhibits no 1p-1h and no 2p-2h excitations. These equations have to be solved to find the unknown amplitudes t_i^a and t_{ij}^{ab} . Then one can use these amplitudes and compute the correlation energy from the first line of Eq. (13).

We note that in the CCSD approximation the reference state is not an exact eigenstates. Rather, it is decoupled from simple states but \bar{H} still connects this state to 3p-3h, and 4p-4h states etc.

Good reference state

At this point, it is important to recall that we assumed starting from a “good” reference state. In such a case, we might reasonably expect that the inclusion of 1p-1h and 2p-2h excitations could result in an accurate approximation. Indeed, empirically one finds that CCSD accounts for about 90% of the corelation energy, i.e. of the difference between the exact energy and the Hartree-Fock energy. The inclusion of triples (3p-3h excitations) typically yields 99% of the correlation energy.

We see that the coupled-cluster method in its CCSD approximation yields a similarity-transformed Hamiltonian that is of a two-body structure with respect to a non-trivial vacuum. When viewed in this light, the coupled-cluster method “transforms” an A -body problem (in CCSD) into a two-body problem, albeit with respect to a nontrivial vacuum.

Above we argued that a similarity transformation preserves all eigenvalues. Nevertheless, the CCD correlation energy is not the exact correlation energy. Explain!

The CCD approximation does not make $|\Phi_0\rangle$ an exact eigenstate of $\overline{H_N}$; it is only an eigenstate when the similarity-transformed Hamiltonian is truncated to at most 2p-2h states. The full $\overline{H_N}$, with $T = T_2$, would involve six-body terms (do you understand this?), and this full Hamiltonian would reproduce the exact correlation energy. Thus CCD is a similarity transformation plus a truncation, which decouples the ground state only from 2p-2h states.

Computing the similarity-transformed Hamiltonian

The solution of the CCSD equations, i.e. the second and third line of Eq. (13), and the computation of the correlation energy requires us to compute matrix elements of the similarity-transformed Hamiltonian (7). This can be done with the Baker-Campbell-Hausdorff expansion

$$\overline{H_N} = e^{-T} H_N e^T \quad (16)$$

$$= H_N + [H_N, T] + \frac{1}{2!} [[H_N, T], T] + \frac{1}{3!} [[[H_N, T], T], T] + \dots \quad (17)$$

The cluster operator

We now come to a key element of coupled-cluster theory: the cluster operator (9) consists of sums of terms that consist of particle creation and hole annihilation operators (but no particle annihilation or hole creation operators). Thus, all terms that enter T commute with each other. This means that the commutators in the Baker-Campbell-Hausdorff expansion can only be non-zero because each T must connect to H_N (but no T with another T). Thus, the expansion is finite.

Non-hermitian Hamiltonian

We see that the (disadvantage of having a) non-Hermitian Hamiltonian $\overline{H_N}$ leads to the advantage that the Baker-Campbell-Hausdorff expansion is finite, thus leading to the possibility to compute $\overline{H_N}$ exactly.

Similarity transformed Hamiltonian

We write the similarity-transformed Hamiltonian as

$$\overline{H_N} = \sum_{pq} \overline{H}_q^p a_q^\dagger a_p + \frac{1}{4} \sum_{pqrs} \overline{H}_{rs}^{pq} a_p^\dagger a_q^\dagger a_s a_r + \dots \quad (18)$$

with

$$\overline{H}_q^p \equiv \langle p | \overline{H_N} | q \rangle, \quad (19)$$

$$\overline{H}_{rs}^{pq} \equiv \langle pq | \overline{H_N} | rs \rangle. \quad (20)$$

If we truncate the operator T at the level of singles and doubles, the CCSD Eqs. (13) for the amplitudes can be written as $\overline{H}_i^a = 0$ and $\overline{H}_{ij}^{ab} = 0$.

CCD Approximation

In what follows, we will consider the coupled cluster doubles (CCD) approximation. This approximation is valid in cases where the system cannot exhibit any particle-hole excitations (such as nuclear matter when formulated on a momentum-space grid) or for the pairing model (as the pairing interactions only excites pairs of particles). In this case $t_i^a = 0$ for all i, a , and $\bar{H}_i^a = 0$. The CCD approximation is also of some sort of leading order approximation in the Hartree-Fock basis (as the Hartree-Fock Hamiltonian exhibits no particle-hole excitations).

Deriving the CCD equations

Let us consider the matrix element \bar{H}_{ij}^{ab} . Clearly, it consists of all diagrams (i.e. all combinations of T_2 , and a single F or V that have two incoming hole lines and two outgoing particle lines. Write down all these diagrams.

We start systematically and consider all combinations of F and V diagrams with 0, 1, and 2 cluster amplitudes T_2 .

The algebraic expression

$$\begin{aligned}\overline{H}_{ij}^{ab} &= \langle ab | V | ij \rangle + P(ab) \sum_c f_c^b t_{ij}^{ac} - P(ij) \sum_k f_j^k t_{ik}^{ab} \\ &+ \frac{1}{2} \sum_{cd} \langle ab | V | cd \rangle t_{ij}^{cd} + \frac{1}{2} \sum_{kl} \langle kl | V | ij \rangle t_{kl}^{ab} + P(ab)P(ij) \sum_{kc} \langle kb | V | cj \rangle \\ &+ \frac{1}{2} P(ij)P(ab) \sum_{kcl} \langle kl | V | cd \rangle t_{ik}^{ac} t_{lj}^{db} + \frac{1}{2} P(ij) \sum_{kcl} \langle kl | V | cd \rangle t_{ik}^{cd} t_{lj}^{ab} \\ &+ \frac{1}{2} P(ab) \sum_{kcl} \langle kl | V | cd \rangle t_{kl}^{ac} t_{ij}^{db} + \frac{1}{4} \sum_{kcl} \langle kl | V | cd \rangle t_{ij}^{cd} t_{kl}^{ab}.\end{aligned}$$

CCD for the pairing Hamiltonian

You learned about the pairing Hamiltonian earlier in the second midterm. Convince yourself that this Hamiltonian does not induce any 1p-1h excitations. Let us solve the CCD equations for this problem. This consists of the following steps

1. Write a function that compute the potential, i.e. it returns a four-indexed array (or tensor). We need $\langle ab|V|cd\rangle$, $\langle ij|V|kl\rangle$, and $\langle ab|V|ij\rangle$. Why is there no $\langle ab|V|id\rangle$ or $\langle ai|V|jb\rangle$?
2. Write a function that computes the Fock matrix, i.e. a two-indexed array. We only need f_a^b and f_i^j . Why?
3. Initialize the cluster amplitudes according and solve the equations for the amplitudes iteratively.

Solving the CCD equations for the pairing problem

The Hamiltonian is

$$H = \delta \sum_{p=1}^{\Omega} (p-1) \left(a_{p+}^\dagger a_{p+} + a_{p-}^\dagger a_{p-} \right) - \frac{g}{2} \sum_{p,q=1}^{\Omega} a_{p+}^\dagger a_{p-}^\dagger a_{q-} a_{q+}. \quad (21)$$

Exact diagonalization

```
#!/usr/bin/python
from sympy import *
from pylab import *
import matplotlib.pyplot as plt

ga = linspace(-1,1,20)
e1 = []

for g_val in ga:
    H1 = matrix([[2-g_val, -g_val/2., -g_val/2., -g_val/2., -g_val/2.,
                  [-g_val/2., 4-g_val, -g_val/2., -g_val/2., -g_val/2.,
                   [-g_val/2., -g_val/2., 6-g_val, 0, -g_val/2.,
                    [-g_val/2., -g_val/2., 0, 6-g_val, -g_val/2.,
                     [-g_val/2., 0, -g_val/2., -g_val/2., 8-g_val,
                      [0, -g_val/2., -g_val/2., -g_val/2., -g_val/2., -g_val/2.]]]]])
    u1, v1 = linalg.eig(H1)
    e1.append(min(u1)))
exact = e1 - (2-ga)
print(exact)

plt.axis([-1,1,-0.5,0.05])
plt.xlabel(r'Interaction strength, $g$', fontsize=16)
plt.ylabel(r'Correlation energy', fontsize=16)
exact = plt.plot(ga, exact, 'b-*', linewidth = 2.0, label = 'Exact')
plt.legend()
plt.savefig('pairing.pdf', format='pdf')
plt.show()
```

Python code

```
## Coupled clusters in CCD approximation
## Implemented for the pairing model of Lecture Notes in Physics 936,
import numpy as np
```

```
def init_pairing_v(g,pnum,hnum):
    """
    returns potential matrices of the pairing model in three relevant
    param g: strength of the pairing interaction, as in Eq. (8.42)
    param pnum: number of particle states
    param hnum: number of hole states

    return v_pppp, v_pphh, v_hhhh: np.array(pnum,pnum,pnum,pnum),
           np.array(pnum,pnum,hnum,hnum),
           np.array(hnum,hnum,hnum,hnum),
           The interaction as a 4-indexed tens
    """
    v_pppp=np.zeros((pnum,pnum,pnum,pnum))
    v_pphh=np.zeros((pnum,pnum,hnum,hnum))
    v_hhhh=np.zeros((hnum,hnum,hnum,hnum))

    gval=-0.5*g
    for a in range(0,pnum,2):
        for b in range(0,pnum,2):
            v_pppp[a,a+1,b,b+1]=gval
            v_pppp[a+1,a,b,b+1]=-gval
            v_pppp[a,a+1,b+1,b]=-gval
            v_pppp[a+1,a,b+1,b]=gval
```