Shell-model Hamiltonians from Density Functional Theory

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The density functional theory of nuclear structure provides a many-particle wave function that is useful for static properties, but an extension of the theory is necessary to describe correlation effects or other dynamic properties. Here we propose a procedure to extend the theory by mapping the properties of the self-consistent mean-field Hamiltonian onto an effective shell-model Hamiltonian with two-body interactions. In this initial study, we consider the sd-shell nuclei \(^{20}\text{Ne},^{24}\text{Mg},^{28}\text{Si}, \) and \(^{36}\text{Ar}.\) Our first application is in the framework of the USD shell-model Hamiltonian, using its mean-field approximation to construct an effective Hamiltonian and partially recover correlation effects. We find that more than half of the correlation energy is due to the quadrupole interaction. We then follow a similar procedure but using the SLy4 Skyrme energy functional as our starting point and truncating the space to the spherical \(sd\) shell. The constructed shell-model Hamiltonian is found to satisfy minimal consistency requirements to reproduce the properties of the mean-field solution. The quadrupolar correlation energies computed with the mapped Hamiltonian are reasonable compared with those computed by other methods.

PACS numbers: 21.60.-n, 21.60.Jz, 21.60.Cs, 21.10.Dr

I. INTRODUCTION

Quantitative theory of nuclear structure has followed two different paths. One path is a self-consistent mean-field theory (SCMF)\textsuperscript{1}, also called density functional theory,\textsuperscript{1} and the other is the configuration-interaction shell model (CISM) approach.\textsuperscript{2} The former requires only a few parameters and has broad applicability across the table of nuclides, while the latter is specific to regions of the nuclides and as usually formulated requires many parameters to specify the Hamiltonian. However, the SCMF is of limited accuracy for calculating binding energies, and the treatment of excitations requires extensions of the theory. In contrast, the CISM gives a rather accurate and almost\textsuperscript{2} complete description of the low-energy spectroscopy. Brown and Richter\textsuperscript{4} proposed to use the SCMF to determine the single-particle Hamiltonian of the CISM so as to gain advantages of both approaches. In this work we go farther, using the SCMF to also determine parts of the two-body interaction Hamiltonian.

Our work has several motivations. One is to develop a well-justified and accurate theory of nuclear masses. SCMF is systematic with a strong theoretical foundation, but it may have reached its limits of accuracy at a level of an MeV rms residual error over the table of known masses. Some effects of correlations are missing from the theory, and indeed better accuracy can be achieved when phenomenological terms are included to take them into account.\textsuperscript{3} A systematic calculation of those correlation energies would be possible if we could find a way to map the SCMF on a shell-model Hamiltonian.\textsuperscript{3}

Another motivation is to bring more precision to shell-model calculations of one-body observables such as quadrupole moments and transition intensities. For example, the SCMF is well adapted to computing quadrupole matrix elements because of the large single-particle space that can be treated. The CISM requires severe truncation of the space, and a systematic mapping procedure would determine the effective operators to be used in the smaller truncated shell-model space.

Finally, such a mapping would be helpful to construct a global theory of nuclear level densities. While the overall behavior of level densities can be derived from the independent particle shell model, there are significant interaction effects. The shell model Monte Carlo (SMMC) method has been efficient for calculating level densities in large shell-model spaces,\textsuperscript{3,4,5} but it requires as input a parameterized one- and two-body shell-model Hamiltonian. It should be mentioned that the SMMC requires for its efficiency interactions that have a specific sign.\textsuperscript{4}

\textsuperscript{1} This terminology is used because of the similarity to the density functional theory of many-electron systems.\textsuperscript{2}
\textsuperscript{2} The qualification is due to the common occurrence of intruder states.

\textsuperscript{3} The SCMF energy functionals in use are fitted to the binding energies, and in principle they should be refitted if the correlation energies is added as a separate contribution. In practice, the refit does not change the parameters significantly so the original functional may be used in the determination of the Fock-space Hamiltonian.

\textsuperscript{4} Interactions containing small terms having the bad sign can be treated as well with the method of Ref. \textsuperscript{6}.
These conditions are satisfied for the Hamiltonian we will construct here.

Here we propose a mapping procedure and apply it to sd shell nuclei. This region of the table of nuclides is a paradigm for the CISM. For example, the USD Hamiltonian \[10\] defined in the sd shell space reproduces very well binding energies, excitation energies, and transition rates between states. For the actual SCMF that one might wish to map, we will use one of the Skyrme family of energy functionals. The corresponding SCMF equations can be solved with computer codes that are publicly available \[11, 12\]; in particular we shall use in this work the Brussels-Paris code \[12, 13\], together with the SLY4 parameterization of the Skyrme energy functional \[14\]. This functional is fit to a number of empirical properties of nuclei and with a slight readjustment gives a 1.7 MeV rms fit to masses of \[\sim 600\] even-even nuclei \[15, 16\].

The outline of this paper is as follows. In Section II we discuss the main idea and procedure for constructing the SCMF to CISM map. Our starting point is the SCMF, which minimizes an energy functional in the presence of an external field \(E\) as the total energy \(E = \sum \phi K \cdot \phi\). Our task is to use this information to construct a Hamiltonian of the form \(\hat{H}\). The minimal requirement of such an effective Hamiltonian is that it reproduces ground-state SCMF one-body observables sufficiently well within a mean-field approximation (in the truncated space). We use this requirement as a guide to construct the effective Hamiltonian.

In the applications discussed in this work, we treat nuclei whose SCMF ground state is deformed. This will guide to construct the effective Hamiltonian.

Finally, in Section VII we discuss possible future developments of the theory introduced here.

## II. METHODOLOGY

Our goal is to choose a suitable truncated shell-model space and construct within it a spherical shell-model Hamiltonian containing one- and two-body terms. We write this CISM Hamiltonian in the form \(\hat{H} = \sum \epsilon_{\alpha j} a_{\alpha j}^\dagger a_{\alpha j} - \frac{1}{2} \sum \kappa g_{\kappa} \hat{A}_\kappa \cdot \hat{A}_\kappa\). The first step is the construction of a spherical single-particle basis. Due to the rotational invariance of the energy functional, a spherically symmetric density matrix \(\rho\) at a given iteration step leads to a one-body Hamiltonian that is also spherically symmetric. For a closed sub-shell

\[\hat{A}_\kappa M = \sum_{\alpha j \alpha j'} f_{\alpha j \alpha j'}^\kappa \sum_{m m'} (j' m' K_\kappa M | j m) a_{\alpha j m}^\dagger a_{\alpha j' m'}, \]

and the scalar product \(H\) in Eq. \(\mathbf{1}\) is \(\hat{A}_\kappa \cdot \hat{A}_\kappa = \sum (\kappa - M) A_{\kappa M} A_{\kappa - M}\). Eq. \(\mathbf{1}\) provides the most natural form to which the SMMC method can be applied. With enough terms in the \(\kappa\) sum, it is sufficiently general to treat any two-body interaction that is rotationally invariant and time-reversal symmetric. The maximal number of terms \(\kappa\) for any given rank \(K_\kappa\) of the multipole operators is the number of orbital pairs whose angular momenta \(j, j'\) can be coupled to \(K_\kappa\) by angular momentum and parity selection rules. We also note that \(f_{\alpha j \alpha j'}^\kappa\) are proportional to the reduced matrix elements of \(\hat{A}_\kappa\), i.e., \(f_{\alpha j \alpha j'}^\kappa = (\alpha j | \hat{A}_\kappa | \alpha j')/\sqrt{2j + 1}\). In the development below we do not need to distinguish orbitals with different quantum numbers \(\alpha\), and we will drop that index.

Our starting point is the SCMF, which minimizes an energy functional for a configuration of single-particle orbitals \(\phi_k\) (i.e., a Slater determinant) or the BCS generalization thereof. Besides the ground-state minimum, one can obtain constrained solutions by minimizing the energy functional in the presence of an external field \(\lambda Q\), where \(Q\) is the mass quadrupole operator defined below and \(\Lambda\) is its strength. The SCMF delivers a set of orbitals \(\delta_{\kappa}\) and corresponding single-particle energies \(\epsilon_{\kappa}\), as well as the total energy \(E^\Lambda\). Our task is to use this information to construct a Hamiltonian of the form \(\mathbf{1}\). The minimal requirement of such an effective Hamiltonian is that it reproduces ground-state SCMF one-body observables sufficiently well within a mean-field approximation (in the truncated space). We use this requirement as a guide to construct the effective Hamiltonian.

In the applications discussed in this work, we treat nuclei whose SCMF ground state is deformed. This will simplify the mapping to some extent because we do not have to impose a constraining field.

The first step is the construction of a spherical single-particle basis. Due to the rotational invariance of the energy functional, a spherically symmetric density matrix at a given iteration step leads to a one-body Hamiltonian.
nucleus (e.g., $^{28}$Si) the density matrix in the next iteration step will remain spherically symmetric. Thus, for a closed sub-shell nucleus, if we start from a rotationally invariant single-particle density and iterate, we will eventually converge to a spherical solution. For an open shell nucleus, we have to choose which $n$ orbitals of the valence partially occupied $j$-shell are filled. Such a choice will break the spherical symmetry of the density and lead to a deformed solution. To overcome this problem, one may use the uniform filling approximation in which the last $n$ particles are distributed uniformly over the corresponding $2j + 1$ orbitals (i.e., we assume occupations of $v^2_{jm} = n/(2j + 1)$). As long as this approximation is used in each HF iteration, we will converge to a spherical solution. In practice, a pairing interaction or BCS-like matrix equation $U_{nm}$ is important to choose the truncated single-particle space such that the truncated ground-state density matrix is given by $\rho_{nm}$ satisfies some minimal properties required for a density matrix $\rho$ that is the solution of the CISM Hamiltonian in a mean-field approximation. One such requirement is that the number of particles be correct; this is checked with the formula $\text{tr} \rho = N$.\(^8\)

This property is satisfied exactly in the full space with $N$ being the total number of particles, and the truncated $\rho$ should satisfy this condition approximately with $N$ being the number of valence particles. Another requirement is that the density matrix represent a single configuration (i.e., a Slater determinant). This can be expressed in the matrix equation $\rho \rho = \rho \ . \quad (6)$

Equivalently, the requirement is that the eigenvalues of the density matrix be zero or one (with exactly $N$ eigenvalues equal to one). We will examine the eigenvalue spectrum of our mapped density matrices to see how well this requirement is satisfied.

The mean-field Hamiltonian in the SCMF ground-state basis is the single-particle matrix $h_{nm}$ with elements $h_{nm,kk'} = \epsilon'_{k} \delta_{kk'}$. Its matrix representation in the truncated spherical basis is given by $h = U h_{nm} U$.

Let us expand $h$ in multipoles, writing $h = \sum_{K} h^{(K)}$.

For example, the expectation value of one-body observables in the truncated space by using this truncated density $P \rho_{nm} P$. For example, the expectation value of the mass quadrupole operator $\hat{q} = \sum_{jm,j'm'} q_{jm,j'm'} \delta_{jm} \delta_{j'm'}$ in the truncated space may be computed as

$$\langle \hat{q} \rangle = \text{tr}(q P \rho_{nm} P) = \sum_{jm,j'm'} q_{jm,j'm'} \rho_{jm,j'm'} \ . \quad (4)$$

where $q_{jm,j'm'} = \langle jm | \hat{Q} | j'm' \rangle$ are the matrix elements of the mass quadrupole operator $\hat{Q} = \sum_{i}(2z_i^2 - x_i^2 - y_i^2)$ in the single-particle space.\(^8\)

Since the transformation $U$ from the deformed basis into the truncated spherical basis may not be unitary, it is important to choose the truncated single-particle space such that the truncated ground-state density matrix $P \rho_{nm} P$ satisfies some minimal properties required for a density matrix that is the solution of the CISM Hamiltonian in a mean-field approximation. One such requirement is that the number of particles be correct; this is checked with the formula $\text{tr} \rho = N$.\(^5\)

We now assume that the SCMF ground state is axially symmetric, so that only the $M = 0$ components of $h^{(K)}$ are non-vanishing. We shall also restrict ourselves to even-even nuclei, which by time-reversal invariance permits only even multipoles $K$. The lowest multipole is the monopole $h^{(0)}$ from which we obtain the spherical single-particle energies $\epsilon_j^{(0)}$.

$$h^{(0)}_{jm,j'm'} = \epsilon_j^{(0)} \delta_{jj'} \delta_{mm'} \ . \quad (9)$$

These single-particle energies are given by $\epsilon_j^{(0)} = \sum_m h_{jm,jm}/(2j + 1)$.

\(^8\) In the following we will omit the projector $P$ in the definition of the truncated $\rho$ and it should be understood from the context whether $\rho$ denotes the complete or projected density matrix.
When the ground state is deformed, the \( K = 2 \) part of the single-particle Hamiltonian will be substantial and can be used to determine an effective interaction in that quadrupolar channel. Defining the second-quantized one-body tensor operators (see Eq. (2))

\[
\hat{h}^{(K)}_M = \sum_{j,j'} \frac{\langle j | \hat{h}^{(K)} | j' \rangle}{\sqrt{2j + 1}} \sum_{mm'} (j' mm' K M jm) \hat{a}^{+}_{jm} \hat{a}_{j'm'},
\]

we consider the following effective CISM Hamiltonian

\[
\hat{H} = \hat{h}^{(0)} - \frac{1}{2} g \hat{h}^{(2)},
\]

(10)

The Hamiltonian \( \hat{h} \) has the form \( \hat{h} \equiv \hat{h} \) with a single multipole term \( K \). Assuming axial symmetry, the Hartree Hamiltonian of \( \hat{H} \) is given by

\[
\hat{h}^{(0)} = \sum_{M} (-)^M \hat{h}^{(2)}_M \sum_{bd} \hat{h}^{(2)}_{M;bd} \rho_{bd},
\]

(11)

where \( \hat{h}^{(2)}_M \equiv \langle a | \hat{h}^{(2)} | c \rangle \). If the mean-field solution \( \rho_{ac} \) in \( \hat{H} \) possesses axial symmetry, only the \( M = 0 \) component of \( \hat{h}^{(2)}_M \) contributes to the direct term, while all values of \( M \) can contribute to the sum in the exchange term.

We have used the Hartree approximation to motivate our choice of the effective interaction \( \hat{H} \). However, in practice we include the Fock term in the Hamiltonian to determine the density matrix and deformation energy of the system. Using \( \{jm\} \) to denote the spherical orbitals, the Hartree-Fock Hamiltonian of \( \hat{H} \) is given by

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Another important quantity is the total energy of the system. The absolute SCMF energies need not be reproduced by the mapped Hamiltonian, but the energy difference between the mean-field minimum \( E_{mf} \) and the spherical state \( E_{sph} \) should come out the same. We call this the deformation energy \( E_{def} \),

\[
E_{def} = E_{mf} - E_{sph}.
\]

(13)

A deformation energy can also be calculated by solving the CISM Hamiltonian \( \hat{H} \) in the mean-field approximation. The above procedure to determine \( g \) does not guarantee that the deformation energy of the original SCMF theory is reproduced by the mapped CISM Hamiltonian. We fine tune the value of \( g \) so as to match the deformation energies of the SCMF theory and of the CISM Hamiltonian. As shown below, the deformation energy of the CISM Hamiltonian is very sensitive to \( g \), and a relatively small change from its value determined by \( \hat{H} \) will suffice to reproduce the correct deformation energy. On the other hand, the quadrupole moment and the ground density matrix (in the truncated space) are less sensitive to \( g \) and are still expected to match approximately.

\[9\] The spherical state is always a stationary solution of the mean-field equations (in the uniform filling approximation), but it may not be lowest state satisfying the constraint \( \langle Q \rangle = 0 \). In that case, one has to find it in some other way. In this section we have found the spherical solution by starting from a spherical density...
The spherical state has a HF energy of $E_{\text{sp}} = -126.03$ MeV and zero quadrupole moment.

Find $g = 0.0232$ MeV/$b^4$ and the deformed state has a quadrupole moment of $-18.2$ $b^2$, about 10% lower than the actual minimum of the full USD Hamiltonian.

\[
\begin{array}{cccccc}
\text{Nucleus} & \text{Interaction} & g & E_{\text{ph}} & E_{\text{mf}} & E_{\text{def}} & \langle Q \rangle & E_{\text{corr}} \\
\hline
^{20}\text{Ne} & \text{USD} & \hat{h}^{(2)} & 0.051 & -25.46 & -36.38 & 10.92 & 15.4 & 4.1 \\
^{24}\text{Mg} & \text{USD} & \hat{h}^{(2)} & 0.0261 & -68.20 & -80.17 & 11.97 & 18.0 & 6.9 \\
^{28}\text{Si} & \text{USD} & \hat{h}^{(2)} & 0.0232 & -126.03 & -129.61 & 3.58 & -20.3 & 6.3 \\
^{36}\text{Ar} & \text{USD} & \hat{h}^{(2)} & 0.062 & -222.75 & -226.56 & 3.82 & -13.5 & 4.0 \\
\end{array}
\]

We have made similar calculations for the other deformed $N = Z$ even-even nuclei in the $sd$ shell. This excludes $^{32}$S, which is found to be spherical in the HF approximation. The results are summarized in Table I. We determine $g$ by matching the deformation energy of the USD Hamiltonian, and in all cases we find the quadrupole moment of the CISM Hamiltonian to be within $\sim 3\% - 10\%$ of the quadrupole moment of the USD Hamiltonian (where both moments are calculated in the HF approximation).

In Fig. 2 we show the occupations $\langle \hat{n}_j \rangle = \sum_m \langle \hat{n}_{jm} \rangle$ of the spherical orbitals versus the spherical single-particle energy $\epsilon_j^{(0)}$ in the deformed 4N $sd$-shell nuclei. The occupation numbers for the effective Hamiltonians are comparable to the occupations for the USD interaction.

An important use of the effective CISM Hamiltonian will be to calculate the correlation energy $E_{\text{corr}}$, defined as the difference between the mean-field energy $E_{\text{mf}}$ and the CISM ground-state energy $E_{gs}$,

\[
E_{\text{corr}} = E_{\text{mf}} - E_{gs}.
\]

The quadrupolar correlation energies extracted from the effective CISM Hamiltonians are plotted versus mass number $A$ in Fig. 2 (solid squares) and compared with the full USD correlation energies (open circles). Note that the quadrupole-quadrupole interaction is responsible for more than one half of the total correlation energy in all cases but $^{36}$Ar.

**IV. MAPPING THE SCMF**

The SCMF of interest are based on local or nearly local energy functionals and use large bases to describe the single-particle orbitals. This introduces technical problems in determining the spherical orbitals as well as conceptual problems associated with the truncation of the space. This section is devoted to these issues. Some of
FIG. 2: Occupation numbers \( \langle n_j \rangle \) of the spherical orbitals in the deformed mean-field solution as a function of the spherical single-particle energy \( \varepsilon_j^{(0)} \) in four \( sd \)-shell nuclei \((^{20}\text{Ne}, ^{24}\text{Mg}, ^{28}\text{Si} \text{ and } ^{36}\text{Ar})\). Results for the effective Hamiltonian \( I^{11} \) (solid squares) are compared with results for the USD interaction (open circles). Also shown are results for the effective interaction \( I^{12} \) discussed in Section \( II \) (solid triangles).

FIG. 3: Correlation energies \( E_{\text{corr}} \) versus mass number \( A \) in deformed \( N=Z \) \( sd \) shell nuclei. The symbols are as in Fig. 2. Quadrupolar correlations are responsible for more than 50% of the full correlation energies (with the exception of \( ^{36}\text{Ar} \)).

The details are specific to the \( \text{ev8} \) SCMF code \( I^{12} \) that we use for our numerical computations.

Since the effective CISM Hamiltonian \( \hat{H} \) must be expressed in a spherical shell basis, we need to find the spherical SCMF solution to construct the basis. We will denote the single-particle wave functions of this spherical solution by \( \phi_{\alpha jm} \) and truncate the space according to the needs of the CISM. Whether or not a solution is spherical can be easily determined by the \( 2j + 1 \) degeneracy in the single-particle energies. The \( \langle \hat{Q}\rangle = 0 \) HF ground state may or may not be spherical. Usually one includes a pairing field in the SCMF, which stabilizes a spherical solution for \( \langle \hat{Q}\rangle = 0 \). Our starting wave functions are obtained from the archive \( I^{18} \), which were calculated including a delta-function pairing interaction with a realistic strength. We take the SCMF wave function corresponding to \( \langle \hat{Q}\rangle = 0 \) (which is spherical), and perform a few unconstrained iterations with the pairing interaction turned off. However, we include BCS-like orbital occupation factors \( \varepsilon_j = (1 - \varepsilon_j - \mu)\sqrt{(\varepsilon_j - \mu)^2 + \Delta^2}/2 \) with a fixed pairing gap \( \Delta \) (\( \mu \) is the chemical potential). This procedure is applied for several values of \( \Delta \), e.g., 2, 1 and 0.5 MeV. The resulting total spherical energies are found to have an approximate linear dependence on \( \Delta \) and we have used a linear extrapolation to estimate the value at \( \Delta = 0 \). We take this extrapolated value to be the HF energy of the spherical solution.

We use the code \( \text{ev8} \), which assumes that the SCMF solution has time-reversal symmetry. The orbitals then come in degenerate time-reversed pairs, so only half of them are considered, e.g., the positive \( z \)-signature \( I^{11} \) orbitals. For an axially symmetric solution, \( J_z = m \) is a good quantum number and the positive \( z \)-signature orbitals have \( m = 1/2, -3/2, 5/2, -7/2, \ldots \). As a practical matter, there may not always be a clean separation between different \( m \) states (belonging to a given \( j \)-shell) and even between orbitals of different \( j \)-shells, due to degeneracies and numerical imprecision. In practice, we first determine \( j \) by identifying, within the manifold of positive \( z \)-signature states, the \( j + 1/2 \) nearly degenerate states with a given parity. We then diagonalize the \( (j + 1/2) \times (j + 1/2) \) matrix representing the axial quadrupole operator \( \hat{Q}_0^{(2)} = Q \) to obtain states of good \( |m| \). Using the Wigner-Eckart theorem, we have

\[
\langle jm|\hat{Q}_0^{(2)}|jm\rangle = c_j(j||\hat{Q}_0^{(2)}||j)(3m^2 - j(j + 1))
\]

with \( c_j = 2[(2j - 1)2j(2j + 1)(2j + 3)]^{-1/2} \). Thus by arranging the eigenvalues of \( \hat{Q} \) in ascending order, we identify the orbitals with \( |m| = j, j - 1, \ldots, 1/2 \).\(^{12}\)

Each good-\( |m| \) orbital constructed above is determined up to an overall phase. However, in the spherical shell model, the relative phases of the states \( |jm\rangle \) within a given \( j \) manifold should be chosen such that the states follow the standard sign convention, i.e., \( J_z|jm\rangle = \sqrt{j(j + 1) - m(m + 1)}jm \pm 1 \). For the case with time-reversal symmetry, the transformation matrix that diago-

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\( ^{11} \) The \( z \)-signature is the quantum number \( (\pm 1) \) that corresponds to the discrete symmetry transformation \( e^{i\pi J_z(-1/2)} \).

\( ^{12} \) We note that the reduced matrix element \( \langle jj'|\hat{Q}_0^{(2)}||jj'\rangle \) is negative so that \( \langle jj'|\hat{Q}_0^{(2)}||jj'\rangle \) is negative. Indeed, in an orbital with good quantum numbers \( nlj \), \( \langle nljj'\hat{Q}_0^{(2)}\rangle nljj' = -\frac{1}{2j+1}\langle r^2\rangle_{nl} < 0 \).
nalizes $\hat{Q}_Q^{(2)}$ is real. Thus each orbital $|jm\rangle$ is determined up to an overall sign.

These signs can be determined by calculating the matrix elements of $\hat{Q}_Q^{(2)}$ and requiring their sign to be consistent with the signs determined by the Wigner-Eckart theorem

$$\langle jm|\hat{Q}_Q^{(2)}|jm'\rangle = \frac{(jm')^2\mu|jm\rangle}{\sqrt{2j+1}} (j||\hat{Q}_Q^{(2)}||j)$$

(17)

where $(j||\hat{Q}_Q^{(2)}||j) < 0$. Starting from the state $|m| = j$ and using the ladder operators $\hat{Q}_Q^{(2)}$, we can successively descend to states with $|m| = j - 1, \ldots, 1/2$. At each step we determine the correct sign of the state to be consistent with the sign of the corresponding Clebsch-Gordan coefficient in (14). We note that since only positive signature orbitals are used, the sign of $m$ alternates in the above process and therefore we need to calculate matrix elements of the type $\langle jm|\hat{Q}_Q^{(2)}|jm'\rangle$ (where $\bar{m}$ is the time-reversed state of $m$).

In practice, we calculate the matrix elements $\langle ja|\hat{Q}_Q^{(2)}|j\bar{b}\rangle$ in the original basis $|ja\rangle$ obtained with ev8 (i.e., before diagonalizing $Q_0$) and then transform both $|ja\rangle$ and $|j\bar{b}\rangle$ to obtain the desired matrix elements $\langle jm|\hat{Q}_Q^{(2)}|jm'\rangle$ (note that the same transformation matrix applies to both $|ja\rangle \rightarrow |jm\rangle$ and $|j\bar{b}\rangle \rightarrow |j\bar{m}\rangle$).

The ev8 code economizes on wave function storage by using only the amplitudes associated with the first octant, $x, y, z > 0$. The symmetry properties of the wave functions under the $x = 0, y = 0$ and $z = 0$ plane reflections should then be taken onto account in the evaluation of the quadrupole matrix elements; see the Appendix for details. In the code and its output, the orbital wave functions are represented on a lattice in coordinate space. Once we have determined the standard spherical orbitals $\phi_{jm}$ and the deformed orbitals $\phi_k$, we can calculate the overlap matrix $U_{jk, jm} = \langle kjm | jm \rangle$ by taking the scalar product of the respective orbitals on the lattice. The spherical-basis matrices of the (deformed) ground-state density matrix $\rho$ and the single-particle Hamiltonian $h$ are calculated from (15) and (17), respectively.

V. 28Si in the SCMF

We now carry out the mapping for the SCMF of $^{28}\text{Si}$ following the procedures described in Sections IV and IV. We use the SCMF code ev8, which takes as input a Skyrme parameterization of the mean-field energy and a possible pairing interaction. In this work we use the SLy4 parameter set [14]. The final energies in this work are calculated in the HF approximation, with no pairing contribution.\(^{13}\)

\(^{13}\) As discussed earlier, we may include a pairing interaction when searching for the spherical solution but the energy is extrapolated to the limit of no pairing.

The energy landscape of $^{28}\text{Si}$ calculated using ev8 with a constraining mass quadrupole field is shown in Fig. 4. Qualitatively, the landscape is very similar to that found for the USD interaction in Fig. 1. The ground state at the minimum of the curve has an oblate deformation with a value $\langle \hat{Q} \rangle = -89.2 \text{ fm}^2$. This is larger than the value found for the USD interaction, which is to be expected because the USD is defined in a restricted space. There are two other stationary points in the landscape, namely the spherical saddle point and a shallow prolate minimum at $\langle \hat{Q} \rangle \approx 30 \text{ fm}^2$. The deformation energy of the oblate solution $E_{def} = E_{mf} - E_{sph} \approx 1.71 \text{ MeV}$ is lower than the deformation energy found for the USD interaction.

A standard spherical basis $|jm\rangle$ is constructed as described in Section IV. The reduced matrix elements of the mass quadrupole operator $\hat{Q}$ in the spherical basis are then extracted from

$$\langle j||\hat{Q}||j'\rangle = \frac{5}{\sqrt{2j+1}} \sum_m (2\,0\,j'm|jm\rangle)(jm||\hat{Q}||j'm) \, .$$

(18)

It is interesting to compare the reduced matrix elements [15] to those of the simple spherical Woods-Saxon (WS) model, taking the parameters of the Woods-Saxon plus spin-orbit potential from Bohr and Mottelson [17]. Fig. 5 shows the ratio $\langle j||\hat{Q}||j\rangle_{\text{SLy4}}/\langle j||\hat{Q}||j\rangle_{\text{WS}}$ of the quadrupole reduced matrix elements in the Sly4 SCMF and in the WS model versus $\langle j||\hat{Q}||j\rangle_{\text{WS}}$. We only show results for the valence sd shell composed of the $0d_{5/2}, 1s_{1/2}$ and $0d_{3/2}$ orbitals. The solid circles correspond to diagonal elements ($j = j'$) and open triangles to off-diagonal elements. The values of this ratio vary in the range $\sim 0.95 - 1.1$ (dotted lines in Fig. 5). Thus,
quadrupole renormalization effects in shell-model calculations are more influenced by the truncation than by the particular treatment of the spherical mean field. Let us next examine those truncation effects.

A. Truncation effects and operator rescaling

As mentioned earlier, the transformation from the deformed mean-field orbitals to the spherical basis is not unitary owing to truncation effects. The severity of the truncation can be assessed in several ways. The first is the trace of the truncated single-particle density matrix, Eq. (5), which should equal the number of particles in the truncated space. Each solid circle corresponds to a spherical single-particle state of protons or neutrons. The untruncated moment of -89.2 fm^2 is shown by the arrow on the right.

FIG. 6: Quadrupole moment of the SCMF ground state of ^28Si using the SLy4 Skyrme energy functional and different truncations in the spherical basis. The horizontal axis is the maximum single-particle energy in the truncated space. Each solid circle corresponds to a spherical single-particle state of protons or neutrons. The untruncated moment of -89.2 fm^2 is shown by the arrow on the right.

describing |⟨Q⟩| as a function of the maximal single-particle energy used to define the spherical model space. With a cutoff of +15 MeV unbound orbital energy, the quadrupole moment is reduced from its full value of -89 fm^2 by about 7%. Excluding all positive energy orbitals, the moment is -69 fm^2, a reduction of ~ 25%. The truncation that excludes all orbitals above the N,Z = 20 shell closure yields ⟨Q⟩ = -54.4 fm^2. Finally, we find a slightly lower value of ⟨Q⟩ = -52.4 fm^2 when truncating to the valence sd shell. Thus, within a shell-model theory in the truncated valence sd shell, the quadrupole operator should be rescaled by 89.4/52.4 ≈ 1.71.

B. Mapped Hamiltonian

To obtain the mapped Hamiltonian, we first calculate the transformation matrix U between the deformed and spherical bases, using the output orbital wave functions from ev8. We then find the deformed single-particle density matrix Eq. (3) and single-particle Hamiltonian Eq. (7) in the spherical basis. Next, we perform the multipole decomposition of the single-particle Hamiltonian \( h = h^{(0)} + h^{(2)} \) using [3] and the Wigner-Eckart theorem (to convert the reduced matrix elements to matrix elements) to extract effective spherical shell orbital energies \( e^{(0)}_{j} \) (see Eq. (9)) and an effective quadrupole field \( h^{(2)} \). From this point on the procedure is identical to what we did for the USD Hamiltonian. We construct the effective CISM Hamiltonian [11] with a coupling constant \( g \). A crude estimate for \( g \) in the Hartree approximation is \( g \approx -1/\hbar_{0}^{2} = 0.046 \) MeV/fm^4. To determine \( g \) more precisely we match the deformation energy of the effective CISM Hamiltonian [11] to its value 1.71 MeV.
found in the SCMF theory (of SLy4). The deformation energy of the CISM Hamiltonian for a given value of $g$ is calculated from the spherical and deformed (oblate) solutions of Eq. (14) in the HF approximation. This procedure gives us $g = 0.0594$ MeV/fm$^4$.

The mapped Hamiltonian can be tested by solving it in the HF approximation and comparing with the SCMF. The deformation energy is the same by construction. We obtain a quadrupole moment of $-55$ fm$^2$, which is $\sim 5\%$ larger than the SCMF quadrupole moment (projected on the $sd$ shell). We also find the spherical occupations of the deformed HF ground state of the CISM Hamiltonian to be rather close to the spherical occupations of the deformed SCMF minimum (see Table IV below).

C. Hartree approximation

In the previous section, we solved the CISM Hamiltonian in the HF approximation, i.e. including exchange. In this section we briefly examine the simpler Hartree approximation, because it allows some insight into the parametric sensitivities. Neglecting the exchange interaction, the dependence of the deformation energy $E_{\text{def}}$ on the strength parameter $g$ can be conveniently determined in two steps. As a first step, the single-particle Hamiltonian

$$\hat{h}_{\text{eff}} = \hat{h}^{(0)} + \xi \hat{h}^{(2)}$$

is solved as function of a coupling parameter $\xi$, say in the range $0 < \xi < 2$. One needs to keep the expectation values of $\hat{h}^{(0)}$ and $\hat{h}^{(2)}$ as a function of $\xi$, calculated with the ground-state density matrix $\rho_\xi$ of Eq. (19). For the second step, one determines the ground-state energy $E(g)$ by finding the minimum in

$$E(g) = \min_\xi \left( \langle \hat{h}^{(0)} \rangle_\xi - \frac{1}{2} g \langle \hat{h}^{(2)} \rangle_\xi^2 \right).$$

The minimization in Eq. (20) with respect to $\xi$ is equivalent to the self-consistent condition $g \langle \hat{h}^{(2)} \rangle_\xi = \xi$. The deformation energy is then given by

$$E_{\text{def}}(g) = E(0) - E(g).$$

Fig. 7 shows $\langle \hat{h}^{(0)} \rangle_\xi$ (left panel) and $\langle \hat{h}^{(2)} \rangle_\xi$ (right panel) versus $\xi$. The value of $\xi$ that minimizes the Hartree energy is determined through the competition between the two terms in Eq. (20). Performing next the minimization to get $E(g)$, we find the deformation energy curve shown in the left panel of Fig. 8. The panel on the right in Fig. 8 shows the quadrupole moment of the solution as a function of $g$. The discontinuity of $\langle \hat{Q} \rangle$ at $g \approx 0.037$ MeV$^{-1}$ describes a first order spherical to deformed shape transition. Note that the deformation energy $E_{\text{def}}(g)$ in the deformed phase is a steep function of $g$; thus $g$ is well constrained by the fitting procedure. The solid triangles in Fig. 9 indicate the SLy4 SCMF point (in which the value of $g = 0.0466$ MeV$^{-1}$ is fitted to give the correct deformation energy).

The Hartree deformation energy $E_{\text{def}}$ versus $\langle \hat{Q} \rangle$ in the truncated $sd$ space is plotted in Fig. 9. The solid triangle is the SLy4 SCMF point. It lies very close to the curve extracted from the effective Hamiltonian, showing that the effective theory closely reproduces the quadrupole moment in the truncated space.

D. Correlation energy

With the mapped shell-model Hamiltonian in hand, we are now in a position to compute the correlation energy as defined in Eq. (15). The HF energy of (11) is found to be $E_{\text{HF}} = -147.44$ MeV. The ground-state energy $E_{\text{gs}}$ is calculated using the shell-model code oxbash, putting in the orbital single-particle energies and interaction matrix elements from the mapped Hamiltonian (11). We find $E_{\text{gs}} = -153.93$ MeV. Anticipating calculations in much larger shell-model spaces, we have also computed $E_{\text{gs}}$ with the SMMC. Taking $\beta = 6$ MeV$^{-1}$ and a time slice of $\Delta \beta = 1/32$ MeV$^{-1}$ we find (us-
Truncation effects on the deformed SCMF density matrix are summarized in Table II. The eigenvalues and trace of the truncated density matrix $P \rho_{\text{mf}} P$ (projected on the valence $sd$ shell) are listed separately for protons and neutrons (there are 6 positive $z$-signature eigenvalues for each type of nucleon). We see that the HF condition $\rho_z^2 = \rho$ holds to a rather good accuracy for all four nuclei (all eigenvalues are very close to either 1 or 0). The condition that the trace is given by the number of valence particles is also satisfied quite well.

Table III compares results for the SCMF with the SLy4 Skyrme force and for the HF of the mapped CISM Hamiltonian \cite{ref11} with $\vartheta$ determined by matching the deformation energy of both theories. Shown are the spherical and mean-field total energies, deformation energy and the quadrupole moment $\langle \hat{Q} \rangle$ in the truncated $sd$ shell-model space. We observe that the quadrupole moment of the effective theory deviates by at most 5\% from its value in the SCMF (for $^{28}\text{Si}$).

We also show in Table III the total SCMF quadrupole
TABLE III: SCMF with the SLy4 Skyrme energy functional and the mapped Hamiltonian Eq. (11) for deformed $4N$ $sd$ shell nuclei. All energies are in MeV, $\langle \hat{Q} \rangle$ is in fm$^2$ and $g$ is in MeV/fm$^4$.

| Nucleus | Interaction | $g$ | $E_{\text{ sph}}$ | $E_{\text{ mf}}$ | $E_{\text{ def}}$ | $\langle \hat{Q} \rangle$ | $\langle \hat{Q} \rangle_{\text{ def}}$ | $E_{\text{ corr}}$ |
|---------|-------------|-----|------------------|-----------------|-----------------|-----------------|-----------------|--------------------|
| $^{20}$Ne | SLy4 $\hat{h}^{(2)}$ | 0.077 | -151.83 | -157.43 | 5.6 | 49.0 | 84.0 | 2.39 |
| $^{24}$Mg | SLy4 $\hat{h}^{(2)}$ | 0.0394 | -187.20 | -195.92 | 8.73 | 59.6 | 111.9 | 3.50 |
| $^{28}$Si | SLy4 $\hat{h}^{(2)}$ | 0.0594 | -233.19 | -234.90 | 1.71 | 52.4 | -89.2 | 4.55 |
| $^{36}$Ar | SLy4 $\hat{h}^{(2)}$ | 0.117 | -303.16 | -305.43 | 2.27 | 43.4 | -74.7 | 1.42 |

TABLE IV: Spherical occupations $\langle \hat{n}_j \rangle$ for deformed $4N$ nuclei in the $sd$ shell. The SCMF occupations (computed for the SLy4 Skyrme energy functional) are compared with the occupations for the mapped Hamiltonian Eq. (11) both in the HF approximation and in shell-model calculations.

| Nucleus | Interaction | $p$ $0d_{5/2}$ | $p$ $1s_{1/2}$ | $p$ $0d_{3/2}$ | $n$ $0d_{5/2}$ | $n$ $1s_{1/2}$ | $n$ $0d_{3/2}$ |
|---------|-------------|----------------|----------------|----------------|-----------------|----------------|----------------|
| $^{20}$Ne | SLy4 $\hat{h}^{(2)}$ (HF) | 1.481 | 0.376 | 0.096 | 1.486 | 0.374 | 0.095 |
| $^{24}$Mg | SLy4 $\hat{h}^{(2)}$ (HF) | 3.290 | 0.411 | 0.215 | 3.293 | 0.408 | 0.215 |
| $^{36}$Ar | SLy4 $\hat{h}^{(2)}$ (HF) | 5.919 | 1.751 | 2.297 | 5.921 | 1.758 | 2.287 |

Finally, in Table IV we compare the occupations of the spherical valence orbitals of the SLy4 SCMF theory with similar occupations obtained with the mapped Hamiltonian [11]. The latter are calculated in the HF approximation as well as in the shell-model theory from the expectation values of $\hat{n}_j = \sum_m \hat{a}^\dagger_{jm} \hat{a}_{jm}$ in the correlated CISM ground state.

VII. THE $Q \cdot Q$ INTERACTION

In nuclear physics there is a long history of modeling the effective interaction as a quadrupole-quadrupole interaction of the form $Q \cdot Q$, including Elliott’s SU(3) model [20] and the pairing-plus-quadrupole model of Kisslinger and Sorensen [21]. Since it is possible in some contexts to exploit the properties of the quadrupole operator, it is of interest to see how well a quadrupole-quadrupole interaction performs in our context. Thus, we apply the Hamiltonian

$$\hat{H} = \hat{h}^{(0)} - \frac{1}{2} g \hat{Q} \cdot \hat{Q}. \quad (22)$$

in the truncated shell-model space with the matrix elements of $\hat{h}^{(0)}$ and the mass quadrupole operator $\hat{Q}$ determined by the appropriate mean-field theory. We follow the same procedure we used before to determine the coupling constant $g$.

We have determined such an effective $\hat{Q} \cdot \hat{Q}$ interaction for the deformed $4N$ $sd$ shell nuclei $^{20}$Ne, $^{24}$Mg, $^{28}$Si and $^{36}$Ar, using the HF approximation of the USD interaction as the SCMF theory. We observe...
that the quadrupole moments calculated with both effective interactions $\hat{h}^{(1)}$ and $\hat{h}^{(2)}$ are quite close to the USD results. This can be seen in Fig. 10, where $\langle \hat{Q} \rangle$ (in fm$^2$) is plotted versus mass number $A$. Results for the $\hat{Q} \cdot \hat{Q}$ interaction (solid triangles) and the interaction with $\hat{h}^{(2)}$ (solid squares) are compared with the USD results (open circles). The effective Hamiltonians $\hat{h}^{(1)}$ and $\hat{h}^{(2)}$ also yield similar occupations $\langle \hat{n}_j \rangle$ (see Fig. 2) and correlations energies (see Fig. 3).

![FIG. 10: Quadrupole moment in the SCMF method versus mass number $A$ for 4$N$ sd-shell nuclei. Results are shown for two effective CISM Hamiltonians: the $\hat{Q} \cdot \hat{Q}$ interaction (solid triangles) and the $\hat{h}^{(2)} \cdot \hat{h}^{(2)}$ interaction (solid squares). They are compared with results for the USD Hamiltonian (open circles).](image)

**VIII. PROSPECTS AND CONCLUSIONS**

We are encouraged by the results we found for constructing a isoscalar quadrupolar interaction from the SCMF in the sd-shell, but much remains to be done to demonstrate that the method will be useful in deriving a global theory of correlation energies. We list some of the major tasks that should be addressed:

- So far, we have carried out the mapping only for nuclei that have deformed HF ground states. If the lowest-energy mean-field solution is spherical, we can use a constraining quadrupole field $\lambda \hat{Q}$ to deform the HF ground state and use this deformed solution to construct the effective CISM Hamiltonian. In particular, for a given strength $\lambda$ of the field, we can compare the deformation energy of the constrained SCMF theory with the deformation energy of the effective Hamiltonian in the constrained HF approximation. The Hartree-Fock Hamiltonian $\hat{h}^{(1)}$ of the effective CISM Hamiltonian is now replaced by

$$h_{ac}^\lambda = h_{ac}^{(0)} - g \sum_M (-)^M h_{M;ac}^{(2)} \left( \sum_{bd} h_{-M;bd}^{(2)} \rho_{db}^\lambda \right) + g \sum_M (-)^M \sum_{bd} h_{M;ad}^{(2)} h_{-M;bc}^{(2)} \rho_{db}^\lambda + \lambda Q_{ac},$$

where $\rho_{ac}^\lambda$ is the self-consistent density matrix in the spherical basis (here $a = (jm)$) and $\hat{Q}$ is the axial quadrupole operator. A practical problem is how large should we take $\lambda$ to be. One possibility is to use the value of $\lambda$ that gives the deformed $J = 0$ minimum.\(^{14}\)

- The real challenge of a global theory is the heavy, open-shell nuclei, and we should ask whether any severe problems will arise when we attempt to map the SCMF onto the CISM Hamiltonian. As a concrete example, consider the nucleus $^{162}$Dy. It is strongly deformed, having an intrinsic quadrupole moment around 18 bn. The SCMF using the SLy4 interaction reproduces this value very well.\(^{18}\) The deformation energy when pairing is not included is quite large (about 20 MeV). The $J = 0$ projection and quadrupole fluctuations (in the presence of a delta pairing force) contribute an additional correlation energy of $\approx 3.4$ MeV.\(^{10}\)

To use an CISM Hamiltonian to calculate deformation and correlation energies, one would need to truncate to a basis of the full shells, i.e., $N = 82 - 126$ for neutrons and $Z = 50 - 82$ for protons plus possibly some intruder states. Thus, a first test would be to see how well the consistency conditions Eqs. 6 and 10 are satisfied in such a truncated space.

The next question that arises is the practicality of various implementations of the CISM Hamiltonian for solving in such large spaces. There are several approaches that have promise, including the shell model Monte Carlo (SMMC) method,\(^{6,9,22}\) the Monte Carlo shell model\(^{28}\) and direct matrix methods such as those used by the Strasbourg group\(^{21}\). One could also consider more approximate theories of the correlation energy, such as the RPA.\(^{27}\)

- The mapping should be extended to include all other parts of the interaction that we believe are well-determined in the SCMF, in particular the interaction in the isovector monopole and quadrupole densities. This should be straightforward to carry out, requiring only that nuclei with $N \neq Z$ be considered in a more general parameterization of the interaction.

\(^{14}\) Nuclei with spherical HF minimum acquire deformation after applying a $J = 0$ projection.\(^{18}\)
The pairing interaction is very important in nuclear structure but is notoriously difficult to specify in detail, i.e., its density dependence and its spatial range (or equivalently the orbital space cut-off). Most SCMF energy functionals include a pairing interaction, but only an overall strength can be reliably determined in phenomenological theories. Thus, absent an \textit{ab initio} theory of the pairing interaction, we cannot expect to have a reliable decomposition of the pairing fields in the mean-field Hamiltonian. Still, it should be possible to determine a strength parameter controlling the overall magnitude of pairing effects. Note that in the presence of pairing, the single-particle density matrices of the SCMF theory have the BCS form. Thus, to parameterize the density matrices, one would use the HFB or the HF-BCS approximations for the self-consistent mean-field theory.

An unsatisfactory aspect of our mapping is the lack of unitarity of the transformation from the deformed basis to the truncated spherical basis. It would be useful to investigate mapping formalisms that yield density matrices that satisfy exactly the criteria we have discussed, i.e., Eqs. \ref{eq:unitary} and \ref{eq:signature}.

Despite this long list of tasks to be done, we believe that the methods discussed here can be developed to produce a well-justified global theory of shell-model correlation energies, starting from a global SCMF theory (e.g., a Skyrme energy functional). Particularly encouraging is the finding that quadrupole correlations account for more than half the correlation energy in \textit{sd} shell nuclei.

\section*{Acknowledgments}

We thank M. Bender and P.H. Heenen for instructions in using the SCMF code \texttt{ev8}. Y.A. would like to acknowledge the hospitality of the Institute of Nuclear Theory in Seattle where part of this work was completed. This work was supported by the U.S. Department of Energy under Grants FG02-00ER41132 and DE-FG-02-91ER40608.

\section*{Appendix: wave functions and matrix elements in the $z$-signature representation}

A deformed single-particle wave function in the density functional theory is a two-component complex spinor $\psi = \begin{pmatrix} \psi_+ \\ \psi_- \end{pmatrix}$, or alternatively a four-component real spinor $\psi = \begin{pmatrix} \psi_1 & \psi_2 \\ \psi_3 & \psi_4 \end{pmatrix}$ with $\psi_1 = \text{Re}\psi_+ + \psi_2 = \text{Im}\psi_+ ; \psi_3 = \text{Re}\psi_- + \psi_4 = \text{Im}\psi_-$. The SCMF orbitals are chosen to have good parity $\pi$ and good $z$-signature \cite{13}. Such states have well-defined symmetry properties under each the three plane reflections $x \rightarrow -x$; $y \rightarrow -y$; $z \rightarrow -z$ (see Table 1 in Ref. \cite{16}). Therefore it is sufficient to store their values in one octant (e.g., $x, y, z > 0$).

The scalar product of two spinors $\psi$ and $\phi$ is a complex number with $\text{Re} (\psi \dagger \phi) = \sum_i \psi_i \phi_i$ and $\text{Im} (\psi \dagger \phi) = \psi_1 \phi_2 - \psi_2 \phi_1 + \psi_3 \phi_4 - \psi_4 \phi_3$.

The find a standard basis of spherical single-particle orbitals $|j m\rangle$, it is necessary to calculate matrix elements of the quadrupole operators $\hat{Q}_{\pm}^{(2)}$ (see Section \\ref{sec:quadrupole}). We denote the $j + 1/2$ orbitals with good $j$ and positive $z$-signature by $|ja\rangle$. The quadrupole matrix elements are obviously real and given by

$$
\langle ja|\hat{Q}_{\pm}^{(2)}|jb\rangle = 8 \int_{x,y,z>0} dx\, dy\, dz \, (2z^2 - x^2 - y^2) \, \text{Re} (\psi \dagger \phi) \, ,
$$

where $\psi$ and $\phi$ are the spinors describing $|ja\rangle$ and $|jb\rangle$, respectively.

To calculate $\langle ja|\hat{Q}_{\pm}^{(2)}|jb\rangle$ we use the symmetry properties of the orbitals under plane reflections and find

$$
\langle ja|xz|jb\rangle = 8 \int_{x,y,z>0} dx\, dy\, dz \, xz \, \text{Re} (\psi \dagger \phi) \, ,
$$

$$
\langle ja|yz|jb\rangle = 8i \int_{x,y,z>0} dx\, dy\, dz \, yz \, \text{Im} (\psi \dagger \phi) \, ,
$$

Since the time-reversed spinor of $\phi$ is given by $\bar{\phi} = \begin{pmatrix} \phi_1 \\ -\phi_3 \\ \phi_2 \\ -\phi_4 \end{pmatrix}$, we have

$$
\text{Re} (\psi \dagger \bar{\phi}) = \psi_1 \phi_3 - \psi_3 \phi_1 + \psi_4 \phi_2 - \psi_2 \phi_4 \, ,
$$

$$
\text{Im} (\psi \dagger \bar{\phi}) = \psi_3 \phi_2 - \psi_2 \phi_3 + \psi_4 \phi_1 - \psi_1 \phi_4 \, .
$$

The matrix elements of $\hat{Q}_{\pm}^{(2)}$ are calculated from \ref{eq:Q2}, using $Q_{\pm}^{(2)} = \mp \sqrt{6} \sum_i (x_i z_i \pm i y_i z_i)$ (they are all real).

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