Natural orbitals for the \textit{ab initio} no-core configuration interaction approach

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\textbf{I. INTRODUCTION}

The goal of \textit{ab initio} nuclear theory \cite{1,11,12} is to predict the behavior of the nuclear many-body system starting from underlying internucleon interactions \cite{11,14}. However, the nuclear many-body problem lives in an infinite-dimensional space. Thus, in practical numerical computations, the problem must be replaced by an approximate, truncated representation, and, given finite computational resources, can only be solved with finite accuracy.

This accuracy may be expected to depend critically upon the choice of many-body basis used to define the truncated space for the problem. The many-body basis is in turn generated from some underlying set of single-particle states. More specifically, given the rotational invariance of the nuclear problem, we consider some underlying set of single-particle orbitals, of definite angular momentum $j$. While the choice of orbitals has been a central concern in quantum many-body calculations for the electron structure of atoms and molecules \cite{15}, it has been largely neglected in \textit{ab initio} nuclear many-body calculations.

In the no-core configuration interaction (NCCI), or no-core shell model (NCSM), approach \cite{11,9}, the many-body basis consists of antisymmetrized products (Slater determinants) of single-particle states. The many-body problem is then recast as a Hamiltonian matrix eigenproblem with occupation of low-lying orbitals, thus accelerating convergence in a configuration-interaction basis, while also possibly providing physical insight into the single-particle structure of the many-body wave function. We describe the implementation of natural orbitals in the NCCI framework, examine the nature of the natural orbitals thus obtained, the properties of the resulting many-body orbitals, and the convergence of observables. After taking $^3$He as an illustrative testbed, we explore aspects of NCCI calculations with natural orbitals for the ground state of the $p$-shell neutron halo nucleus $^8$He.

Calculations. This choice is motivated in part by technical convenience. Namely, two-body matrix elements of translationally invariant operators such as the Hamiltonian are conveniently evaluated in the oscillator basis, via the Moshinsky transformation \cite{16} from the relative oscillator basis. Furthermore, an exact separation of the center-of-mass motion is obtained with an oscillator basis truncated according to the $N_{\text{max}}$ scheme, \textit{i.e.}, by total number of oscillator excitations \cite{17,18}.

The calculated results for energies, electromagnetic observables, \textit{etc.}, from an \textit{ab initio} NCCI calculation depend on the truncated space in which this calculation is carried out. As $N_{\text{max}}$ is increased towards infinity, the calculated results in principle converge towards those which would be obtained in the full, untruncated space for the nuclear many-body problem. However, a rapid growth in dimension of the many-body space, with increasing $N_{\text{max}}$ and number of nucleons, limits the accuracy which can be obtained.

Despite the computationally-convenient properties of the harmonic-oscillator orbitals, within a many-body calculation, there is no reason to presume that they are “optimal” as the underlying single-particle basis for expanding the many-body wave function. Moreover, in at least one way, they are qualitatively mismatched to the problem. Notably, as the solutions to the infinitely-bound harmonic oscillator problem, the oscillator functions fall off at large distance with Gaussian asymptotics, \textit{i.e.}, $\sim e^{-\kappa r}/(2\kappa^2)$. However, the nuclear attraction is of finite-range. Consequently, the single-particle wave functions arising in mean-field descriptions of the nucleus instead fall off exponentially, \textit{i.e.}, $\sim e^{-\kappa r}$. While a suitable fall-off can be recovered, out to any finite distance of relevance, by taking a superposition of oscillator functions, doing so may require a large number of oscillator functions (see, \textit{e.g.}, Fig. 4 of Ref. \cite{19}).
We are therefore motivated to look beyond the traditional harmonic oscillator many-body basis, to increase the accuracy attainable for a given NCCI problem dimension. In the present work, we explore the improvement which may be obtained by optimizing the choice of underlying orbitals used to construct the basis configurations. While we might simply prescribe a set of orbitals of some analytic form (e.g., Refs. [20, 21]), in the hopes that these might provide some benefit relative to the harmonic oscillator orbitals, a more informed choice can be obtained by first carrying out some preliminary many-body calculation, and using the resulting information on the many-body wave function for guidance in constructing the orbitals.

In this spirit, the natural orbitals [22, 26] have been used extensively in atomic and molecular electron-structure theory [15, 25], and have also found application in the nuclear problem [27, 31]. They are constructed in a way intended to reduce the number of antisymmetrized product states required for an accurate representation of the many-body wave function, thereby accelerating the convergence of its description in a configuration interaction basis [32, 33].

Natural orbitals are defined with reference to some many-body state \(|\Psi\rangle\) — not necessarily a single Slater determinant, but a general, correlated many-body state. The corresponding set of natural orbitals is obtained by diagonalizing the one-body density matrix of \(|\Psi\rangle\). The eigenvectors define the natural orbitals, and the corresponding eigenvalues represent the mean occupations of these orbitals within the reference many-body state \(|\Psi\rangle\).

In order to find the true natural orbitals for a given nuclear state, say, the ground state, we would have to have first solved the full many-body problem for this state, thence obtaining the densities. However, even from an approximate initial solution for the many-body wave function, which yields approximate densities, we may still obtain approximate natural orbitals. It is these which we may attempt to use in constructing an improved basis for the many-body calculation.

Here we explore the use of natural orbitals in NCCI calculations. The initial many-body calculation, providing the densities used to define the natural orbitals, is a traditional \(N_{\text{max}}\)-truncated oscillator-basis calculation. The natural orbitals for a subsequent many-body calculation are thus obtained as a unitary transformation on the original oscillator orbitals. In addition to illustrating the convergence properties of the resulting NCCI calculations, we attempt to illuminate the properties of the natural orbitals and probe some of the implications for center-of-mass motion.

Preliminary results of the present work were reported in Refs. [34, 35]. Complementary approaches have since also been explored where natural orbitals for use in \textit{ab initio} nuclear many-body calculations are obtained from solutions of a spatially-localized two-body (deuteron) problem [36] or from many-body perturbation theory for closed-shell nuclei [37, 38]. The implications of natural orbitals for wave function entanglement in NCCI calculations have also been examined [39].

Whereas the preliminary results presented in Refs. [34, 35] were based on the earlier JISP16 interaction [29], the present examples are based on NCCI calculations using the Daejeon16 internucleon interaction [40]. Relative to JISP16, Daejeon16 has the advantage of providing both faster convergence of calculated observables and improved agreement with experimental binding and excitation energies [41].

We first review the framework for calculations with natural orbitals: defining how symmetry-adapted natural orbitals (of definite angular momentum and parity) are extracted from the density matrix (Sec. 1.1A) and outlining how these are obtained and used within the NCCI framework (Sec. 1.1B). Then, to see how the formalism is reflected in actual NCCI calculations, we take \(^3\)He as the simplest nontrivial example: examining the convergence of energy and radius observables for \(^3\)He (Sec. 1.1B), inspecting the radial wave functions of the natural orbitals themselves (Sec. 1.1B), and diagnosing the center-of-mass motion of the many-body wave function (Sec. 1.1C). After establishing this baseline, we explore aspects of NCCI calculations for the neutron halo properties of \(^6\)He (Sec. 1.4).

II. NATURAL ORBITALS

A. Natural orbitals and rotational symmetry

Recall that we seek orbitals which will provide rapid convergence in a finite basis of antisymmetrized product states. Our many-body basis is built out of an ordered set of single-particle orbitals where we favor “lower-lying” orbitals (and disfavor “higher-lying” orbitals) when deciding which orbitals to use in constructing basis states. We would therefore be best served by a set of orbitals such that the “lower-lying” orbitals contribute disproportionately to the most important antisymmetrized products.

It is therefore natural to construct orbitals in a way that maximizes the mean occupation of the lowest-lying orbitals — and, correspondingly, minimizes the mean occupation of the higher-lying orbitals — in the many-body state. In a particle-hole picture, this may be thought of as minimizing the depletion of the Fermi sea. Natural orbitals, in a well-defined sense, accomplish this goal.

Suppose we are interested in finding single-particle states in which to efficiently represent a particular many-body state \(|\Psi\rangle\). The single-particle properties of \(|\Psi\rangle\) are described by its \textit{one-body density operator} \(\hat{\rho}\) [42], which is an operator on the single-particle space. \textit{Natural orbitals} are, quite simply, obtained as its eigenstates.

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1 Such a one-body density operator, derived from a pure state of a many-body system, is properly known as a \textit{one-body reduced density operator} [43].
Taken in this traditional sense \[22,24\], the natural “orbitals” are not orbitals per se, as usually construed in nuclear physics. They are, rather, simply a set of independently-defined single-particle states, unrelated to each other by any explicit symmetry constraint. However, one may refine the definition of the natural orbitals, so as to manifestly respect the symmetries of the system \[25,44\]. In the case of the rotationally-invariant nuclear problem, the resulting symmetry-adapted natural orbitals become orbitals in the usual nuclear-physics sense, of \(nlj\) orbitals \[45\]. In the following, we first review the formulation of natural orbitals in the traditional sense, \textit{i.e.}, without explicitly embedding the nuclear symmetries, then establish the symmetry-adapted natural orbitals appropriate to nuclear NCCI calculations.

Although the definition of \(\hat{\rho}_\Psi\) as an operator is independent of the choice of basis for the single-particle space, this operator may be expressed in terms of any discrete single-particle basis as

\[
\hat{\rho}_\Psi = \sum_{\alpha\beta} |\alpha\rangle \langle \alpha| \rho_{\alpha\beta} |\beta\rangle \langle \beta|, \tag{1}
\]

Here, the labels \(\alpha\) and \(\beta\) specify the single-particle basis states, \textit{e.g.}, for the nuclear problem, they may represent the magnetic substates \(\alpha = (n,\ell, m, \alpha)\) of \(nlj\) orbitals \[45\], while \(a_\alpha^\dagger\) and \(a_\alpha\) represent the creation and annihilation operators, respectively, for a nucleon in state \(\alpha\).

The eigenstates \(|\phi_i\rangle\) of \(\hat{\rho}_\Psi\) are what we take as the natural orbitals for the many-body reference state \(|\Psi\rangle\). In terms of this eigenbasis, the expression (1) for \(\hat{\rho}_\Psi\) reduces to the familiar canonical form for a density operator as a real linear combination of projection operators (\textit{e.g.}, Ref. \[46\]),

\[
\hat{\rho}_\Psi = \sum_i \lambda_i |\phi_i\rangle \langle \phi_i|, \tag{2}
\]

where the \(\lambda_i\) are the corresponding real eigenvalues for the \(|\phi_i\rangle\) (we rely here on the observation that \(\hat{\rho}_\Psi\) is a self-adjoint operator).

If we work in terms of a discrete basis for the single-particle space, \(\hat{\rho}_\Psi\) is represented as the one-body density matrix \(\rho\), with matrix elements \(\rho_{\alpha\beta} = \langle \alpha|\hat{\rho}_\Psi|\beta\rangle\), which may be read off from (1) as

\[
\rho_{\alpha\beta} = \langle \Psi|a_\beta^\dagger a_\alpha|\Psi\rangle. \tag{3}
\]

The operator eigenproblem for \(\hat{\rho}_\Psi\) reduces to the matrix eigenproblem for \(\rho\). The eigenvectors then express the natural orbitals \(|\phi_i\rangle\) in terms of the underlying basis. Changing to a natural orbital basis for the single-particle space makes the density matrix diagonal, with entries \(\lambda_i\), as is apparent from (2).

A diagonal matrix element \(n_\alpha = \rho_{\alpha\alpha}\) is simply the expectation value of the number operator \(\hat{N}_\alpha = a_\alpha^\dagger a_\alpha\), and thus represents the mean occupation of the single-particle state \(\alpha\) in the many-body state \(|\Psi\rangle\). Thus, the eigenvalues \(\lambda_i\) for the natural orbitals \(|\phi_i\rangle\) of a reference state \(|\Psi\rangle\) represent their mean occupations in this reference state, \textit{i.e.}, \(n_{\phi_i} = \lambda_i\). Consequently, these eigenvalues must satisfy the properties expected for mean occupations: \(0 \leq \lambda_i \leq 1\) and \(\sum_i \lambda_i = A\), where \(A\) is the number of nucleons in the system.

To see the relevance of the natural orbitals to the problem of identifying an optimal basis of antisymmetrized product states, first consider the case where \(|\Psi\rangle\) itself a single antisymmetrized product, specifically, of the first \(A\) single-particle states taken from some particular discrete basis \(|\{\alpha_i\}\rangle\), \textit{i.e.}, \(|\Psi\rangle = |\alpha_1, \alpha_2, \ldots, \alpha_A\rangle\). The one-body density matrix taken in this basis is already diagonal, with occupation numbers \(n_{\alpha_i} = 1\) for occupied states or 0 for unoccupied states \[42,47\].

If we were instead working in terms of some other single-particle basis \(|\{\beta_i\}\rangle\), \(|\Psi\rangle\) would not manifestly be represented as a single antisymmetrized product state. However, evaluating the density matrix in this basis \(|\{\beta_i\}\rangle\), and diagonalizing the resulting matrix, will recover the \(|\{\alpha_i\}\rangle\) basis as the natural orbital basis, thereby revealing \(|\Psi\rangle\) as a single antisymmetrized product state. (More properly, it will recover the \(|\{\alpha_i\}\rangle\) basis to within an arbitrary freedom of choice of basis within the spaces spanned by the occupied and unoccupied orbitals separately, as each of these represents a degenerate eigenspace of the density operator, with eigenvalues 1 and 0, respectively.) Such a transformation back to a single antisymmetrized product state is possible if and only if the density matrix has eigenvalues which are all either 0 or 1 \[43\].

Of course, we are more generally interested in many-body states which incorporate correlations. There is no single-particle basis in which such a state can be represented as a single antisymmetrized product, and the eigenvalues of the one-body density operator are no longer simply 0 and 1.

However, the transformation to the natural orbital basis still generates single-particle states for which the mean occupations “fall as quickly as possible,” in a very particular sense. Namely, we order the natural orbitals \(|\phi_i\rangle\) by decreasing eigenvalue \(\lambda_1 \geq \lambda_2 \geq \ldots\), that is, in order of decreasing mean occupation \(n_{\phi_i} = \lambda_i\). The total mean occupation of the first \(q\) single-particle states in any basis is \(n_{\leq q} = \sum_{i=1}^q \rho_{ii}\), and the total mean occupation of the first \(q\) natural orbitals, in particular, is \(n'_{\leq q} = \sum_{i=1}^q \lambda_i\). By a general property of traces of Hermitian matrices \[48\], the partial trace (sum of the first \(q\) diagonal entries) in any basis is bounded from above by the partial trace in the eigenbasis (sum of the first \(q\) eigenvalues). Thus,

\[
n_{\leq q} = \sum_{i=1}^q \rho_{ii} \leq \sum_{i=1}^q \lambda_i = n'_{\leq q}. \tag{4}
\]

That is, for any \(q\), the natural orbitals constitute the basis which maximizes the total mean occupation of the first \(q\) single-particle states \[42\].

The “naive” or generic natural orbitals as defined
above, by simply diagonalizing $\rho$ without further precautions, fail to take into account the symmetry properties of the system. Despite their name, these natural orbitals are simply an independent set of single-particle states, without well-specified quantum numbers, rather than orbitals per se, in the sense that “orbitals” are usually meant in rotationally-invariant problems, as we now elaborate.

Consider, in particular, the symmetries present in nuclear configuration-interaction calculations. To ensure that the many-body space supports states of definite angular momentum and parity, the single-particle states used to build the basis configurations are not arbitrary, but must form orbitals in the traditional shell-model sense. An orbital is a set of magnetic substates $|nljm\rangle$ ($m = -j, \ldots, +j$), which together form an angular momentum multiplet of definite $j$ and definite parity $P = (-)^j$. (Since $l$ and $j$ can differ only by $1/2$, the condition of definite parity is sufficient to also enforce definite $l$.) The different magnetic substates of the orbital are related to each other by angular momentum ladderings and share the same radial wave function $R_{nlj}$.

The many-body reference state $|\Psi_{JM}^P\rangle$ for a nuclear configuration-interaction calculation will have definite angular momentum $J$, projection $M$ (assuming the problem is formulated in the $M$ scheme [49]), and parity $P$. These properties of the reference state serve to impose some, but not all, of the requisite properties for the natural orbitals to constitute true $nlj$ orbitals (or the $m$-substates thereof). By inspection of $|\Psi_{JM}^P\rangle$, and the additive nature of the $m$ quantum number, it is clear that the density matrix for a reference state $|\Psi_{JM}^P\rangle$ cannot connect single-particle states $|nljm\rangle$ with different $m$. Similarly, by the multiplicative nature of parity, it cannot connect single-particle states of different parity.

However, the density matrix will in general connect single-particle states $|nljm\rangle$ with different $j$, leading to natural orbitals without definite angular momentum.$^2$ Due to such considerations, calculations involving natural orbitals are instead commonly based on symmetry-adapted natural orbitals [25, 43]. These are obtained by diagonalizing only that part of the one-body density matrix which is invariant under the action of the symmetry group, namely, for the present problem, angular momentum (and parity).

We construct a rotational scalar one-body density matrix $\tilde{\rho}$ in terms of the spherical tensor coupled product $^3$ of the creation and annihilation operators for an orbital.$^4$ This rotational scalar one-body density matrix has elements

$$\tilde{\rho}_{ab} = \langle \Psi_{JM}^P | [a_b^\dagger \tilde{a}_a]_{00} | \Psi_{JM}^P \rangle,$$

or, equivalently, in terms of the original, uncoupled one-body density matrix elements defined in $^5$, $\tilde{\rho}_{ab} = \delta_{\alpha a} \delta_{\beta b} \sum_m \rho_{(n_a la jm)(n_b lb jm)}$, where we adopt the notation $j \equiv (2j + 1)^{1/2}$.

This scalar density matrix $\tilde{\rho}$ is now simply a matrix with respect to orbitals (labeled by $a$), rather than their magnetic substates (labeled by $\alpha$). The matrix elements $\tilde{\rho}_{ab}$ must be independent of the magnetic substate $M$ of the reference state, since they are given in $^3$ as the matrix element of a scalar operator in the many-body space.$^6$

Nonzero matrix elements $\tilde{\rho}_{ab}$ only arise between orbitals of the same angular momentum ($j_a = j_b$), parity, and thus (as argued above) orbital angular momentum ($l_a = l_b$). That is, the scalar one-body density matrix is block diagonal in $lj$. Symmetry-adapted natural orbitals, obtained as eigenvectors of $\tilde{\rho}$, may thus be found by diagonalizing independently within each $lj$ subspace. The resulting natural orbitals are related to the underlying orbitals simply by a unitary transformation

$$|\phi_{n'ljm}\rangle = \sum_n A_{nn'} |nljm\rangle$$

on the radial wave functions $R_{nlj}$ within each $lj$ space separately.

The total number operator, summed over all magnetic substates of an orbital, is $N_a = j_a [a_a^\dagger \tilde{a}_a]_{00}$. Thus, the diagonal matrix elements $\tilde{\rho}_{aa}$ of the scalar density matrix are proportional to the mean occupation of the orbital $a$,

$$n_a = j_a \tilde{\rho}_{aa},$$

which ranges from 0 to the degeneracy $2j_a + 1$ of the orbital. For the symmetry-adapted natural orbital $\phi_a$, the corresponding eigenvalue $\lambda_a$ is then proportional to the mean occupation of the orbital.$^6$

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$^2$ Only for the special case of a reference state with $J = 0$ do spherical tensor selection rules prevent the density matrix from connecting and thus mixing single-particle states of different $j$. Even here, caution would be necessary in diagonalizing $\rho$, as it would contain redundant $ljm$ blocks, one for each $m = -j, \ldots, j$. Diagonalizing these blocks together would lead to degeneracies and thus ambiguity (and, in general, undesirable $m$-mixing) in the choice of eigenstates within each degenerate eigenspace, while diagonalizing each block of definite $m$ independently would still fail to enforce consistent phase relations between the $m$-substates of an $nlj$ orbital.

$^3$ The creation and annihilation operators for the magnetic substates $\alpha = (n_a la jm_a)$ of an orbital $a = (n_a la j)$ together constitute spherical tensors $a_a^\dagger$ and $\tilde{a}_a$ with components $\langle a_a^\dagger |m_{a_1} la jm |a_a\rangle = a_{m_{a_1} la jm_a}$ and $\langle \tilde{a}_a |m_{a_1} la jm |a_a\rangle = (-)^{j_a + m_{a_1}} a_{m_{a_1} la jm_a}$, respectively [43].

$^4$ Alternatively, the vestigial reference to the $M$ quantum number in $^3$ can be eliminated by recourse to the Wigner-Eckart theorem [50], which gives $\tilde{\rho}_{ab} = J^{-1} \langle \Psi_{JM}^P | [a_b^\dagger \tilde{a}_a]_{00} | \Psi_{JM}^P \rangle$.

$^5$ If the spherical tensor annihilation operator in footnote $^3$ is instead defined with the common alternative phase convention $(a_a^\dagger)_{ja, m_a} = (-)^{j_a - m_{a_1}} a_{m_{a_1} la jm_a}$, which differs by an overall sign, then we instead have $N_a = -j_a [a_a^\dagger \tilde{a}_a]_{00}$, and $n_a = -j_a \tilde{\rho}_{aa}$. 
Ordering the natural orbitals by decreasing eigenvalue, separately within each $lj$ subspace, serves to define a radial $n$ quantum number, which is now simply a counting index with no strict relation to the number of radial nodes. Ordering by decreasing eigenvalue or, equivalently, decreasing mean occupation, again serves to maximize the mean occupation of the “lower-lying” orbitals, as in [4], but now only within each $lj$ subspace.

For the rotationally-invariant many-body problem with symmetry-adapted natural orbitals, in contrast to the situation above for “naive” natural orbitals, we would not in general expect the transformation to natural orbitals to reveal a many-body reference state to be a single antisymmetrized product state. Even for a pure shell-model configuration, i.e., defined by a specific distribution of nucleons over $nljm$ orbitals, a state of definite $J$ is in general obtained as a linear combination of many such antisymmetrized product states, involving different choices of occupied $m$-substates for each orbital, as required to couple the angular momenta of the individual nucleons to yield resultant total angular momentum $J$ [49]. Transformation to the symmetry-adapted natural orbitals serves to reveal if a reference state can be represented, not as a single antisymmetrized product state, but rather as a pure shell-model configuration, for some choice of basis orbitals. More generally, it serves to allow the expansion of the many-body wave function in terms of fewer low-lying configurations.

B. Natural orbitals in the NCCI framework

In NCCI calculations [10], the many-body basis consists of antisymmetrized product states built from some underlying orbitals, usually those of the three-dimensional isotropic harmonic oscillator. The nuclear Hamiltonian

$$H = T_{\text{intr}} + V + a N_{\text{c.m.}}.$$  

(8)

is then represented as a matrix in terms of this basis. Here $T_{\text{intr}}$ is the two-body intrinsic kinetic energy operator [18] [52] [53], $V$ represents the internucleon interaction (typically limited to two-body or three-body contributions), and the final Lawson term [49] [54] [55], proportional to the number operator $N_{\text{c.m.}}$ for center-of-mass oscillator quanta, optionally serves to control the center-of-mass motion (as discussed further below).

The NCCI many-body basis states are defined as antisymmetrized products of single-particle states described by quantum numbers $nljm$, where $n$ is the radial quantum number, $l$ the orbital angular momentum, $j$ the resultant angular momentum after coupling to spin, and $m$ its projection. Each product state thus has definite total angular momentum projection $M = \sum_{i=1}^{A} m_i$ and parity $P = \Pi_{i=1}^{A} (-1)^{l_i}$. In a typical $M$-scheme calculation [49], the basis is restricted to fixed $M$ and $P$. The individual basis states do not have definite angular momentum, but, since the Hamiltonian is rotationally invariant, states of definite total angular momentum $J \geq |M|$ emerge from the diagonalization.

In the usual case where we adopt oscillator orbitals, each orbital is furthermore identified by its oscillator major shell, or number of oscillator quanta, $N = 2n + l$ [45]. An antisymmetrized product state then has $N = \sum_{i=1}^{A} N_i$ oscillator quanta overall, where $N_i = 2n_i + l_i$ represents the number of oscillator quanta contributed by the $i$th particle. The total number of quanta may be reexpressed as $N = N_0 + N_{\text{ex}}$, where $N_0$ is the number of quanta in the lowest filling of oscillator shells permitted by the Pauli principle for the given nucleus, so that $N_{\text{ex}}$ then represents the number of excitation quanta relative to this lowest filling.

The $N_{\text{max}}$ truncation scheme restricts the basis configurations to those with $N_{\text{ex}} \leq N_{\text{max}}$, that is, limiting the total number of excitation quanta. Thus, $N_{\text{max}} = 0$ yields a traditional “0$\hbar \omega$” shell model space, in which all nucleons are restricted to the valence shell (and an inert core). Since the parity of a harmonic oscillator configuration is $P = (-)^{N_0+N_{\text{ex}}}$, a basis consisting of configurations with $N_{\text{ex}} = 0, 2, \ldots, N_{\text{max}}$ (with $N_{\text{max}}$ even) yields a truncated space of the same parity as the lowest oscillator configuration (normal parity), while a basis consisting of configurations with $N_{\text{ex}} = 1, 3, \ldots, N_{\text{max}}$ (with $N_{\text{max}}$ odd) yields a truncated space of the opposite parity (nonnormal parity) [56]. The growth in dimension of the nuclear many-body space with increasing $N_{\text{max}}$ is illustrated in Fig. 1 for selected nuclides through the lower $sd$ shell.

The truncated space spanned by such a basis, and thus the results of an NCCI calculation, depend on both the many-body basis truncation parameter $N_{\text{max}}$ and the oscillator length $b$ of the underlying oscillator single-particle basis. This length scale is commonly quoted as an oscillator energy $\hbar \omega$, in terms of which $b = (\hbar c)/[(m_N c^2)(\hbar \omega)]^{1/2}$, where $m_N$ is the mean nucleon mass ($m_N c^2 \approx 938.92$ MeV). See, e.g., Refs. [37] [38] for illustrations of convergence of observables with respect to these basis parameters.

The $N_{\text{max}}$ truncation for the oscillator basis holds a special place in NCCI calculations due to its properties

6 The notable exception is a closed-shell configuration, for which the resulting $J = 0$ state is indeed simply an antisymmetrized product state.
FIG. 1. Dimension of the NCCI many-body space as a function of the number of oscillator excitations \( N_{\text{max}} \) included in the basis, including for \(^{8,6}\)He (highlighted). The dimension of the FCI space constructed from the same orbitals is also shown for \(^4\)He (dotted gray line). Dimensions are those obtained with \( M \)-scheme bases (\( M = 0 \) for even-mass nuclei, or \( M = 1/2 \) for odd-mass nuclei) for the normal-parity space.

regarding center-of-mass motion. The physically-relevant degrees of freedom for describing nuclear structure and excitations reside in the motion of the nucleons relative to their common center of mass, rather than in the motion of this center of mass relative to the laboratory frame. However, given that the NCCI approach is formulated in terms of antisymmetrized products of single-particle states defined with respect to the laboratory frame, the center-of-mass coordinate cannot be strictly eliminated as a degree of freedom in the many-body wave function. Nonetheless, this motion can at least be brought into a known, controlled form.

Namely, the \( N_{\text{max}} \) truncation, in particular, ensures that nuclear eigenstates can be obtained exhibiting an exact separation between a pure oscillator \( 0s \) wave function for the center of mass coordinate (i.e., the center-of-mass degree of freedom is frozen into its zero point motion) and an intrinsic wave function for the motion of the nucleons relative to each other (see Sec. II.B of Ref. [18] for a detailed explanation of the reasoning). The Lawson term in [8] selects such eigenstates with \( 0s \) center-of-mass motion, by shifting any remaining states involving center-of-mass excitation out of the low-lying spectrum. Thus, states involving excitation of the intrinsic wave function are cleanly separated from what would otherwise be a thicket of spurious excitations in the calculated spectrum (see Fig. 8 of Ref. [21] for an illustration of the effect on the spectrum). Moreover, such factorization greatly simplifies the calculation of certain observables, including the r.m.s. radius, electric monopole (\( E0 \)), magnetic dipole (\( M1 \)), and electric quadrupole (\( E2 \)) observables [18].

Here it is important to note that the factorized \( 0s \) center-of-mass wave function thus obtained has an oscillator parameter \( \hbar \omega_{c.m.} = \hbar \omega \) which is determined by the oscillator parameter of the underlying single-particle basis. Equivalently, in terms of oscillator lengths, the \( 0s \) wave function in the center-of-mass coordinate has an \( \hbar \omega \)-dependent oscillator length \( \hbar c.m. = A^{-1/2} \) (see Sec. F.3 of Ref. [18]). Thus, many-body calculations carried out in \( N_{\text{max}} \)-truncated oscillator bases of different \( \hbar \omega \) result in different “spectator” center-of-mass motions. That is, the many-body eigenstates obtained using these different bases may converge towards the same intrinsic structure with increasing \( N_{\text{max}} \), but not the same center-of-mass wave function. This will be important to keep in mind when interpreting the \( \hbar \omega \)-dependence of the natural orbitals thus obtained (as in Sec. III.B below).

If we move beyond the traditional oscillator basis in \( N_{\text{max}} \) truncation, as we must to make use of natural orbitals, we forsake the formal comfort of having a guaranteed exact center-of-mass factorization. However, in practice, an approximate factorization may still be obtained [20, 61–63], either since it naturally emerges in the calculation (as explored for the natural orbital basis in Sec. III.C below) or with some help from a Lawson term. Furthermore, the impact upon observables of any spurious contribution may be mitigated through judicious use of translationally-invariant intrinsic operators [18]. Indeed, alternate choices both for orbitals and for truncation have already been applied in NCCI calculations. For instance, orbitals defined in terms of the Laguerre functions [64–66], a standard set of basis functions in electron-structure theory [15], have been explored [20, 21].

For the many-body truncation, calculations have also been performed using the so-called full configuration interaction (FCI) truncation [15], which simply retains all configurations built by distributing nucleons over the given set of orbitals (this is simply the traditional fermionic many-body space obtained from a given set of single-particle states [67]). In the context of NCCI calculations, the FCI basis is taken as all configurations involving a given set of oscillator shells. However, convergence with respect to the many-body basis size is found to be much slower than for traditional \( N_{\text{max}} \) calculations [68]. More general many-body truncation schemes\(^8\) are also feasible, e.g., in which orbitals are weighted by measures other than the number of oscillator quanta [20] or in which the basis configurations are selected through more sophisticated importance criteria [71].

\[^8\] Here we specifically have in mind truncation schemes for a traditional configuration interaction basis of antisymmetrized product states. It should be noted that symmetry-adapted coupling schemes for NCCI calculations, based on SU(3) [8] or Sp(3, 3) [69] symmetry groups, are subject to truncation schemes of a different nature, as these schemes involve a change of basis, before truncation, to correlated many-body basis states.
Regardless of basis choice, the essential inputs into the construction of the Hamiltonian matrix in the NCCI basis are the two-body matrix elements of this Hamiltonian (assuming the internucleon interaction $V$ is two-body, or three-body matrix elements, if the interaction is three-body, etc.). These must be obtained for the given choice of orbitals. The rest of the Hamiltonian construction follows from the standard treatment of $n$-body operators in second quantization \cite{77}. The eigenproblem is thus cast as a large, sparse matrix diagonalization problem, which is solved numerically using, e.g., the Lanczos algorithm \cite{39} \cite{72}.

One-body densities are readily extracted from the resulting wave functions. These densities are commonly used for the computation of one-body observables, such as matrix elements of electromagnetic operators for moments and transitions \cite{45}, and as inputs to reaction calculations \cite{73}. More precisely, while the electromagnetic operators, taken properly in the center-of-mass frame, involve two-body or higher contributions, they may effectively be replaced by one-body operators when the center-of-mass motion has the harmonic-oscillator 0\textsuperscript{th} form noted above \cite{13} \cite{74}. The scalar densities \cite{9}, in particular, are also the necessary ingredient for deducing natural orbitals appropriate to the NCCI framework (Sec. II A).

Our procedure is thus to carry out an initial NCCI calculation in a traditional $N_{\text{max}}$-truncated oscillator basis. One of the calculated eigenstates, say, the ground state, is taken as the reference state for generating natural orbitals, and the relevant scalar densities are extracted.

To see which oscillator orbitals contribute to the resulting natural orbitals, note that, in an $N_{\text{max}}$-truncated NCCI basis, the configurations involve nucleons reaching orbitals with $N_{\text{max}}$ quanta above the valence shell. The active orbitals thus have $N \leq N_v + N_{\text{max}}$, where $N_v$ is the number of oscillator quanta for the valence shell (e.g., $N_v = 0$ for an "s-shell" nucleus, or $N_v = 1$ for a "p-shell" nucleus). The calculated scalar densities reflect only these active orbitals, and the natural orbitals resulting from diagonalizing the resulting density matrix represent mixtures of only these orbitals, that is, oscillator orbitals of the same $(ij)$ and with $N \leq N_v + N_{\text{max}}$.

The resulting natural orbitals are again labeled by quantum numbers $nlj$, where now the radial quantum number $n$ no longer necessarily reflects the number of nodes in the radial wave function but simply reflects the chosen ordering of natural orbitals by decreasing eigenvalue (i.e., decreasing mean occupation) as discussed above (Sec. II A). For example, consider an $N_{\text{max}} = 4$ calculation for the s-shell nucleus $^4\text{He}$. Within the $s_1/2$, or $(l,j) = (0,1/2)$, subspace, the resulting scalar densities connect the $0s_1/2$, $1s_1/2$, and $2s_1/2$ orbitals ($N = 0, 2, 4$, respectively), and diagonalizing the scalar density matrix thus mixes these orbitals to define natural orbitals $0s_1/2$, $1s_1/2$, and $2s'_1/2$.

It is now straightforward to carry out an NCCI calculation in a new basis, formed from antisymmetrized products of natural orbitals. The same many-body machinery is used as in the original oscillator-basis calculation. It is merely necessary to carry out a change of basis on the two-body matrix elements of the Hamiltonian \cite{9} (see Sec. III C of Ref. \cite{79}). Only a finite set of two-body matrix elements in the oscillator basis are required as input to the transformation, since, as just noted, each natural orbital is obtained from a finite set of underlying oscillator orbitals. Then, evaluation and diagonalization of the many-body Hamiltonian matrix proceed as before.

However, in defining an NCCI calculation in terms of natural orbitals, a fundamental question arises as to how to truncate the many-body basis. The choice may be expected to profoundly affect the results and, in particular, determine how the accuracy obtained from the many-body calculation relates to basis size.

The transformation from oscillator orbitals to natural orbitals is simply a unitary transformation on the single-particle space. More specifically, this transformation is restricted to the low-lying subspace spanned by oscillator states with $N \leq N_v + N_{\text{max}}$. Since the many-body basis consists of antisymmetrized products of the single-particle orbitals, a change of basis on the single-particle space inherently induces a change of basis on the many-body product space.

However, if all antisymmetrized products are retained, as in the FCI truncation, then, while the basis itself may change, the many-body space spanned by this basis is invariant under such a rearrangement of the single-particle space. Thus, an FCI calculation based on the original oscillator orbitals, or on natural orbitals obtained by a unitary transformation of these, yield identical results. No benefit in convergence is achieved. The truncated many-body spaces obtained before and after transformation to natural orbitals only differ when the set of antisymmetrized product states constituting the many-body basis is truncated in a nontrivial fashion, that is, to a proper subspace of the FCI space (as compared in Sec. III A below). The dimension of the $N_{\text{max}}$ truncated space and the enveloping FCI space involving the same orbitals (dotted line) for $^3\text{He}$ may be compared in Fig. 1.

An obvious, though not necessarily optimal, choice of many-body truncation scheme, as adopted here, is to simply carry over the formal structure of the $N_{\text{max}}$ truncation. The natural orbitals are already identified by $nlj$ labels, where, again, $n$ reflects the chosen ordering by decreasing occupation in the reference state. For each of these orbitals, we may simply define a weighting label $N = 2n + l$ (as in Refs. \cite{20} \cite{37}), although this label no longer has any direct meaning in terms of oscillator quanta. We then proceed as before, by treating this label as an additive quantity, thereby defining $N = \sum_{i=1}^{A} N_i$.
for a many-body configuration, and imposing a nominal \(N_{\text{max}}\) truncation on the configurations. This truncation no longer has any direct connection to the oscillator excitation quanta in the system, nor does it guarantee exact center-of-mass separability. However, conveniently for purposes of comparison, the dimension of the problem is exactly as it was for the original \(N_{\text{max}}\)-truncated oscillator basis (Fig. 1).

III. ILLUSTRATION OF NATURAL ORBITALS IN NCCI CALCULATIONS: \(^3\)He

A. Convergence of observables

To see how the formalism just elaborated (Sec. II) is reflected in actual NCCI calculations, let us now examine the convergence of observables in illustrative NCCI calculations, making use of symmetry-adapted natural orbitals. Here we take \(^3\)He as the simplest nontrivial example. The comparatively slow growth of dimension with \(N_{\text{max}}\) for this nuclide (Fig. 1) means that essentially converged results can be obtained, as a reference against which the convergence of lower-\(N_{\text{max}}\) results can be compared.

Results for the ground state energy eigenvalue of \(^3\)He are shown in Fig. 2 first as obtained in the oscillator basis [Fig. 2(a)], then as obtained in the natural orbital basis [Fig. 2(b)]. For these illustrations, we take the Daejeon16 internucleon interaction \(^{10}\), which is based on the two-body part of the Entem-Machleidt (EM) N^3LO chiral EFT interaction \(^{12}\), subsequently softened via a similarity renormalization group (SRG) transformation \(^{77}\) to enhance convergence and then adjusted via a phase-shift equivalent transformation to provide better description of nuclei with \(A \leq 16\). Calculations are obtained using the many-body code MFDn \(^{78, 79}\), along with codes for the transformation of two-body matrix elements from the oscillator basis to the natural-orbital basis \(^{80}\), and no Lawson term [see \(^{5}\)] is included in the Hamiltonian for the calculations in the natural-orbital basis. We also show the experimental binding energy \(^{76}\) for comparison.

The eigenvalues obtained in the oscillator-basis calculations [Fig. 2(a)] follow a familiar convergence pattern (e.g., Refs. \(^{58, 77}\)). Each curve represents calculations at fixed \(N_{\text{max}}\) (from 8 to 16), for varying \(h\omega\), and has a variational minimum with respect to \(h\omega\), which arises in the vicinity of \(h\omega = 12.5\text{ MeV}\) for this particular nuclide, state, and interaction. Increasing \(N_{\text{max}}\), at given \(h\omega\), strictly expands the space in which the calculation is carried out, and is thus guaranteed by the variational principle to monotonically lower the ground state energy. Convergence towards the true eigenvalue, as would be obtained in the full, untruncated many-body space, is signalled by insensitivity to the basis truncation \(N_{\text{max}}\) (compression of successive curves), as well as local insensitivity to the oscillator parameter \(h\omega\) (flattening of the curves). For the ground state energy, this manifests as compression of the curves against a variational floor.

For each of these oscillator-basis calculations, the resulting one-body densities yield a set of approximate natural orbitals, which define the natural orbital basis, which we then use for a subsequent many-body calculation, as outlined in Sec. IIIB. For the resulting energies [Fig. 2(b)], each curve again represents calculations at fixed \(N_{\text{max}}\), now in the sense of the nominal \(N_{\text{max}}\) truncation scheme for natural orbitals (Sec. IIIB).

Comparing the overall shapes of the curves, of \(E\) vs. \(h\omega\), in Fig. 2, we may observe that the natural-orbital basis provides an overall flattening, or reduced depen-
decreases on $\hbar \omega$, in the vicinity of the variational minimum. However, for a more direct quantitative comparison of the results obtained with the two bases, the approximately exponential nature of the convergence with $N_{\text{max}}$ means that comparison can be carried out more readily on a logarithmic scale. To provide a meaningful zero point for the logarithmic scale, we must take the residual with respect to a “converged” reference value $E_{\text{ref}}$, which we obtain from higher-$N_{\text{max}}$ calculations (for $N_{\text{max}} \approx 24$, the energy in the vicinity of the variational minimum is converged to the keV scale). The energies, thus recast as residuals, are shown on a logarithmic scale in Fig. 2(c), for the results obtained both with the oscillator (filled circles) and natural-orbital (open squares) bases. To provide clear separation in the plot, only the $N_{\text{max}} = 8, 14, 16$ results are shown.

At lower $N_{\text{max}}$, as exemplified by the $N_{\text{max}} = 8$ results (dotted lines) in Fig. 2(c), there is little distinction between the results obtained in oscillator and natural-orbital bases. This is perhaps to be expected. In the limit of $N_{\text{max}} = 0$, the bases for the oscillator and subsequent natural-orbital calculations are strictly identical. More generally, a low-$N_{\text{max}}$ underlying oscillator calculation provides little opportunity for high-$N$ orbitals to appear in the densities and thus natural orbitals.

At higher $N_{\text{max}}$, as exemplified by the $N_{\text{max}} = 14$ and 16 results (dashed and solid lines, respectively) in Fig. 2(c), one way of comparing the results is to measure the advance obtained by the transformation to natural orbitals in terms of the equivalent increase in $N_{\text{max}}$ required with a traditional oscillator basis to achieve the same advance. In this sense, for calculations in the vicinity of the variational minimum, the energies obtained with natural orbitals are approximately “one step” in $N_{\text{max}}$ ahead of those obtained with oscillator orbitals. Away from the variational minimum, the advantage provided by the natural orbitals is more marked, reflecting the comparative $\hbar \omega$-independence already noted for these calculations in the natural-orbital basis.

Alternatively, we may assess the results of the change of basis in terms of the fraction by which it reduces the residual, i.e., how far it brings us towards the true value which would be obtained in the full, untruncated space. On a logarithmic scale, a given downward vertical shift represents a given fractional reduction. Comparing the $N_{\text{max}} = 16$ results obtained in the two bases, we may observe an approximately uniform downward shift, across the range of $\hbar \omega$, representing a reduction in the residual by a factor of $\sim 3$ (a somewhat greater reduction is attained with the natural-orbital basis for $\hbar \omega \approx 10 \text{ MeV}$).

However, there is an obvious bound on the improvement which may be expected from the transformation to the natural-orbital basis derived from an $N_{\text{max}}$-truncated oscillator basis calculation. Recall that the active orbitals in the oscillator-basis calculation and the subsequent natural orbitals span the same single-particle space. Both the $N_{\text{max}}$-truncated oscillator basis and the nominally $N_{\text{max}}$-truncated constructed from the ensuing natural orbitals span subspaces of the same enveloping FCI space defined by those orbitals (Sec. II B). This FCI space is, in general, of much higher dimension [68]. E.g., for an $N_{\text{max}} = 10$ calculation for $^3\text{He}$, which has dimension $2.6 \times 10^4$, the FCI space consists of all product states involving orbitals through the $N = 10$ oscillator shell, which has the substantially larger dimension $2.8 \times 10^6$ (Fig. 1). We might hope that the $N_{\text{max}}$-truncated natural orbital basis might allow us to reach comparable accuracy in a much smaller space, but it cannot access any components of the true wave function which lie outside of the FCI truncated space.

For the ground state energy, in particular, the result in the FCI space provides a variational lower bound on the results in the subspaces. Thus, it is informative to compare the improvement obtained with natural orbitals to the maximum improvement which could be obtained in the enveloping FCI space. The calculated $^3\text{He}$ ground state energies obtained in the oscillator basis (filled circles) and natural-orbital basis (open squares) are compared with the variational bound provided by the enveloping FCI space (filled triangles) in Fig. 2. Here again, as in Fig. 2(c), values are shown as residuals relative to the true energy, on a logarithmic scale.

At low $N_{\text{max}}$, as exemplified by the $N_{\text{max}} = 4$ results (dotted lines), a factor of $\sim 3$ improvement is possible within the FCI space, near the variational minimum and over most of the $\hbar \omega$ range shown. Yet, as already noted, the transformation to natural orbitals conveys negligible benefit, at least within the nominal $N_{\text{max}}$ truncation scheme.

At higher $N_{\text{max}}$, as exemplified by the $N_{\text{max}} = 10$ results shown in Fig. 2(c), the advantage provided by the transformation to natural orbitals is more marked, reflecting the comparative $\hbar \omega$-independence already noted for these calculations in the natural-orbital basis.
The calculated results for $r_p$ are shown in Fig. 4, where the values obtained with the oscillator (filled circles) and natural-orbital (open squares) bases are overlaid. An approach to convergence is signaled by the “shouldering” of the curves, to form a region of local $\hbar \omega$-independence (flattening) and compression of curves for successive $N_{\text{max}}$ against each other. The value for $r_p$ deduced from the experimental $r_c$ \cite{83} is shown for comparison (filled diamond).

The oscillator-basis calculations for the radius are already atypically well-converged for $^3\text{He}$ (compare, e.g., Sec. IV below). Note the highly expanded vertical scale in Fig. 4 (on the scale of 0.1 fm overall). For the underlying oscillator calculations (filled circles), the various curves for different $N_{\text{max}}$ (from 10 to 16) cross in the vicinity of $\hbar \omega = 10 \text{MeV}$. (Such crossings have been suggested, purely heuristically, as a means of estimating the true radius as it would be obtained in the full, untruncated space \cite{77, 80, 81}, though in practice this prescription must be treated with caution \cite{21}.)

The subsequent calculations in the natural-orbital basis (open squares) do not share such a sharply-defined crossing point. Rather, they more clearly demonstrate the traditional hallmarks of convergence, namely, flattening and compression of the curves. For instance, the $N_{\text{max}} = 16$ curve varies by $\lesssim 0.04$ fm over the range of $\hbar \omega$ from 10 MeV to 20 MeV, while the $N_{\text{max}} = 14$ and 16 curves differ by less than $\lesssim 0.01$ fm over this same range. As a consequence of this flattening, by the high end of the $\hbar \omega$ range shown ($\hbar \omega = 25 \text{MeV}$), the calculations in the natural-orbital basis lie two steps in $N_{\text{max}}$ “ahead” of the calculation in the oscillator basis. The question, of course, is how this difference in convergence behavior actually aids in the problem of direct interest in less well-converged cases, which is to accurately estimate the true value of the observable, as it would be found in the full, untruncated space.

### B. Natural orbitals

Let us now examine the natural orbitals obtained (and subsequently used) in the present $^3\text{He}$ calculations, with the aim of understanding their dependence on the underlying oscillator calculation and thereby also of gaining some insight into their influence on the convergence of observables in the many-body calculation. Recall that the natural orbitals in these many-body calculations are approximations to the “true” natural orbitals for the $^3\text{He}$ ground state, since they are deduced from the approximate $^3\text{He}$ ground state densities obtained in finite, truncated oscillator-basis NCCI calculations (Sec. II.B). The densities, and thus the resulting natural orbitals, depend upon both the $N_{\text{max}}$ and $\hbar \omega$ of the underlying oscillator-basis calculation.

Recall, furthermore, that the symmetry-adapted natural orbitals (Sec. II.A) appropriate to NCCI calculations preserve the $l$ and $j$ quantum numbers, changing only...
The radial wave function, by “mixing” underlying oscillator orbitals of different \( n \) within an \( l_j \) space. We focus first on the \( 0s_{1/2} \) orbital, as this is the notionally “occupied” orbital in a simple shell-model picture, and is indeed still the most heavily-occupied orbital in the actual NCCI calculations. We then explore the properties of the notionally “unoccupied” excited orbitals. While the occupations of these excited (or notionally unoccupied) orbitals are comparatively small, it is these orbitals which drive the convergence of the many-body calculation in a natural-orbital basis.

The radial wave function for the \( 0s_{1/2} \) natural orbital for protons, in particular, is shown in Fig. 5 where its dependence on the \( N_{\text{max}} \) and \( \hbar \omega \) of the underlying oscillator calculation is mapped out. (The behavior for the neutron \( 0s_{1/2} \) orbital is qualitatively similar.) Here, the radial wave function is plotted as the radial probability density \( P(r) = r^2 |\psi(r)|^2 \), from \( N_{\text{max}} = 2 \) (dotted lines) to \( N_{\text{max}} = 16 \) (solid lines), separately for \( \hbar \omega = 9 \text{ MeV} \) [Fig. 5(a)], 15 MeV [Fig. 5(b)], and 25 MeV [Fig. 5(c)]. The 0s radial function for the underlying oscillator basis is also shown for comparison (thick gray line). Note that the natural orbital obtained from an \( N_{\text{max}} = 0 \) oscillator calculation is still simply this oscillator function, as the resulting densities do not mix the fully-occupied \( s \)-shell orbitals with the fully-unoccupied higher orbitals.

The densities, and thus the resulting natural orbitals, are expected to eventually converge with increasing \( N_{\text{max}} \). Such is indeed seen in Fig. 5 if we examine the curves within a given panel, \( i.e., \) obtained for different \( N_{\text{max}} \) but at a given choice of \( \hbar \omega \). On this scale, the shape of the radial wave function appears to change comparatively little for \( N_{\text{max}} \) beyond about 4 or 6.

The \( \hbar \omega \) dependence is more subtle. All observables (energies, electromagnetic matrix elements, radii, etc.)
FIG. 7. Radial wave functions for the $^3$He $s$-, $p$-, and $sd$-shell natural orbitals, for both protons (short dashed lines) and neutrons (long dashed lines), plotted as the radial probability density $P(r)$. These are obtained from the underlying oscillator-basis calculation near the variational minimum ($\hbar\omega = 15$ MeV) and at high $N_{\text{max}}$ ($N_{\text{max}} = 16$). The corresponding oscillator radial functions for $\hbar\omega = 15$ MeV (thick gray lines) are shown for comparison. The mean occupation $n_a$ for each natural orbital, from the corresponding eigenvalue of the scalar density matrix, is indicated by the filling of the bar at top right (upper bar for protons, lower bar for neutrons).

obtained from the densities retain some $\hbar\omega$ dependence at finite $N_{\text{max}}$ due to their sensitivity to the $N_{\text{max}}$- and $\hbar\omega$-dependent intrinsic structure of the approximate $^3$He ground state obtained in a truncated oscillator calculation. At finite $N_{\text{max}}$, some $\hbar\omega$ dependence of the natural orbitals may similarly be expected to arise from such sensitivity to the $N_{\text{max}}$- and $\hbar\omega$-dependence of the calculated intrinsic structure. This dependence is expected to ultimately disappear with increasing $N_{\text{max}}$, as the intrinsic structure converges.

However, recall (Sec. II B) that even in the large $N_{\text{max}}$ limit the natural orbitals for the NCCI problem are not uniquely defined. Rather, they may be expected to have an inherent $\hbar\omega$-dependence arising from the center-of-mass zero-point motion of the reference many-body state, which varies with the $\hbar\omega$ of the underlying oscillator basis. Thus, it should not be surprising that, even at high $N_{\text{max}}$, the natural orbitals obtained from underlying oscillator-basis calculations with different $\hbar\omega$ do not coincide. Compare the solid curves in the different panels of Fig. 5. These clearly do not coincide, with the location of the maximum moving to smaller radius with increasing $\hbar\omega$.

To characterize how the radial wave functions for the natural orbitals at high $N_{\text{max}}$ (solid lines) differ qualitatively from those of the underlying oscillator functions (thick gray lines), in Fig. 5, we shall find it convenient to separately consider the central region of the wave function and its large-radius tail (porous though this distinction may be). Let us first consider the central region, that is, around the peak in the wave function. For the $0s_{1/2}$ natural orbital obtained from the reference wave function calculated in an $\hbar\omega = 15$ MeV oscillator basis [Fig. 5(b)], there is little apparent change going from the underlying oscillator function to the natural orbital. For the natural orbital obtained in an $\hbar\omega = 9$ MeV oscillator basis [Fig. 5(a)], which has a longer oscillator length [recall $b \propto (\hbar\omega)^{-1/2}$], the peak shifts inward, to lower radius, relative to the underlying oscillator function, though not all the way to the peak location for $\hbar\omega = 15$ MeV [Fig. 5(b)]. Alternatively, for the natural orbital obtained in an $\hbar\omega = 25$ MeV oscillator basis, which has a shorter oscillator length [Fig. 5(c)], the peak shifts outward, to larger radius, relative to the underly-
ing oscillator function, though again not all the way to the peak location for \( h\omega = 15 \text{ MeV} \).

Either way, a portion of the effect of transforming from the underlying oscillator basis to natural orbitals is to “dilate” the radial function to more closely resemble a 0s oscillator function of \( h\omega \approx 15 \text{ MeV} \). The effect is to moderate the change in characteristic length scale for the natural orbitals, as the \( h\omega \) for the underlying oscillator basis is varied, as compared to the change in oscillator length for the underlying oscillator orbitals themselves. This reduced \( h\omega \) dependence of the orbitals (at least in the central region) presumably contributes to the reduction in \( h\omega \) dependence found for the observables in the calculations based on the natural-orbital basis (Sec. IIIA).

A simple and intuitive explanation for this behavior of the orbitals is that the natural orbitals are the result of a compromise between the intrinsic structure and center-of-mass motion embodied within the reference wave function. The intrinsic structure is described well by nucleons occupying orbitals resembling an \( h\omega = 15 \text{ MeV} \) 0s\(_{1/2}\) oscillator orbital, regardless of underlying oscillator basis. But the center-of-mass motion of the reference wave function for \( h\omega = 9 \text{ MeV} \) is well described by nucleons in an \( h\omega = 9 \text{ MeV} \) 0s\(_{1/2}\) oscillator orbital. The resulting 0s\(_{1/2}\) natural orbital lies somewhere inbetween. Similarly, the center-of-mass motion of the reference wave function for \( h\omega = 25 \text{ MeV} \) is well described by nucleons in an \( h\omega = 25 \text{ MeV} \) 0s\(_{1/2}\) oscillator orbital, and the resulting 0s\(_{1/2}\) natural orbital lies somewhere inbetween.

Turning now to the tail region of the orbital, the natural question is the extent to which the natural orbitals take on the exponential asymptotics anticipated from the mean-field description of the nucleus. Recall that these asymptotics are expected to be particularly important for the convergence of long-distance observables (Sec. IIIA).

The asymptotic behavior is more readily apparent if we replot the radial probability densities on a logarithmic scale, as in Fig. 6. A tail with exponential asymptotics appears as a straight line on such a plot, while a tail with the Gaussian asymptotics characteristic of the oscillator functions appears as downward-curving parabola, as seen for the underlying oscillator radial functions (grey lines). We may observe that the tail “grows in”, with the inclusion of additional oscillator functions, so that exponential asymptotics (i.e., straight-line falloff on the log plot) are gradually established, extending to larger radii with increasing \( N_{\text{max}} \). (One may compare to Fig. 4 of Ref. [11], for a classic illustration of an exponential tail growing in for a Hartree-Fock orbital, or to Fig. 1 of Ref. [21], for the schematic example of a Woods-Saxon orbital expanded in an oscillator basis [15].) The emergence of exponential asymptotics is most clearly visible for the \( h\omega = 15 \text{ MeV} \) natural orbitals [Fig. 6(b)], where the progression from the underlying oscillator orbital to the true, high-\( N_{\text{max}} \) natural orbital is not complicated by a significant radial shift in the peak location.

NCCI calculations for \(^3\text{He}\) in a natural-orbital basis involve, of course, not just the notionally occupied 0s\(_{1/2}\) orbital, but also basis configurations incorporating the higher, notionally unoccupied, natural orbitals, as well. Some of the low-lying natural orbitals are shown in Fig. 7 for both protons (short dashed lines) and neutrons (long dashed lines). Here we follow the analogy to an oscillator basis, by focusing on natural orbitals with \( nlj \) quantum numbers corresponding to the traditional \( N = 0 \) (s), 1 (p), and 2 (sd) oscillator shells. We focus on the natural orbitals obtained from the \( h\omega = 15 \text{ MeV} \) oscillator-basis calculation at \( N_{\text{max}} = 16 \), so that the proton 0s\(_{1/2}\) orbital here corresponds to the highest-\( N_{\text{max}} \) case shown in Fig. 5(b). Again, the underlying oscillator orbital is shown for comparison (thick gray lines).

Let us first consider the “occupations” \( n \) of these orbitals in the reference wave function, which we know from the corresponding eigenvalues of the scalar density matrix (Sec. IIIA). (Such occupations provide only an estimate of the occupation in any subsequent many-body calculation using the natural-orbital basis.) The occupations are shown graphically at the top of each panel in Fig. 7 but at this scale are indistinguishable from those of the traditional shell model description (in which \( n_{0s_{1/2}} = 1 \) for the neutrons, \( n_{0s_{1/2}} = 2 \) for the protons, and all other orbitals are unoccupied). More precisely, for the present illustrative calculation, we have \( n_{0s_{1/2}} \approx 0.96 \) for the neutrons and \( n_{0s_{1/2}} \approx 1.92 \) for the protons. The next most occupied orbitals are the p-shell orbitals and the 1s\(_{1/2}\) orbital of the sd shell, with mean occupations of \( \sim 10^{-2} \), while occupations fall off towards \( \sim 10^{-3} \) and below for higher orbitals.

Overall, in Fig. 7 the general impression is that the natural orbitals simply “tweak” the oscillator radial functions, with modest shifts to the peak location and overall shape (again, a linear scale does not do justice to changes in the asymptotics). The difference in proton and neutron structure in the reference many-body calculation for \(^3\text{He}\) is manifest in the differences between corresponding proton and neutron natural orbitals. The distinction is most striking for the proton 0p\(_{1/2}\) orbital, which is shifted to markedly larger radii than the corresponding neutron orbital (which remains close to the underlying oscillator function). In general, the proton radial functions develop more pronounced tails than the neutron orbitals, visible even on a linear scale, suggestive of Coulomb repulsion effects.

In atomic and molecular electron structure theory, it is recognized that an important characteristic of the natural orbitals, including the unoccupied orbitals, is their tendency to remain localized in the region of high particle density [23]. This is to be contrasted with the unoccupied (virtual) Hartree-Fock orbitals, which instead provide an expansion of the continuum.

It is thus worth elaborating on an essential difference between natural orbitals and Hartree-Fock orbitals (e.g., Ref. [42]). The unoccupied natural orbitals are well-defined, from the densities of the reference many-body calculation. In contrast, the basic variational condition for the Hartree-Fock ground state focuses entirely
on optimizing the occupied orbitals, so as to minimize the energy in a single Slater determinant. The unoccupied orbitals are entirely unconstrained by this variational condition (except insofar as they must span an orthogonal complement to the occupied orbitals). The iterative calculational procedure for obtaining Hartree-Fock orbitals introduces a single-particle eigenproblem (involving Hartree and exchange potentials), intended to yield the occupied orbitals. While the set of solutions can be extended to provide a definition (one particular choice) for the unoccupied Hartree-Fock orbitals, it is not at all obvious that these unoccupied Hartree-Fock orbitals should be particularly well-suited for efficiently expanding the many-body wave functions in a configuration-interaction basis.

C. Center-of-mass factorization

A factorized and well-controlled center-of-mass motion is important, as discussed in Sec. [113] if the results of the many-body calculation are to be of practical use, beyond limited calculations for ground-state observables. Recall that the $N_{\text{max}}$-truncated oscillator basis is special, in that the many-body wave functions resulting from NCCI calculations with this basis factorize into intrinsic and center-of-mass parts, and the center-of-mass part can be selected to have pure oscillator $0s$ zero-point motion in the center-of-mass coordinate. Such exact factorization is no longer guaranteed, and no longer to be expected, if we move away from the $N_{\text{max}}$-truncated oscillator basis. However, approximate factorization may arise, with or without the persuasion of a Lawson term in the Hamiltonian. Let us therefore diagnose the center-of-mass motion which arises in our present calculations with the natural-orbital basis, and how it depends upon the choice of underlying oscillator basis.

For many purposes, we might be satisfied by factorization involving an arbitrary center-of-mass wave function. For instance, angular momentum selection rules which allow the intrinsic electromagnetic operators to be replaced with one-body operators in practical calculations [112] require factorization with an $s$-wave center-of-mass wave function, as $|\Psi_J\rangle = |\Psi_{J}^{\text{intr}}\rangle \times |\Psi_{0s}^{\text{c.m.}}\rangle$, but not specifically an oscillator $0s$ wave function. However, in practice, we do not have a good way to measure how well a many-body wave function factorizes, unless the factorization specifically involves $0s$ harmonic-oscillator motion.

Specifically, the expectation value of the center-of-mass number operator $N_{\text{c.m.}}$ allows us to measure deviations from pure $0s$ center-of-mass motion [20 61 105]. Such $0s$ center-of-mass motion then incidentally implies factorization as $|\Psi_J\rangle = |\Psi_{J}^{\text{intr}}\rangle \times |\Psi_{0s}^{\text{c.m.}}\rangle$. The definition of a center-of-mass harmonic-oscillator number operator depends upon the oscillator parameter $\hbar \omega_{\text{c.m.}}$, taken for the center-of-mass motion:

$$N_{\text{c.m.}}(\hbar \omega_{\text{c.m.}}) = c_{\text{c.m.}}^\dagger \cdot c_{\text{c.m.}}$$

$= \frac{1}{2} (\hbar \omega_{\text{c.m.}})^{-1} (\hbar c)^2 A(m_{N} c^2) K^2$

$+ \frac{1}{2} (\hbar \omega_{\text{c.m.}}) A(m_{N} c^2) R^2 - \frac{3}{2},$  \hspace{1cm} (9)

where $c_{\text{c.m.}}^\dagger$ and $c_{\text{c.m.}}$ are the center-of-mass ladder operators (see Sec. F.3 of Ref. [113] for definitions), $K^2 = |k_{\text{c.m.}}|^2$ is the squared magnitude of the center-of-mass momentum vector or, more precisely, wave vector, where $p_{\text{c.m.}} = \hbar k_{\text{c.m.}}$, and $R^2 = |x_{\text{c.m.}}|^2$ is the squared magni-
FIG. 9. Dependence of the approximate 0 \text{s} center-of-mass motion of the calculated \(^{4}\text{He}\) ground state (and its degree of contamination) on the \(h\omega\) of the underlying oscillator basis, in calculations with a natural-orbital basis, as measured by (a) \(\tilde{h}\omega_{\text{c.m.}}\) and (b) \(\tilde{N}_{\text{c.m.}}\). Results are shown for calculations with \(N_{\text{max}} = 4\) (dotted lines) through \(N_{\text{max}} = 16\) (solid lines, highlighted).

Figure 9 displays the dependence of the approximate \(0\) \text{s} center-of-mass motion on \(h\omega\) for the \(^{4}\text{He}\) ground state wave function, calculated in a natural-orbital basis. The motion is quantified by the expectation values of the position operators, \(\langle \tilde{r}_{j} \rangle\), and the corresponding minimized measure of the center-of-mass motion, \(\tilde{N}_{\text{c.m.}} \equiv \langle N_{\text{c.m.}} (\tilde{\omega}_{\text{c.m.}}) \rangle\), is given by

\[
\tilde{N}_{\text{c.m.}} = \langle (K^2) (R^2) \rangle^{1/2} - \frac{3}{2},
\]

with \(\langle K^2 \rangle\) and \(\langle R^2 \rangle\) calculated in a natural-orbital basis, where \(\tilde{\omega}_{\text{c.m.}}\) represents the center-of-mass oscillation of the \(^{4}\text{He}\) ground state wave function, calculated in a NCCI many-body code, for \(N_{\text{max}} = 4\) (dotted line) through \(N_{\text{max}} = 16\) (solid line). The value of \(\tilde{N}_{\text{c.m.}}\) is given by

\[
\tilde{N}_{\text{c.m.}} = \langle N_{\text{c.m.}} (\tilde{\omega}_{\text{c.m.}}) \rangle.
\]

With this in mind, let us now examine the center-of-mass motion for the \(^{3}\text{He}\) ground state wave functions obtained in a natural-orbital basis. The values of \(\langle N_{\text{c.m.}} (\tilde{\omega}_{\text{c.m.}}) \rangle\), as we sweep \(\tilde{\omega}_{\text{c.m.}}\) in \([9]\), are shown in Fig. 8. Each curve is simply determined analytically, by \([9]\), taking the calculated \((K^2)\) and \((R^2)\) for the corresponding wave function. We again (as in Fig. 5) take \(h\omega = 9 \text{ MeV}\) [Fig. 8(a)], 15 MeV [Fig. 8(b)], and 25 MeV [Fig. 8(c)] as representative values for the oscillator parameter of the underlying oscillator basis (namely, below, near, and above the variational energy minimum, respectively).

For \(N_{\text{max}} = 0\) (thick gray line), in Fig. 8 recall that the natural orbitals are simply the original oscillator orbitals, with oscillator parameter \(h\omega\), and calculations in the natural-orbital basis are simply calculations in the oscillator basis. The center-of-mass motion is thus pure \(0\) \text{s} motion, with \(\tilde{\omega}_{\text{c.m.}} = h\omega\) (vertical dotted line), for which \(\tilde{N}_{\text{c.m.}} = 0\). (In fact, curves identical to that shown would be obtained for any of the \(N_{\text{max}}\)-truncated oscillator-basis calculations with this same \(h\omega\).)

Then, for the calculations in a natural-orbital basis proper, with \(N_{\text{max}} = 4\) (dotted line) through \(N_{\text{max}} = 16\) (solid line), in Fig. 8 there is no \(\tilde{\omega}_{\text{c.m.}}\) for which \(\langle N_{\text{c.m.}} \rangle\) vanishes. Rather, the location of the minimum in \(\langle N_{\text{c.m.}} \rangle\), given by \([10]\) and \([11]\), is marked by a dot.

For \(h\omega = 9 \text{ MeV}\) [Figs. 8(a)], there is an initial discontinuity going from the oscillator basis to a natural-orbital basis, where \(\tilde{N}_{\text{c.m.}}\) jumps to \(\gtrsim 10^{-2}\) for \(N_{\text{max}} = 4\), then steadily decreases again, converging to a value \(\sim 10^{-3}\).

The optimal \(\tilde{\omega}_{\text{c.m.}}\) for recognizing this approximate factorization is \(\tilde{\omega}_{\text{c.m.}} \approx 9.4 \text{ MeV}\), slightly above the \(h\omega\) of the underlying oscillator basis (\(h\omega = 9 \text{ MeV}\)).
Moving to the other side of the variational minimum in \( h\omega \), for \( h\omega = 25\text{ MeV} \) [Figs. 8(c)], there is again an initial discontinuity, with \( N_{\text{c.m.}} \sim 10^{-2} \) for \( N_{\text{max}} = 4 \), and converging towards \( \lesssim 10^{-3} \). Here the optimal \( h\omega_{\text{c.m.}} \) for recognizing this approximate factorization is \( h\omega_{\text{c.m.}} \approx 19\text{ MeV} \), notably below the \( h\omega \) of the underlying oscillator basis (\( h\omega = 25\text{ MeV} \)).

Finally, for \( h\omega = 15\text{ MeV} \) [Figs. 8(b)], near the variational energy minimum, after \( N_{\text{c.m.}} \) initially jumps to \( N_{\text{c.m.}} \lesssim 10^{-2} \) for \( N_{\text{max}} = 4 \), it then immediately drops to \( N_{\text{c.m.}} \approx 10^{-3} \) for higher \( N_{\text{max}} \). The location of the minimum drifts slightly downward, from the \( h\omega \) of the underlying oscillator basis (\( h\omega = 15\text{ MeV} \)), toward \( h\omega_{\text{c.m.}} \approx 12.5\text{ MeV} \).

Thus, in each case, regardless of the \( h\omega \) for the underlying oscillator basis, a reasonably pure \( 0s \) center-of-mass motion spontaneously emerges for the \(^6\text{He} \) ground state, as recognized when the appropriate choice \( h\omega_{\text{c.m.}} \) of oscillator parameter is used in measuring the center-of-mass motion, implying also a high degree of center-of-mass factorization. Furthermore, in each case, this \( h\omega_{\text{c.m.}} \) for which \( 0s \) motion is most closely realized differs from the \( h\omega \) of the underlying oscillator basis.

To more systematically map out the behaviors we have just seen, \( h\omega_{\text{c.m.}} \) and \( N_{\text{c.m.}} \) are shown as functions of the underlying oscillator basis \( h\omega \) in Fig. 9. For large \( N_{\text{max}} \), the dependence of \( h\omega_{\text{c.m.}} \) on \( h\omega \) [Fig. 9(a)] is nearly linear, but of shallower slope than the reference line \( h\omega_{\text{c.m.}} = h\omega \) (dashed diagonal line). The oscillator parameter \( h\omega_{\text{c.m.}} \) for the center-of-mass motion matches that of the underlying oscillator basis for the natural orbitals in the vicinity of \( h\omega = 10\text{ MeV} \) to \( 12.5\text{ MeV} \). In this range of \( h\omega \), at high \( N_{\text{max}} \), one also observes that the purest \( 0s \) center-of-mass motion is obtained [Fig. 9(b)], with \( N_{\text{c.m.}} \lesssim 10^{-4} \).

A rough intuitive understanding of the center-of-mass motion, in particular, the behavior of the preferred \( h\omega_{\text{c.m.}} \) observed in Fig. 9(a), follows from the \( h\omega \)-dependence noted above for the natural orbitals themselves (Sec. II B). Recall the tendency, observed in Fig. 5 for natural orbitals obtained from a low-\( h\omega \) underlying oscillator basis [Fig. 5(a)] to still resemble oscillator orbitals, but of a somewhat higher \( h\omega \), closer to \( h\omega \approx 15\text{ MeV} \), and for natural orbitals obtained from a high-\( h\omega \) underlying oscillator basis [Fig. 5(c)] to resemble oscillator orbitals of a lower \( h\omega \), again closer to \( h\omega \approx 15\text{ MeV} \). To the extent that the low-lying natural orbitals resemble oscillator orbitals of some \( h\omega \), then a (nominally \( N_{\text{max}} \)-truncated) calculation in such a natural-orbital basis may be expected to have similar properties to an \( N_{\text{max}} \)-truncated calculation in an oscillator basis of this same \( h\omega \). It is thus perhaps not surprising that \( h\omega_{\text{c.m.}} \) of the center-of-mass function follows the same overall trend as the “effective” \( h\omega \) of the natural orbitals.

However, we must always keep in mind that \( \langle N_{\text{c.m.}} \rangle \) is, strictly, only a measure of center-of-mass contamination, relative to harmonic-oscillator \( 0s \) motion, and therefore only incidentally provides an upper bound on the break-down of center-of-mass factorization. Nonzero \( \langle N_{\text{c.m.}} \rangle \) could reflect that factorization is broken, but it could also simply mean that we have factorization which is of a more difficult form to recognize, since the center-of-mass motion is not simply described by a \( 0s \) oscillator wave function.

Further, for the present many-body calculations in the natural-orbital basis, recall that we have included no Lawson center-of-mass term (Sec. II) in the Hamiltonian. For now, we are thus identifying the center-of-mass motion which emerges spontaneously when we diagonalize a translationally-invariant intrinsic Hamiltonian, restricted to the particular truncated many-body space of these calculations. Starting from this baseline, one may then explore the effect of including a Lawson term, which is expected to refine the center-of-mass motion, at some cost to the convergence of the intrinsic motion (see Ref. 35 for initial examples of such calculations). Here one might more naturally choose an \( h\omega_{\text{c.m.}} \) parameter for the Lawson term which reinforces the center-of-mass motion as it already spontaneously emerges in the natural-orbital basis (\( h\omega_{\text{c.m.}} = h\omega_{\text{c.m.}} \)) rather than simply matching the oscillator parameter the underlying oscillator basis (\( h\omega_{\text{c.m.}} = h\omega \)).

IV. NATURAL ORBITALS AND HALO STRUCTURE: \(^6\text{He} \)

A. Convergence of observables

For a halo nucleus, such as \(^6\text{He} \), the connection between natural orbitals and the single-particle structure of the nucleus should be particularly revealing. The nat-
atural orbitals occupied by halo nucleons may be expected to reflect the large-distance behavior which generates the halo. The ground state of $^6$He is understood to be characterized, consisting of a $^4$He (or $\alpha$) core plus two weakly-bound neutrons. This leads to a spatially-extended neutron distribution, with possible correlations in the motion of the halo neutrons. The weak binding is reflected in a small two-neutron separation energy ($\approx 0.97$ MeV), while the extended spatial structure is reflected in a marked increase in r.m.s. radius observables from $^4$He to $^6$He. Having already explored the basic properties of NCCI calculations in a natural-orbital basis for $^3$He (Sec. III), we will take these as a baseline for comparison for $^6$He. Let us first consider the calculated energy and radius observables for $^6$He, with a natural-orbital basis, then (in the subsequent Sec. III B below) the radial wave functions of the orbitals themselves.

The $^6$He ground state energy is shown in Fig. 10, as calculated with oscillator (solid circles) and natural-orbital (open squares) bases. Here we consider truncations through $N_{\text{max}} = 14$, again with the Daejeon16 interaction. The experimental binding energy is shown for comparison (filled diamond).

The energy obtained with natural orbitals, in Fig. 10 is consistently lower than that obtained in the underlying oscillator basis, and is thus, by the variational principle, closer to the true energy in the full many-body space. In the vicinity of the variational minimum, the energy obtained with natural orbitals is approximately “one step” in $N_{\text{max}}$ ahead of that obtained with oscillator orbitals. This relation strictly holds for the highest $N_{\text{max}}$ shown in Fig. 10 (i.e., the energy obtained in the natural-orbital basis for $N_{\text{max}} = 12$ lies below that obtained in the oscillator basis for $N_{\text{max}} = 14$). The $h\omega$ dependence of the calculated energy is, again, much reduced in the natural-orbital basis, so the improvement of the natural-orbital results over the oscillator-basis results becomes more marked as we move away from the variational energy minimum and towards the extreme values of $h\omega$ shown in Fig. 10.

Whereas for $^3$He we could benchmark the calculated energies against an effectively converged value obtained at much higher $N_{\text{max}}$, as in Fig. 2(c), we no longer have this luxury for $^6$He, where the growth in dimension with $N_{\text{max}}$ is much more rapid (Fig. 1). We must simply compare the calculations obtained with oscillator and natural-orbital bases, and for different $N_{\text{max}}$, against each other.

The overall scale of the change in calculated energy with $N_{\text{max}}$ for $^6$He is much larger than for $^3$He. In the vicinity of the variational energy minimum, the change in calculated energy with each step in $N_{\text{max}}$ is $\lesssim 0.1$ MeV (Fig. 10), compared to steps of $\approx 0.001$ MeV for comparable $N_{\text{max}}$ in $^3$He (Fig. 2). This difference might be...
taken to reflect the greater complication in reproducing a higher-A system in general, as well as the challenging halo structure of $^6$He in particular.

However, in judging convergence, what is important is not only the size of the change between values calculated with successive $N_{\text{max}}$, but how this change decreases with $N_{\text{max}}$. A convenient baseline against which to compare the convergence of the ground state energy is the hypothesis of exponential convergence with respect to $N_{\text{max}}$,

$$E(N_{\text{max}}) = E_\infty + a \exp(-cN_{\text{max}}),$$

where $E_\infty$ is then the full-space value \cite{57}. The calculated values approach the full-space value in a geometric progression with successive steps in $N_{\text{max}}$. For exponential convergence, the residual $\delta E(N_{\text{max}}) \equiv E(N_{\text{max}}) - E_\infty$ of the calculated energy relative to the full-space value, considered above for $^3$He (Sec. III A), is given by $\delta E(N_{\text{max}}) = a \exp(-cN_{\text{max}})$, and thus decreases by a constant factor $e^{-2c}$ with each (even) step in $N_{\text{max}}$. On a logarithmic plot of the residual, as we considered for $^3$He in Fig. 2(c), this appears as equally spaced steps with respect to $N_{\text{max}}$, as was indeed approximately noted for $^3$He (Sec. III A).

For $^6$He, we have no converged value with respect to which to take residuals, and thus cannot generate a logarithmic plot of residuals as in Fig. 2(c). Nonetheless, we can still compare successive calculated values of the energy, for successive truncations $N_{\text{max}}$, and consider their difference $\Delta E(N_{\text{max}}) = E(N_{\text{max}}) - E(N_{\text{max}} - 2)$. For exponential convergence, the ratio of successive steps

$$\eta(N_{\text{max}}) \equiv \frac{E(N_{\text{max}}) - E(N_{\text{max}} - 2)}{E(N_{\text{max}} - 2) - E(N_{\text{max}} - 4)},$$

is simply a constant $\eta = e^{-2c}$, independent of $N_{\text{max}}$. That is, $\eta = 0.5$ corresponds to a step size in $E$ which is halved with each successive step in $N_{\text{max}}$, and a smaller value of $\eta$ corresponds to a more rapid exponential decay towards the full-space value. Such differences which decrease by a constant ratio again appear, on a logarithmic plot, to move downward by equal increments with each step in $N_{\text{max}}$.

We thus consider a plot of $\log(\Delta E)$ for the $^6$He ground state energy, in Fig. 11. The overall convergence behavior is qualitatively similar for calculations in oscillator (solid circles) and natural orbital (open squares) bases. The spacing between curves for successive $N_{\text{max}}$ is roughly uniform with $N_{\text{max}}$, but decreases gradually for higher $N_{\text{max}}$, i.e., the convergence “slows down” relative to exponential convergence. For the energies calculated in either basis, the step size $\Delta E$ decreases by a factor of $\sim 0.4$–0.6 with each step in $N_{\text{max}}$. At low $N_{\text{max}}$, $\Delta E$ in either basis is roughly comparable. However, for high $N_{\text{max}}$, the curves representing $\Delta E$ for the calculations in the natural-orbital basis lie approximately one step in $N_{\text{max}}$ ahead of those for the oscillator basis. Near the variational minimum in energy ($\hbar \omega \approx 15$ MeV), this is consistent with the observation from above, that the natural-orbital basis improves on the best oscillator-basis energy by about one step in $N_{\text{max}}$. But this observation holds uniformly over a wide range extending to higher $\hbar \omega$, as well (at lower $\hbar \omega$, the $\Delta E$ obtained with the natural orbital basis falls off much more sharply with $N_{\text{max}}$).

We now consider the r.m.s. radii, which provide measures of the halo structure. The calculated values of both $r_p$ and $r_n$, for the $^6$He ground state, are shown in Fig. 12. Note that the point-proton r.m.s. radius $r_p$, the point-neutron r.m.s. radius $r_n$, and the matter (or total point-nucleon) radius $r_m$ form a redundant set of observables, related by $Ar_m^2 = Zr_p^2 + Nr_n^2$. It is thus worth briefly reviewing the physical significance of these observables, in the context of $^6$He \cite{21,83}.

Although $r_p$ does not directly measure neutron halo structure, it is nonetheless indirectly sensitive to this structure, and it is accessible to electromagnetic measurement, through its simple relation to the charge radius. It is important to keep in mind that $r_p$, as calculated here and as accessed in experiment, is defined relative to the common center of mass of the protons and neutrons (see, e.g., Refs. 6, 18, 21, 83). In the cluster halo description of $^4$He, the $\alpha$ recoils against the halo neutrons, which consequently displaces the center of mass of the $\alpha$ (and thus of the protons) relative to this common center of mass. This induces an increase in $r_p$ going from $^4$He to $^6$He. (There may also be contributions from modifications to the structure of the $\alpha$ particle itself, or “core polarization” \cite{83}.) Experimentally, the increase in $r_p$ from 1.462(6) fm for $^4$He to 1.934(9) fm for $^6$He \cite{83,93,94} is taken as a principal indicator of halo structure in $^6$He.

Then, both $r_n$ and $r_m$ include direct contributions from the halo neutrons. While $r_n$ is more selectively a measure of the neutron distribution, it is $r_m$ which is extracted from nuclear reaction cross section or proton-nucleus elastic scattering measurements. The results thereby obtained for the $\alpha$ isotopes are model-dependent and contradictory (see Sec. III A of Ref. 21 for an overview). They variously suggest $r_n \approx 2.3$ fm–2.7 fm in $^4$He, corresponding to an increase relative to $^4$He of $\approx 50$–90%. Subject to these uncertainties, the increased matter radius in $^6$He is again taken as an indicator of halo structure.

For the calculated $r_p$ (lower curves in Fig. 12), the results obtained in the natural-orbital basis yield reduced $\hbar \omega$ dependence relative to those obtained in the oscillator basis, much as already seen for $^3$He (Fig. 4). At the extremes in $\hbar \omega$ shown in Fig. 12, the calculations in the natural-orbital basis thus lie several steps in $N_{\text{max}}$ “ahead” of the oscillator-basis calculations. Again, we show the value of $r_p$ deduced from the experimental $r_n$, \cite{83} for comparison (filled diamond).

For $^6$He, the curves of radius vs. $\hbar \omega$ exhibit recognizable crossing points regardless of which basis is used. Recall (Sec. III A) that these crossing points have been suggested as a heuristic estimator of the full-space value. These crossing points are displaced in $\hbar \omega$ relative to each other — from $\hbar \omega \approx 10$ MeV for the oscillator basis down...
to $\hbar \omega \approx 7$ MeV for the natural-orbital basis — but occur at comparable values for the observable ($r_p \approx 1.9$ fm), consistent with the experimental value noted above.

Naturally, given the halo structure, the calculated values for $r_n$ (upper curves in Fig. 12) are larger than for $r_p$, the $\hbar \omega$ dependence is stronger, and the changes in calculated value with each step in $N_{\text{max}}$ is larger. Again, crossing points are obtained for calculations in both the oscillator and natural-orbital bases, shifted towards lower $\hbar \omega$ (longer oscillator length) than for $r_p$, namely $\hbar \omega \approx 9$ MeV for the oscillator basis, and approaching $\hbar \omega \approx 6$ MeV for the natural-orbital basis. This shift is perhaps not surprising given the larger radial extent of the structure being described. These two crossing points again occur at comparable values for $r_n$, in the range $r_n \approx 2.6$ fm–2.7 fm. (In conjunction with the above value for $r_p$, this suggests $r_m \approx 2.4$ fm–2.5 fm.) The highest $N_{\text{max}}$ curves for the natural orbital calculations develop a flat “shoulder”, varying by $\lesssim 0.05$ fm over several steps in $\hbar \omega$. This range of calculated $r_n$ values is compatible with the range $r_n \approx 2.5$ fm–3.0 fm suggested by the range of experimental matter radii (discussed above) in conjunction with $r_p$.

The transformation to a natural-orbital basis clearly does not definitively solve the problem of convergence for the r.m.s. radius observables. Nonetheless, it does contribute to taming the convergence behavior for these observables.

### B. Natural orbitals

Turning to the natural orbitals themselves, the radial wave functions for the lowest natural orbitals are shown in Fig. 13 for a high-$N_{\text{max}}$ calculation ($N_{\text{max}} = 14$) with $\hbar \omega$ near the variational energy minimum ($\hbar \omega = 15$ MeV). The orbitals shown again correspond, by their $nlj$ labels, to the oscillator orbitals through the $sd$ shell, as in the analogous figure above for $^3$He (Fig. 7). Radial functions are shown for both protons (short dashed lines) and neutrons (long dashed lines), and the oscillator radial functions are again shown for comparison (thick gray lines).

The mean occupations of these natural orbitals, indicated by the bars at top right in each panel of Fig. 13, are not far from what would be expected in a traditional shell model picture. The $s$ shell is almost filled, with an occupation of 1.81 for protons and 1.86 for neutrons. Most of the remaining occupation, out of a total occupation of 2 for the protons and 4 for the neutrons, lies in the $p$ shell. For the neutrons, in particular, the $0p_{3/2}$ orbital, which would be the expected orbital for the two valence neutrons in an extreme noninteracting shell model picture, naturally enough has a mean occupation of 1.81, while the $0p_{1/2}$ orbital accounts for a mean occupation of 0.21.

By contrast, the $1s_{1/2}$ orbital has a mean occupation of $< 0.05$. Thus, the halo neutrons are decisively $p$-shell nucleons. The occupations for the low-lying natural orbitals are higher than for the corresponding oscillator orbitals in the underlying calculation, but only marginally so: the increase in occupation is by $\approx 0.16$ for the neutron $p_{3/2}$ orbital, but only at the level of $\approx 0.01$ for the remaining $s$-shell and $p$-shell orbitals, for both protons and neutrons.

The $0s_{1/2}$ natural orbitals appear virtually unchanged, in Fig. 13 relative to the underlying oscillator orbital, for both protons and neutrons. This is consistent with an unmodified $\alpha$ “core”. However, to examine the large-distance behavior, we turn to logarithmic plots, shown in Fig. 14. Intriguingly, while both the proton and neutron natural orbitals have linear tails on the logarithmic plot, indicating exponential fall-off, the decay constants differ, with a slower fall-off (longer tail) for the neutron orbital.

The $0p_{3/2}$ orbital is of course of special interest, as the orbital “occupied” by the halo neutrons. The peak of the probability distribution, in the central region (Fig. 13), shifts only marginally outward in the radial coordinate, on the scale of $\approx 0.1$ fm. But the tail is noticeably extended even viewed on a linear scale. This is confirmed as a shallow exponential fall-off when viewed on a logarithmic scale (Fig. 14). In contrast, the peak for the “unoccupied” proton $0p_{3/2}$ orbital moves to smaller radius, by a comparable amount, and the tail similarly is retracted (Fig. 13), with a much steeper exponential fall-off (Fig. 14). Similar observations hold for the $0p_{1/2}$ orbital, which, as noted above, is partially occupied by the valence neutrons.

The sd-shell orbitals are notionally “unoccupied” orbitals for both the protons and neutrons. The mean occupations of these orbitals are each $\lesssim 0.05$. The proton orbitals move radially inward, relative to the oscillator orbital, both in terms of peak location and tail extent (Fig. 13). For the neutrons, the behavior is less consistent. The second peak of the $1s_{1/2}$ orbital, as well as the peak of the $0d_{5/2}$ orbital, both move markedly outwards, by $\approx 1$ fm, and the tails of these orbitals are even more exaggeratedly extended than for the neutron $p$-shell orbitals. Yet the neutron $0d_{3/2}$ orbital has a behavior which closely resembles that of the corresponding proton orbital, in the central region at least. Asymptotically, the proton orbitals have similar exponential tails, with faster decay than the neutron orbitals (Fig. 14).

For a “core” orbital, the proton $0s_{1/2}$ orbital, we explore the dependence on the $\hbar \omega$ and $N_{\text{max}}$ of the reference calculation in Fig. 15. The sensitivity of the natural orbital to the spectator $0s$ motion of the center of mass degree of freedom is similar to that already seen for this same orbital in $^3$He, discussed in Sec. III B. Once again, convergence is rapidly reached with increasing $N_{\text{max}}$ for the reference oscillator-basis calculation, while the shape of this converged natural orbital is dependent upon the $\hbar \omega$ of the underlying oscillator-basis calculation, which determines the $\hbar \omega_{\text{cm}}$ of the center-of-mass zero-point motion. In $^6$He (Fig. 15), the peak location for the natural orbital depends on the $\hbar \omega$ of the reference calculation less strongly than for $^3$He (Fig. 5), especially at low $\hbar \omega$.

Then, for the principal “halo” orbital, the neutron $0p_{3/2}$ orbital, the $\hbar \omega$ and $N_{\text{max}}$ dependence is similarly
explored in Fig. 16] now on a logarithmic scale. (The peak location has a similar dependence to that noted above for the proton $0s_{1/2}$ orbital.) The $\hbar \omega = 9$ MeV oscillator basis [Fig. 16(a)], with its comparatively long oscillator length, provides the best support in the tail region, and thus the fastest realization of a region of exponential decay (again, indicated by a straight line on the logarithmic plot). In contrast, the $\hbar \omega = 25$ MeV basis [Fig. 16(c)] yields the slowest grow-in of the exponential tail.

Finally, there is the question of the center-of-mass motion which emerges in these calculations for $^6$He in a natural-orbital basis. We apply the same diagnostics for $^6$He, shown in Fig. 17, as considered earlier for $^3$He in Sec. III C. That is, starting from the natural orbitals obtained from a reference oscillator basis calculation for $^6$He, we carry out the many-body calculation for $^6$He, then evaluate the center-of-mass ($R^2$) and ($K^2$) observables. From these, we deduce the “optimal” value of the $\hbar \omega_{\text{c.m.}}$ parameter for center-of-mass motion, $\hbar \omega_{\text{c.m.}}$, such that the expectation value $\langle N_{\text{c.m.}} \rangle$ of the center-of-mass number operator assumes its minimum value $\tilde{N}_{\text{c.m.}}$.

FIG. 13. Radial wave functions for the $^6$He $s$-, $p$-, and $sd$-shell natural orbitals, for both protons (short dashed lines) and neutrons (long dashed lines), plotted as the radial probability density $P(r)$. These are obtained from the underlying oscillator-basis calculation near the variational minimum ($\hbar \omega = 15$ MeV) and at high $N_{\text{max}}$ ($N_{\text{max}} = 14$). The corresponding oscillator radial functions for $\hbar \omega = 15$ MeV (thick gray lines) are shown for comparison. The mean occupation $n_a$ for each natural orbital, from the corresponding eigenvalue of the scalar density matrix, is indicated by the filling of the bar at top right (upper bar for protons, lower bar for neutrons).

Comparing the $^6$He results for the center-of-mass diagnostics (Fig. 17) to the $^3$He results (Fig. 8), a few features stand out. The dependence of both $\hbar \omega_{\text{c.m.}}$ [Fig. 17(a)] and $\tilde{N}_{\text{c.m.}}$ [Fig. 17(b)] on the reference basis parameters $\hbar \omega$ and $N_{\text{max}}$ is generally smoother for $^6$He than for $^3$He. The zig-zagging irregularities of Fig. 8 are no longer in evidence.

The oscillator parameter $\hbar \omega_{\text{c.m.}}$ for the center-of-mass motion [Fig. 17(a)] again matches that of the underlying oscillator basis for the natural orbitals in the vicinity of $\hbar \omega = 10$ MeV to 12.5 MeV. However, the $\hbar \omega$ dependence of $\hbar \omega_{\text{c.m.}}$ is less steep, above this point, for $^6$He than for $^3$He and, indeed, is continuing to become shallower with increasing $N_{\text{max}}$.

Furthermore, $\tilde{N}_{\text{c.m.}}$ [Fig. 17(b)] is comparatively independent of $\hbar \omega$. By $N_{\text{max}} = 14$, $\tilde{N}_{\text{c.m.}}$ has decreased to $\approx 10^{-3}$ over most of the $\hbar \omega$ range shown, and it continues to decrease with increasing $N_{\text{max}}$. Thus, in short, for practical purposes, a near-pure harmonic-oscillator $0s$ center-of-mass motion is uniformly obtained in the many-body calculations for $^6$He in the natural-orbital basis.
V. CONCLUSION

The nuclear many-body system is highly correlated, and thus inherently requires many antisymmetrized product states (Slater determinants) for its accurate description. No choice of single-particle states can completely obviate the need for superposing antisymmetrized product states in representing a correlated system. Nonetheless, a judicious choice of single-particle basis can accelerate the convergence of the description of the many-body wave function in a configuration-interaction basis.

Natural orbitals, obtained by diagonalizing the (scalar) one-body density matrix, address this aim in a well-defined sense, by maximizing the occupation of low-lying orbitals, or minimizing the depletion of the Fermi sea, in the expansion of a specific many-body reference state. After outlining the procedure for constructing and using natural orbitals within the NCCI framework (Sec. II), we have examined in detail the properties both of the orbitals themselves and the consequent many-body calculations in a natural orbital basis, first for the simple testbed case of $^3$He (Sec. III), then for the halo nucleus $^6$He (Sec. IV).

There are several noteworthy factors (Sec. II) limiting what we might expect to accomplish, in practice, with the natural orbital basis in NCCI calculations. The natural orbitals are only known to limited accuracy, as they are obtained from a reference many-body state which is only an approximation to the true solution of the many-body problem (as it would be obtained in an untruncated space). This reference state is represented in terms of orbitals from a truncated single-particle space, which limits the portion of the single-particle space which the natural orbitals can sample. Moreover, the many-body space in which the reference state is calculated is then subjected to a nontrivial truncation (e.g., by $N_{\text{max}}$), which may be expected to further restrict the fidelity of the reference state found therein and, specifically, the representation of high-lying orbitals in the scalar density obtained from this reference state.

Furthermore, even if the reference state could be found exactly, and its natural orbitals deduced exactly, one-body densities obtained from a laboratory-frame reference state are not uniquely defined by the intrinsic structure. Rather, they reflect some spectator center-of-mass motion arbitrarily superposed on this intrinsic structure. The natural orbitals obtained from these densities are then used in a many-body calculation which, although intended simply to reproduce the intrinsic structure of interest, in practice must yield some possibly complicated combination of intrinsic and center-of-mass motion.
Nonetheless, changing to a many-body basis constructed from natural orbitals does permit an NCCI calculation to probe portions of the many-body space which were not accessible in the original reference calculation. If the initial calculation is in an $N_{\text{max}}$-truncated harmonic oscillator basis, as here, then the calculation in a natural orbital basis brings in highly-excited oscillator configurations which were beyond the limit of the initial calculation.

We find that the transformation from harmonic oscillator orbitals to natural orbitals serves in part to simply accomplish a dilation of the harmonic oscillator basis, from the length scale (or $\hbar\omega$) of the underlying basis, to a more optimal length scale (or $\hbar\omega$). This observation (e.g., Fig. 15) already explains the relative insensitivity of calculated energies and other observables in a natural orbital basis to the $\hbar\omega$ of the underlying oscillator basis: as one varies $\hbar\omega$, the transformation to natural orbitals simply undoes this variation. Such dilation, in itself, merely recovers the results of a harmonic oscillator basis chosen with optimal length scale, rather than improving on it.

More substantially, though, the transformation to natural orbitals provides genuine modifications to the shape of the radial wave functions. Notably, the artificial Gaussian fall-off of the oscillator functions is modified to more closely resemble the exponential fall-off physically expected from the finite range of the nuclear interaction (e.g., Fig. 14). These differences can account for the im-

**FIG. 15.** Radial wave functions obtained for the $^6\text{He}$ proton $0s_{1/2}$ natural orbital, from different underlying oscillator-basis calculations, plotted as the radial probability density $P(r)$. Results are shown as obtained from underlying oscillator-basis calculations with (a) $\hbar\omega = 9\text{ MeV}$, (b) $\hbar\omega = 15\text{ MeV}$, and (c) $\hbar\omega = 25\text{ MeV}$. Radial wave functions are shown for $N_{\text{max}} = 2$ (dotted lines) through $N_{\text{max}} = 14$ (solid lines, highlighted), with the oscillator $0s$ function for the given $\hbar\omega$ (thick gray lines) shown for comparison. The locations of the peaks of the underlying harmonic-oscillator orbital and $N_{\text{max}} = 14$ natural orbital are marked with dashed vertical lines.

**FIG. 16.** Radial wave functions obtained for the $^6\text{He}$ neutron $0p_{3/2}$ natural orbital, from different underlying oscillator-basis calculations, plotted as the radial probability density $P(r)$, on a logarithmic scale. Results are shown as obtained from underlying oscillator-basis calculations with (a) $\hbar\omega = 9\text{ MeV}$, (b) $\hbar\omega = 15\text{ MeV}$, and (c) $\hbar\omega = 25\text{ MeV}$. Radial wave functions are shown for $N_{\text{max}} = 2$ (dotted lines) through $N_{\text{max}} = 14$ (solid lines, highlighted), with the oscillator $0s$ function for the given $\hbar\omega$ (thick gray lines) shown for comparison.
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