#### Nuclear Shell Model

Morten Hjorth-Jensen, National Superconducting Cyclotron Laboratory and Department of Physics and Astronomy, Michigan State University, East Lansing, MI 48824, USA & Department of Physics, University of Oslo, Oslo, Norway

Spring 2015

#### Nuclear shell model

The topics covered in this and next week's lectures can be found in Alex Brown's lecture notes, chapters 5, 18, 21-22. See also Suhonen's chapter's 2 and 5.

### Slater determinants as basis states, Repetition

The simplest possible choice for many-body wavefunctions are **product** wavefunctions. That is

$$\Psi(x_1, x_2, x_3, \ldots, x_A) \approx \phi_1(x_1)\phi_2(x_2)\phi_3(x_3)\ldots$$

because we are really only good at thinking about one particle at a time. Such product wavefunctions, without correlations, are easy to work with; for example, if the single-particle states  $\phi_i(x)$  are orthonormal, then the product wavefunctions are easy to orthonormalize.

Similarly, computing matrix elements of operators are relatively easy, because the integrals factorize.

The price we pay is the lack of correlations, which we must build up by using many, many product wavefunctions. (Thus we have a trade-off: compact representation of correlations but difficult integrals versus easy integrals but many states required.)

#### Slater determinants as basis states, repetition

Because we have fermions, we are required to have antisymmetric wavefunctions, e.g.

$$\Psi(x_1, x_2, x_3, \ldots, x_A) = -\Psi(x_2, x_1, x_3, \ldots, x_A)$$

etc. This is accomplished formally by using the determinantal formalism

$$\Psi(x_1, x_2, \dots, x_A) = \frac{1}{\sqrt{A!}} \det \begin{vmatrix} \phi_1(x_1) & \phi_1(x_2) & \dots & \phi_1(x_A) \\ \phi_2(x_1) & \phi_2(x_2) & \dots & \phi_2(x_A) \\ \vdots & & & & \\ \phi_A(x_1) & \phi_A(x_2) & \dots & \phi_A(x_A) \end{vmatrix}$$

Product wavefunction + antisymmetry = Slater determinant.

#### Slater determinants as basis states

$$\Psi(x_1, x_2, \dots, x_A) = \frac{1}{\sqrt{N!}} \det \begin{vmatrix} \phi_1(x_1) & \phi_1(x_2) & \dots & \phi_1(x_A) \\ \phi_2(x_1) & \phi_2(x_2) & \dots & \phi_2(x_A) \\ \vdots & & & & \\ \phi_A(x_1) & \phi_A(x_2) & \dots & \phi_A(x_A) \end{vmatrix}$$

Properties of the determinant (interchange of any two rows or any two columns yields a change in sign; thus no two rows and no two columns can be the same) lead to the Pauli principle:

- No two particles can be at the same place (two columns the same); and
- No two particles can be in the same state (two rows the same).

#### Slater determinants as basis states

As a practical matter, however, Slater determinants beyond N=4 quickly become unwieldy. Thus we turn to the **occupation** representation or second quantization to simplify calculations. The occupation representation, using fermion creation and annihilation operators, is compact and efficient. It is also abstract and, at first encounter, not easy to internalize. It is inspired by other operator formalism, such as the ladder operators for the harmonic oscillator or for angular momentum, but unlike those cases, the operators do not have coordinate space representations.

Instead, one can think of fermion creation/annihilation operators as a game of symbols that compactly reproduces what one would do, albeit clumsily, with full coordinate-space Slater determinants.

#### Quick repetition of the occupation representation

We start with a set of orthonormal single-particle states  $\{\phi_i(x)\}$ . (Note: this requirement, and others, can be relaxed, but leads to a more involved formalism.) **Any** orthonormal set will do.

To each single-particle state  $\phi_i(x)$  we associate a creation operator  $\hat{a}_i^{\dagger}$  and an annihilation operator  $\hat{a}_i$ .

When acting on the vacuum state  $|0\rangle$ , the creation operator  $\hat{a}_i^{\dagger}$  causes a particle to occupy the single-particle state  $\phi_i(x)$ :

$$\phi_i(x) \rightarrow \hat{a}_i^{\dagger} |0\rangle$$

### Quick repetition of the occupation representation

But with multiple creation operators we can occupy multiple states:

$$\phi_i(x)\phi_j(x')\phi_k(x'') \rightarrow \hat{a}_i^{\dagger}\hat{a}_i^{\dagger}\hat{a}_k^{\dagger}|0\rangle.$$

Now we impose antisymmetry, by having the fermion operators satisfy **anticommutation relations**:

$$\hat{a}_i^{\dagger}\hat{a}_j^{\dagger}+\hat{a}_j^{\dagger}\hat{a}_i^{\dagger}=[\hat{a}_i^{\dagger},\hat{a}_j^{\dagger}]_+=\{\hat{a}_i^{\dagger},\hat{a}_j^{\dagger}\}=0$$

so that

$$\hat{a}_i^{\dagger}\hat{a}_j^{\dagger}=-\hat{a}_j^{\dagger}\hat{a}_i^{\dagger}$$

## Quick repetition of the occupation representation

Because of this property, automatically  $\hat{a}_i^{\dagger}\hat{a}_i^{\dagger}=0$ , enforcing the Pauli exclusion principle. Thus when writing a Slater determinant using creation operators,

$$\hat{a}_i^{\dagger}\hat{a}_j^{\dagger}\hat{a}_k^{\dagger}\dots|0\rangle$$

each index  $i, j, k, \ldots$  must be unique.

We have defined the ansatz for the ground state as

$$|\Phi_0
angle = \left(\prod_{i\leq F}\hat{a}_i^\dagger
ight)|0
angle,$$

where the index i defines different single-particle states up to the Fermi level. We have assumed that we have N fermions. A given one-particle-one-hole (1p1h) state can be written as

$$|\Phi_{i}^{a}\rangle=\hat{a}_{a}^{\dagger}\hat{a}_{i}|\Phi_{0}\rangle,$$

while a 2p2h state can be written as

$$|\Phi_{ii}^{ab}\rangle = \hat{a}_{a}^{\dagger}\hat{a}_{b}^{\dagger}\hat{a}_{j}\hat{a}_{i}|\Phi_{0}\rangle,$$

and a general NpNh state as

$$|\Phi_{iik...}^{abc...}\rangle = \hat{a}_{a}^{\dagger}\hat{a}_{b}^{\dagger}\hat{a}_{c}^{\dagger}\dots\hat{a}_{k}\hat{a}_{j}\hat{a}_{i}|\Phi_{0}\rangle.$$

We can then expand our exact state function for the ground state as

$$|\Psi_0\rangle = C_0|\Phi_0\rangle + \sum_{ai} C_i^a|\Phi_i^a\rangle + \sum_{abii} C_{ij}^{ab}|\Phi_{ij}^{ab}\rangle + \cdots = (C_0 + \hat{C})|\Phi_0\rangle,$$

where we have introduced the so-called correlation operator

$$\hat{C} = \sum_{ai} C_i^a \hat{a}_a^\dagger \hat{a}_i + \sum_{abij} C_{ij}^{ab} \hat{a}_a^\dagger \hat{a}_b^\dagger \hat{a}_j \hat{a}_i + \dots$$

Since the normalization of  $\Psi_0$  is at our disposal and since  $C_0$  is by hypothesis non-zero, we may arbitrarily set  $C_0=1$  with corresponding proportional changes in all other coefficients. Using this so-called intermediate normalization we have

$$\langle \Psi_0 | \Phi_0 \rangle = \langle \Phi_0 | \Phi_0 \rangle = 1,$$

resulting in

We rewrite

$$|\Psi_0\rangle = \mathit{C}_0 |\Phi_0\rangle + \sum_{ai} \mathit{C}_i^a |\Phi_i^a\rangle + \sum_{abii} \mathit{C}_{ij}^{ab} |\Phi_{ij}^{ab}\rangle + \ldots,$$

in a more compact form as

$$|\Psi_0\rangle = \sum_{PH} C_H^P \Phi_H^P = \left(\sum_{PH} C_H^P \hat{A}_H^P\right) |\Phi_0\rangle,$$

where H stands for  $0, 1, \ldots, n$  hole states and P for  $0, 1, \ldots, n$  particle states. Our requirement of unit normalization gives

$$\langle \Psi_0 | \Phi_0 \rangle = \sum_{PH} | \mathit{C}_H^P |^2 = 1,$$

and the energy can be written as

$$E = \langle \Psi_0 | \hat{H} | \Phi_0 \rangle = \sum_{l} C_H^{*P} \langle \Phi_H^P | \hat{H} | \Phi_{H'}^{P'} \rangle C_{H'}^{P'}.$$

Normally

$$E = \langle \Psi_0 | \hat{H} | \Phi_0 \rangle = \sum_{PP'HH'} C_H^{*P} \langle \Phi_H^P | \hat{H} | \Phi_{H'}^{P'} \rangle C_{H'}^{P'},$$

is solved by diagonalization setting up the Hamiltonian matrix defined by the basis of all possible Slater determinants. A diagonalization is equivalent to finding the variational minimum of

$$\langle \Psi_0 | \hat{H} | \Phi_0 \rangle = \lambda \langle \Psi_0 | \Phi_0 \rangle$$

where  $\lambda$  is a variational multiplier to be identified with the energy of the system. The minimization process results in

$$\delta \left[ \langle \Psi_0 | \hat{H} | \Phi_0 \rangle - \lambda \langle \Psi_0 | \Phi_0 \rangle \right] =$$

$$\sum_{Pll''}\Big\{\delta[C_{H}^{*P}]\langle\Phi_{H}^{P}|\hat{H}|\Phi_{H'}^{P'}\rangle C_{H'}^{P'}+C_{H}^{*P}\langle\Phi_{H}^{P}|\hat{H}|\Phi_{H'}^{P'}\rangle\delta[C_{H'}^{P'}]-\lambda(\delta[C_{H}^{*P}]C_{H'}^{P'}+$$

An alternative way to derive the last equation is to start from

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

and if this equation is successively projected against all  $\Phi_H^P$  in the expansion of  $\Psi$ , then the last equation on the previous slide results. 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  $\hat{\mathcal{C}}$ .

For reasons to come (links with Coupled-Cluster theory and Many-Body perturbation theory), we will rewrite Eq. (  $\ref{eq:coupled}$ ) as a set of coupled non-linear equations in terms of the unknown coefficients  $C_H^P$ .

To see this, we look at the contributions arising from

$$\langle \Phi_H^P | = \langle \Phi_0 |$$

in Eq. (??), that is we multiply with  $\langle \Phi_0 |$  from the left in

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

If we assume that we have a two-body operator at most, Slater's rule gives then an equation for the correlation energy in terms of  $C_i^a$  and  $C_{ii}^{ab}$  only. We get then

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

Inserting the various matrix elements we can rewrite the previous equation as

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

This equation determines the correlation energy but not the coefficients C. We need more equations. Our next step is to set up

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

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

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

We rewrite this equation as

$$C_i^a = -(\langle \Phi_i^a | \hat{H} | \Phi_i^a \rangle^{-1}$$

$$\times \left( \langle i|\hat{f}|a\rangle + \sum_{bj\neq ai} \langle \Phi^a_i|\hat{H}|\Phi^b_j\rangle C^b_j + \sum_{bcjk} \langle \Phi^a_i|\hat{H}|\Phi^{bc}_{jk}\rangle C^{bc}_{jk} + \sum_{bcdjkl} \langle \Phi^a_i|\hat{H}|\Phi^{bcd}_{jkl}\rangle \right)$$
 Since these equations are solved iteratively ( that is we can start with a guess for the coefficients  $C^a_i$ ), it is common to start the iteration by setting

 $C_i^a = -rac{\langle i|f|a
angle}{\langle \Phi_i^a|\hat{H}|\Phi_i^a
angle},$ 

and the denominator can be written as

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

The observant reader will however see that we need an equation for

For  $C_{ik}^{bc}$  we need then

cdkl

$$\langle \Phi^{ab}_{ij}|\hat{H} - E|\Phi_0\rangle + \sum_{kc} \langle \Phi^{ab}_{ij}|\hat{H} - E|\Phi^c_k\rangle C^c_k +$$

$$\sum \langle \Phi_{ij}^{ab} | \hat{H} - E | \Phi_{kl}^{cd} \rangle C_{kl}^{cd} + \sum \langle \Phi_{ij}^{ab} | \hat{H} - E | \Phi_{klm}^{cde} \rangle C_{klm}^{cde} + \sum \langle \Phi_{ij}^{ab} | \hat{H} - E | \Phi_{klm}^{cde} \rangle C_{klm}^{cde} + \sum \langle \Phi_{ij}^{ab} | \hat{H} - E | \Phi_{klm}^{cde} \rangle C_{klm}^{cde} + \sum \langle \Phi_{ij}^{ab} | \hat{H} - E | \Phi_{klm}^{cde} \rangle C_{klm}^{cde} + \sum \langle \Phi_{ij}^{ab} | \hat{H} - E | \Phi_{klm}^{cde} \rangle C_{klm}^{cde} + \sum \langle \Phi_{ij}^{ab} | \hat{H} - E | \Phi_{klm}^{cde} \rangle C_{klm}^{cde} + \sum \langle \Phi_{ij}^{ab} | \hat{H} - E | \Phi_{klm}^{cde} \rangle C_{klm}^{cde} + \sum \langle \Phi_{ij}^{ab} | \hat{H} - E | \Phi_{klm}^{cde} \rangle C_{klm}^{cde} + \sum \langle \Phi_{ij}^{ab} | \hat{H} - E | \Phi_{klm}^{cde} \rangle C_{klm}^{cde} + \sum \langle \Phi_{ij}^{ab} | \hat{H} - E | \Phi_{klm}^{cde} \rangle C_{klm}^{cde} + \sum \langle \Phi_{ij}^{ab} | \hat{H} - E | \Phi_{klm}^{cde} \rangle C_{klm}^{cde} + \sum \langle \Phi_{ij}^{ab} | \hat{H} - E | \Phi_{klm}^{cde} \rangle C_{klm}^{cde} + \sum \langle \Phi_{ij}^{ab} | \hat{H} - E | \Phi_{klm}^{cde} \rangle C_{klm}^{cde} + \sum \langle \Phi_{ij}^{ab} | \hat{H} - E | \Phi_{klm}^{cde} \rangle C_{klm}^{cde} + \sum \langle \Phi_{ij}^{ab} | \hat{H} - E | \Phi_{klm}^{cde} \rangle C_{klm}^{cde} + \sum \langle \Phi_{ij}^{ab} | \hat{H} - E | \Phi_{klm}^{cde} \rangle C_{klm}^{cde} + \sum \langle \Phi_{ij}^{ab} | \hat{H} - E | \Phi_{klm}^{cde} \rangle C_{klm}^{cde} + \sum \langle \Phi_{ij}^{ab} | \hat{H} - E | \Phi_{klm}^{cde} \rangle C_{klm}^{cde} + \sum \langle \Phi_{ij}^{ab} | \hat{H} - E | \Phi_{klm}^{cde} \rangle C_{klm}^{cde} + \sum \langle \Phi_{ij}^{ab} | \hat{H} - E | \Phi_{klm}^{cde} \rangle C_{klm}^{cde} + \sum \langle \Phi_{ij}^{ab} | \hat{H} - E | \Phi_{klm}^{cde} \rangle C_{klm}^{cde} + \sum \langle \Phi_{ij}^{ab} | \hat{H} - E | \Phi_{klm}^{cde} \rangle C_{klm}^{cde} + \sum \langle \Phi_{ij}^{ab} | \hat{H} - E | \Phi_{klm}^{cde} \rangle C_{klm}^{cde} + \sum \langle \Phi_{ij}^{ab} | \hat{H} - E | \Phi_{klm}^{cde} \rangle C_{klm}^{cde} + \sum \langle \Phi_{ij}^{ab} | \hat{H} - E | \Phi_{klm}^{cde} \rangle C_{klm}^{cde} + \sum \langle \Phi_{ij}^{ab} | \hat{H} - E | \Phi_{klm}^{cde} \rangle C_{klm}^{cde} + \sum \langle \Phi_{ij}^{ab} | \hat{H} - E | \Phi_{klm}^{cde} \rangle C_{klm}^{cde} + \sum \langle \Phi_{ij}^{ab} | \hat{H} - E | \Phi_{klm}^{cde} \rangle C_{klm}^{cde} + \sum \langle \Phi_{ij}^{ab} | \hat{H} - E | \Phi_{klm}^{cde} \rangle C_{klm}^{cde} + \sum \langle \Phi_{ij}^{ab} | \hat{H} - E | \Phi_{klm}^{cde} \rangle C_{klm}^{cde} + \sum \langle \Phi_{ij}^{ab} | \hat{H} - E | \Phi_{klm}^{cde} \rangle C_{klm}^{cde} + \sum \langle \Phi_{ij}^{ab} | \hat{H} - E | \Phi_{klm}^{cde} \rangle C_{klm}^{cde} + \sum \langle \Phi_{ij}^{ab} | \hat{H} - E | \Phi_{klm$$

and we can isolate the coefficients  $C_{\nu I}^{cd}$  in a similar way as we did for the coefficients  $C_i^a$ . 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, www can relate this non-linear scheme with Coupled Cluster theory and many-body perturbation theory. These theories will not be discussed in this course

If we use a Hartree-Fock basis, we simplify this equation

$$\Delta E = \sum_{\mathit{ai}} \langle i | \hat{f} | \mathit{a} \rangle \mathit{C}^{\mathit{a}}_{i} + \sum_{\mathit{abij}} \langle i j | \hat{v} | \mathit{ab} \rangle \mathit{C}^{\mathit{ab}}_{ij}.$$

What about

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

and

$$\langle \Phi^{ab}_{ij}|\hat{H}-E|\Phi_0\rangle + \sum_{kc} \langle \Phi^{ab}_{ij}|\hat{H}-E|\Phi^c_k\rangle C^c_k + \sum_{cdkl} \langle \Phi^{ab}_{ij}|\hat{H}-E|\Phi^{cd}_{kl}\rangle C^{cd}_{kl} +$$

$$\sum_{cdeklm} \langle \Phi^{ab}_{ij} | \hat{H} - E | \Phi^{cde}_{klm} \rangle C^{cde}_{klm} + \sum_{cdefklmn} \langle \Phi^{ab}_{ij} | \hat{H} - E | \Phi^{cdef}_{klmn} \rangle C^{cdef}_{klmn} = 0?$$

In project 1 the plan is to construct a working code that constructs the many-body Hamiltonian matrix in a basis of Slater determinants and to find the low-lying eigenenergies. This is referred to as the configuration-interaction method or shell-model diagonalization (or the interacting shell model).

The first step in such codes—and in your project—is to construct the many-body basis.

While the formalism is independent of the choice of basis, the **effectiveness** of a calculation will certainly be basis dependent. Furthermore there are common conventions useful to know.

First, the single-particle basis has angular momentum as a good quantum number. You can imagine the single-particle wavefunctions being generated by a one-body Hamiltonian, for example a harmonic oscillator. Modifications include harmonic oscillator plus spin-orbit splitting, or self-consistent mean-field potentials, or the Woods-Saxon potential which mocks up the self-consistent mean-field.

For nuclei, the harmonic oscillator, modified by spin-orbit splitting, provides a useful language for describing single-particle states. Each single-particle state is labeled by the following quantum numbers:

- Orbital angular momentum /
- ▶ Intrinsic spin s = 1/2 for protons and neutrons
- Angular momentum  $j = l \pm 1/2$
- $\triangleright$  z-component  $i_z$  (or m)
- Some labeling of the radial wavefunction, typically *n* the

In this format one labels states by  $n(I)_j$ , with (I) replaced by a letter: s for I=0, p for I=1, d for I=2, f for I=3, and thenceforth alphabetical.

In practice the single-particle space has to be severely truncated. This truncation is typically based upon the single-particle energies, which is the effective energy from a mean-field potential. Sometimes we freeze the core and only consider a valence space. For example, one may assume a frozen  $^4\text{He}$  core, with two protons and two neutrons in the  $0s_{1/2}$  shell, and then only allow active particles in the  $0p_{1/2}$  and  $0p_{3/2}$  orbits.

Another example is a frozen  $^{16}{\rm O}$  core, with eight protons and eight neutrons filling the  $0s_{1/2}$ ,  $0p_{1/2}$  and  $0p_{3/2}$  orbits, with valence particles in the  $0d_{5/2}$ ,  $1s_{1/2}$  and  $0d_{3/2}$  orbits.

Sometimes we refer to nuclei by the valence space where their last nucleons go. So, for example, we call  $^{12}$ C a p-shell nucleus, while  $^{26}$ Al is an sd-shell nucleus and  $^{56}$ Fe is a pf-shell nucleus.

There are different kinds of truncations.

- ► For example, one can start with 'filled' orbits (almost always the lowest), and then allow one, two, three... particles excited out of those filled orbits. These are called 1p-1h, 2p-2h, 3p-3h excitations.
- Alternately, one can state a maximal orbit and allow all possible configurations with particles occupying states up to that maximum. This is called *full configuration*.
- Finally, for particular use in nuclear physics, there is the *energy* truncation, also called the  $N\hbar\Omega$  or  $N_{max}$  truncation.

Here one works in a harmonic oscillator basis, with each major oscillator shell assigned a principal quantum number  $N=0,1,2,3,\ldots$ 

The  $N\hbar\Omega$  or  $N_{max}$  truncation: Any configuration is given an noninteracting energy, which is the sum of the single-particle harmonic oscillator energies. (Thus this ignores spin-orbit splitting.)

Excited state are labeled relative to the lowest configuration by the number of harmonic oscillator quanta.

This truncation is useful because: if one includes *all* configuration up to some  $N_{max}$ , and has a translationally invariant interaction, then the intrinsic motion and the center-of-mass motion factor. In other words, we can know exactly the center-of-mass wavefunction.

In almost all cases, the many-body Hamiltonian is rotationally invariant. This means it commutes with the operators  $\hat{J}^2$ ,  $\hat{J}_z$  and so eigenstates will have good J, M. Furthermore, the eigenenergies do not depend upon the orientation M.

Therefore we can choose to construct a many-body basis which has fixed M; this is called an M-scheme basis.

Alternately, one can construct a many-body basis which has fixed J, or a J-scheme basis.

The Hamiltonian matrix will have smaller dimensions (a factor of 10 or more) in the *J*-scheme than in the *M*-scheme. On the other hand, as we'll show in the next slide, the *M*-scheme is very easy to construct with Slater determinants, while the *J*-scheme basis states, and thus the matrix elements, are more complicated, almost always being linear combinations of *M*-scheme states. *J*-scheme bases are important and useful, but we'll focus on the simpler *M*-scheme.

The quantum number m is additive (because the underlying group is Abelian): if a Slater determinant  $\hat{a}_i^{\dagger}\hat{a}_j^{\dagger}\hat{a}_k^{\dagger}\dots|0\rangle$  is built from single-particle states all with good m, then the total

$$M = m_i + m_i + m_k + \dots$$

This is *not* true of J, because the angular momentum group SU(2) is not Abelian.

The upshot is that

- ▶ It is easy to construct a Slater determinant with good total *M*;
- ▶ It is trivial to calculate *M* for each Slater determinant;
- ► So it is easy to construct an *M*-scheme basis with fixed total *M*.

Note that the individual M-scheme basis states will not, in general, have good total J. Because the Hamiltonian is rotationally invariant, however, the eigenstates will have good J. (The situation is muddied when one has states of different J that are nonetheless

t	Example: two $j = 1/2$ orbits						
	Index	n	1	j	$m_j$		
_	1			1 /2	1/2		

muex	11	1	J	Шj
1	0	0	1/2	-1/2
2	0	0	1/2	1/2
3	1	0	1/2	-1/2
4	1	0	1/2	1/2

Note: the order is arbitrary. There are  $\begin{pmatrix} 4 \\ 2 \end{pmatrix} = 6$  two-particle

states, which we list with the total M:

Occupied	IVI	
1,2	0	
1,3	-1	
1,4	0	
2,3	0	
2,4	1	
3,4	0	
and 1 each	with	Λ/

and 1 each with  $M = \pm 1$ .

Example: consider using only single particle states from the  $0\,d_{5/2}$  space. They have the following quantum numbers

Index	n	1	j	$m_j$
1	0	2	5/2	-5/2
2	0	2	5/2	-3/2
3	0	2	5/2	-1/2
4	0	2	5/2	1/2
5	0	2	5/2	3/2
6	0	2	5/2	5/2
		/ _	$\overline{}$	

There are 
$$\begin{pmatrix} 6 \\ 2 \end{pmatrix} = 15$$
 two-particle states, which we list with the

total M:

Occupied	М	Occupied	Μ	Occupied	Μ
1,2	-4	2,3	-2	3,5	1
1,3	-3	2,4	-1	3,6	2
1,4	-2	2,5	0	4,5	2
1,5	-1	2,6	1	4,6	3
1,6	0	3,4	0	5,6	4

#### Project

The basic goal of the project is for you to build your own configuration-interaction shell-model code. The code will be fairly basic; it will assume that we have a single species of particles, e.g. only neutrons, and you could, if you wish to, read in uncoupled two-body matrix elements. Furthermore the pieces of the code will not be the most efficient. Nonetheless it will be usable; most importantly, you will gain a good idea of what goes into a many-body shell-model code.

The first step is to construct the M-scheme basis of Slater determinants. Here M-scheme means the total  $J_z$  of the many-body states is fixed.

The steps could be:

- Read in a user-supplied file of single-particle states (examples can be given) or just code these internally;
- Ask for the total M of the system and the number of particles N;
- Construct all the N-particle states with given M. You will validate the code by comparing both the number of states and specific states.

The format of a possible input file could be

Index	n	1	2 <i>j</i>	$2m_j$
1	1	0	1	-1
2	1	0	1	1
3	0	2	3	-3
4	0	2	3	-1
5	0	2	3	1
6	0	2	3	3
7	0	2	5	-5
8	0	2	5	-3
9	0	2	5	-1
10	0	2	5	1
11	0	2	5	3
12	0	2	5	5
Thic ro	nroc	onto	- +h-	1

This represents the  $1s_{1/2}0d_{3/2}0d_{5/2}$  valence space, or just the sd-space. There are twelve single-particle states, labeled by an overall index, and which have associated quantum numbers the number of radial nodes, the orbital angular momentum J, and the angular momentum j and third component  $j_z$ . To keep everything as integers, we could store  $2 \times j$  and  $2 \times j_z$ .

To read in the single-particle states you need to:

- ▶ Open the file
  - Read the number of single-particle states (in the above example, 12); allocate memory; all you need is a single array storing  $2 \times j_z$  for each state, labeled by the index.
- ▶ Read in the quantum numbers and store  $2 \times j_z$  (and anything else you happen to want).

The next step is to read in the number of particles N and the fixed total M (or, actually,  $2 \times M$ ). For this project we assume only a single species of particles, say neutrons, although this can be relaxed. **Note**: Although it is often a good idea to try to write a more general code, given the short time alloted we would suggest you keep your ambition in check, at least in the initial phases of the project.

You should probably write an error trap to make sure N and M are congruent; if N is even, then  $2 \times M$  should be even, and if N is odd then  $2 \times M$  should be odd.

The final step is to generate the set of N-particle Slater determinants with fixed M. The Slater determinants will be stored in occupation representation. Although in many codes this representation is done compactly in bit notation with ones and zeros, but for greater transparency and simplicity we will list the occupied single particle states.

Hence we can store the Slater determinant basis states as sd(i,j), that is an array of dimension  $N_{SD}$ , the number of Slater determinants, by N, the number of occupied state. So if for the 7th Slater determinant the 2nd, 3rd, and 9th single-particle states are occupied, then sd(7,1)=2, sd(7,2)=3, and sd(7,3)=9.

We can construct an occupation representation of Slater determinants by the *odometer* method. Consider  $N_{sp} = 12$  and N = 4. Start with the first 4 states occupied, that is:

▶ 
$$sd(1,:) = 1,2,3,4$$
 (also written as  $|1,2,3,4\rangle$ )

Now increase the last occupancy recursively:

$$ightharpoonup sd(2,:) = 1,2,3,5$$

$$sd(3,:) = 1,2,3,6$$

$$\blacktriangleright$$
  $sd(4,:) = 1,2,3,7$ 

$$\rightarrow$$
  $sd(9,:) = 1, 2, 3, 12$ 

Then start over with

**.** . . .

$$\rightarrow$$
  $sd(10,:) = 1,2,4,5$ 

and again increase the rightmost digit

$$sd(11,:) = 1,2,4,6$$

When we restrict ourselves to an M-scheme basis, we could choose two paths. The first is simplest (and simplest is often best, at least in the first draft of a code): generate all possible Slater determinants, and then extract from this initial list a list of those Slater determinants with a given M. (You will need to write a short function or routine that computes M for any given occupation.) Alternately, and not too difficult, is to run the odometer routine twice: each time, as as a Slater determinant is calculated, compute M, but do not store the Slater determinants except the current one. You can then count up the number of Slater determinants with a chosen M. Then allocated storage for the Slater determinants, and run the odometer algorithm again, this time storing Slater determinants with the desired M (this can be done with a simple logical flag).

Some example solutions: Let's begin with a simple case, the  $0d_{5/2}$  space containing six single-particle states

Index	n	1	j	$m_j$	
1	0	2	5/2	-5/2	
2	0	2	5/2	-3/2	
3	0	2	5/2	-1/2	
4	0	2	5/2	1/2	
5	0	2	5/2	3/2	
6	0	2	5/2	5/2	
For two particles, there are a total of 15 states, which we list here					

with the total M:

▶ 
$$|1,2\rangle$$
,  $M = -4$ ,  $|1,3\rangle$ ,  $M = -3$   
▶  $|1,4\rangle$ ,  $M = -2$ ,  $|1,5\rangle$ ,  $M = -1$ 

► 
$$|1,5\rangle$$
,  $M = 0$ ,  $|2,3\rangle$ ,  $M = -2$ 

$$|1,5\rangle$$
,  $M = 0$ ,  $|2,3\rangle$ ,  $M = -2$   
 $|2,4\rangle$ ,  $M = -1$ ,  $|2,5\rangle$ ,  $M = 0$ 

$$|2,6\rangle, M=1, |3,4\rangle, M=0$$

$$|3,5\rangle$$
,  $M=1$ ,  $|3,6\rangle$ ,  $M=2$ 

$$|45\rangle M = 2 |46\rangle M = 3$$

You should try by hand to show that in this same single-particle space, that for N=3 there are 3 states with M=1/2 and for N=4 there are also only 3 states with M=0.

To test your code, confirm the above.

Also, for the sd-space given above, for N=2 there are 14 states with M=0, for N=3 there are 37 states with M=1/2, for N=4 there are 81 states with M=0.

For our project, we will only consider the pairing model. A simple space is the  $(1/2)^2$  space with four single-particle states

=		_		_
4	1	0	1/2	1/2
3	1	0	1/2	-1/2
2	0	0	1/2	1/2
1	0	0	1/2	-1/2
Index	n	1	S	$m_s$

For N=2 there are 4 states with M=0; show this by hand and confirm your code reproduces it.

Another, slightly more challenging space is the  $(1/2)^4$  space, that is, with eight single-particle states we have

Index	n	1	5	m <sub>s</sub>
1	0	0	1/2	-1/2
2	0	0	1/2	1/2
3	1	0	1/2	-1/2
4	1	0	1/2	1/2
5	2	0	1/2	-1/2
6	2	0	1/2	1/2
7	3	0	1/2	-1/2
8	3	0	1/2	1/2
_ ^/	_	. 1		1.

For N=2 there are 16 states with M=0; for N=3 there are 24 states with M=1/2, and for N=4 there are 36 states with M=0.

In the shell-model context we can interpret this as 4  $s_{1/2}$  levels, with  $m=\pm 1/2$ , we can also think of these are simple four pairs,  $\pm k, k=1,2,3,4$ . Later on we will assign single-particle energies, depending on the radial quantum number n, that is,  $\epsilon_k=|k|\delta$  so that they are equally spaced.

For application in the pairing model we can go further and consider only states with no "broken pairs," that is, if +k is filled (or m=+1/2, so is -k (m=-1/2). If you want, you can write your code to accept only these, and obtain the following six states:

- ► |1, 2, 3, 4⟩,
- |1,2,5,6⟩,
- **▶** |1, 2, 7, 8⟩,
- **▶** |3, 4, 5, 6⟩,
- |3, 4, 7, 8⟩,
- **▶** |5, 6, 7, 8⟩

#### Hints for coding

- Write small modules (routines/functions); avoid big functions that do everything. (But not too small.)
- ▶ Write lots of error traps, even for things that are 'obvious.'
- Document as you go along. For each function write a header that includes:
  - 1. Main purpose of function
  - 2. names and brief explanation of input variables, if any
  - 3. names and brief explanation of output variables, if any
  - 4. functions called by this function
  - 5. called by which functions

#### Hints for coding

- ► When debugging, print out intermediate values. It's almost impossible to debug a code by looking at it—the code will almost always win a 'staring contest.'
- ▶ Validate code with SIMPLE CASES. Validate early and often.

The number one mistake is using a too complex a system to test. For example, if you are computing particles in a potential in a box, try removing the potential—you should get particles in a box. And start with one particle, then two, then three... Don't start with eight particles.

Our recommended occupation representation, e.g.  $|1,2,4,8\rangle$ , is easy to code, but numerically inefficient when one has hundreds of millions of Slater determinants.

In state-of-the-art shell-model codes, one generally uses bit representation, i.e.  $|1101000100...\rangle$  where one stores the Slater determinant as a single (or a small number of) integer.

This is much more compact, but more intricate to code with considerable more overhead. There exist bit-manipulation functions. This is left as a challenge for those of you who would like to study this topic further for the final project to be presented for the oral examination.

We consider a space with  $2\Omega$  single-particle states, with each state labeled by  $k=1,2,3,\Omega$  and  $m=\pm 1/2$ . The convention is that the state with k>0 has m=+1/2 while -k has m=-1/2. The Hamiltonian we consider is

$$\hat{H} = -G\hat{P}_{+}\hat{P}_{-},$$

where

$$\hat{P}_+ = \sum_{k>0} \hat{a}_k^{\dagger} \hat{a}_{-k}^{\dagger}.$$

and  $\hat{P}_- = (\hat{P}_+)^\dagger$ .

This problem can be solved using what is called the quasi-spin formalism to obtain the exact results. Thereafter we will try again using the explicit Slater determinant formalism.

One can show (and this is part of the project) that

$$\left[\hat{P}_+,\hat{P}_-
ight] = \sum_{k>0} \left(\hat{a}_k^\dagger \hat{a}_k + \hat{a}_{-k}^\dagger \hat{a}_{-k} - 1\right) = \hat{N} - \Omega.$$

Now define

$$\hat{P}_z = \frac{1}{2}(\hat{N} - \Omega).$$

Finally you can show

$$\left[\hat{P}_{z},\hat{P}_{\pm}\right]=\pm\hat{P}_{\pm}.$$

This means the operators  $\hat{P}_{\pm}$ ,  $\hat{P}_z$  form a so-called SU(2) algebra, and we can use all our insights about angular momentum, even though there is no actual angular momentum involved (this is similar to project 1).

So we rewrite the Hamiltonian to make this explicit:

$$\hat{\Omega} = \hat{\Omega} \hat{\Omega} \hat{\Omega} = \hat{\Omega} \hat{\Omega} \hat{\Omega} \hat{\Omega} \hat{\Omega} \hat{\Omega} \hat{\Omega}$$

Because of the SU(2) algebra, we know that the eigenvalues of  $\hat{P}^2$  must be of the form p(p+1), with p either integer or half-integer, and the eigenvalues of  $\hat{P}_z$  are  $m_p$  with  $p \geq |m_p|$ , with  $m_p$  also integer or half-integer.

But because  $\hat{P}_z = (1/2)(\hat{N} - \Omega)$ , we know that for N particles the value  $m_p = (N - \Omega)/2$ . Furthermore, the values of  $m_p$  range from  $-\Omega/2$  (for N=0) to  $+\Omega/2$  (for  $N=2\Omega$ , with all states filled).

We deduce the maximal  $p=\Omega/2$  and for a given n the values range of p range from  $|N-\Omega|/2$  to  $\Omega/2$  in steps of 1 (for an even number of particles)

Following Racah we introduce the notation  $p=(\Omega-v)/2$  where  $v=0,2,4,...,\Omega-|N-\Omega|$  With this it is easy to deduce that the eigenvalues of the pairing Hamiltonian are

$$-G(N-v)(2\Omega+2-N-v)/4$$

This also works for N odd, with  $v = 1, 3, 5, \ldots$ 

Let's take a specific example:  $\Omega=3$  so there are 6 single-particle states, and N=3, with  $\nu=1,3$ . Therefore there are two distinct eigenvalues,

$$E = -2G, 0$$

Now let's work this out explicitly. The single particle degrees of freedom are defined as

maex	K	m
1	1	-1/2
2	-1	1/2
3	2	-1/2
4	-2	1/2
5	3	-1/2
6	-3	1/2
		/ - \

There are  $\begin{pmatrix} 6 \\ 3 \end{pmatrix} = 20$  three-particle states, but there are 9 states

with M = +1/2, namely

 $|1,2,3\rangle, |1,2,5\rangle, |1,4,6\rangle, |2,3,4\rangle, |2,3,6\rangle, |2,4,5\rangle, |2,5,6\rangle, |3,4,6\rangle, |4,5,6\rangle, |$ 

In this basis, the operator

$$\hat{P}_{+}=\hat{a}_{1}^{\dagger}\hat{a}_{2}^{\dagger}+\hat{a}_{3}^{\dagger}\hat{a}_{4}^{\dagger}+\hat{a}_{5}^{\dagger}\hat{a}_{6}^{\dagger}$$

From this we can determine that

$$\hat{P}_{-}|1,4,6\rangle = \hat{P}_{-}|2,3,6\rangle = \hat{P}_{-}|2,4,5\rangle = 0$$

so those states all have eigenvalue 0.

Now for further example,

$$\hat{P}_{-}|1,2,3\rangle=|3\rangle$$

SO

$$\hat{P}_{+}\hat{P}_{-}|1,2,3\rangle=|1,2,3\rangle+|3,4,3\rangle+|5,6,3\rangle$$

The second term vanishes because state 3 is occupied twice, and reordering the last term we get

$$\hat{P}_{+}\hat{P}_{-}|1,2,3\rangle = |1,2,3\rangle + |3,5,6\rangle$$

without picking up a phase.

Continuing in this fashion, with the previous ordering of the many-body states (  $|1,2,3\rangle,|1,2,5\rangle,|1,4,6\rangle,|2,3,4\rangle,|2,3,6\rangle,|2,4,5\rangle,|2,5,6\rangle,|3,4,6\rangle,|4,5,6$  the Hamiltonian matrix of this system is

This is useful for our project. One can by hand confirm that there are 3 eigenvalues -2G and 6 with value zero.

Another example Using the  $(1/2)^4$  single-particle space, resulting in eight single-particle states

Index	n	1	S	$m_s$
1	0	0	1/2	-1/2
2	0	0	1/2	1/2
3	1	0	1/2	-1/2
4	1	0	1/2	1/2
5	2	0	1/2	-1/2
6	2	0	1/2	1/2
7	3	0	1/2	-1/2
8	3	0	1/2	1/2
a a d + b a	+	ماناه	را م م ا	. 1

and then taking only 4-particle, M=0 states that have no 'broken pairs', there are six basis Slater determinants:

- ► |1,2,3,4⟩,
- **▶** |1, 2, 5, 6⟩,
- ▶ |1,2,7,8⟩,
- |3, 4, 5, 6⟩,
- ▶ |3, 4, 7, 8⟩,

Now we take the following Hamiltonian

$$\hat{H} = \sum_{n} n \delta \hat{N}_{n} - G \hat{P}^{\dagger} \hat{P}$$

where

$$\hat{N}_n = \hat{a}_{n,m=+1/2}^{\dagger} \hat{a}_{n,m=+1/2} + \hat{a}_{n,m=-1/2}^{\dagger} \hat{a}_{n,m=-1/2}$$

and

$$\hat{\mathcal{P}}^\dagger = \sum_n \hat{a}_{n,m=+1/2}^\dagger \hat{a}_{n,m=-1/2}^\dagger$$

We can write down the  $6\times 6$  Hamiltonian in the basis from the prior slide:

$$H = \left( egin{array}{ccccccc} 2\delta - 2G & -G & -G & -G & -G & 0 \ -G & 4\delta - 2G & -G & -G & -G & -G \ -G & -G & 6\delta - 2G & 0 & -G & -G \ -G & -G & 0 & 6\delta - 2G & -G & -G \end{array} 
ight)$$