Systematics of binding energies and radii based on realistic two-nucleon plus phenomenological three-nucleon interactions

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We investigate the influence of phenomenological three-nucleon interactions on the systematics of ground-state energies and charge radii throughout the whole nuclear mass range from 4 He to 208 Pb. The three-nucleon interactions supplement unitarily transformed two-body interactions constructed within the unitary correlation operator method or the similarity renormalization group approach. To be able to address heavy nuclei as well, we treat the many-body problem in Hartree-Fock plus many-body perturbation theory, which is sufficient to assess the systematics of energies and radii, and limit ourselves to regularized three-body contact interactions. We show that even with such a simplistic three-nucleon interaction a simultaneous reproduction of the experimental ground-state energies and charge radii can be achieved, which is not possible with unitarily transformed two-body interactions alone.

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I. INTRODUCTION

Nuclear structure theory is approaching an era of systematic many-body calculations using nuclear Hamiltonians based on quantum chromodynamics (QCD). An important step along these lines is the formulation of nuclear interactions within chiral effective field theory $[1-3]$, leading to a consistent hierarchy of two-, three-, and many-nucleon interactions starting from the relevant degrees of freedom and symmetries for the low-energy nuclear structure regime. The use of these two-, three-, and many-nucleon interactions in nuclear structure calculations is a formidable task.

In addition to few-body calculations the most promising nuclear structure calculations using the chiral two- plus three-nucleon interaction consistently have been performed in the no-core shell model (NCSM) for mid *p*-shell nuclei [\[4\]](#page-8-0). An immense numerical effort is needed to compute and manage the three-body matrix elements in these calculations, which limits the range of applicability of these calculations at present. Recently, the use of consistent twoplus three-nucleon interactions resulting from a similarity renormalization group (SRG) evolution of the chiral two- plus three-nucleon interaction was demonstrated also in the context of the NCSM [\[5\]](#page-8-0). This approach, a unitary transformation of the chiral Hamiltonian aiming at a prediagonalization that improves the convergence properties of NCSM substantially, holds great potential also for the use in other many-body schemes and will play a significant role in the future. However, the computational effort for including those twoplus three-nucleon interactions into many-body calculations, be it exact or approximate, is still the limiting factor for many applications.

In this article we follow a more pragmatic route to explore the impact of three-body forces in connection with unitarily transformed two-nucleon interactions. We start from the Argonne V18 high-precision two-nucleon potential [\[6\]](#page-8-0), which is still widely used although it does not have the same systematic link to QCD like the chiral effective field theory interactions and is considered phenomenological in this respect. We then use the similarity renormalization group [\[7–11\]](#page-8-0) as well as the unitary correlation operator method (UCOM) [\[10,12](#page-8-0)[–14\]](#page-9-0) to construct a transformed two-nucleon interaction, which has a much better convergence behavior and allows us to use simplified many-body schemes. At this level neither genuine nor induced three-nucleon interactions are included. From various applications of these unitarily transformed two-nucleon interactions we know that there are characteristic deviations of basic nuclear observables from the experimental systematics that might be connected to three-body interactions. For example, unitarily transformed two-body interactions which yield realistic systematics for binding energies tend to underestimate the charge radii [\[10,](#page-8-0)[15\]](#page-9-0). Here we study to what extent these systematic deviations can be cured by including a three-body interaction. Note that we are not aiming at a precision description of individual nuclei but rather the complete systematics from light nuclei, ⁴He, to heavy nuclei, 208Pb.

To facilitate calculations for the full mass range from 4He to ²⁰⁸Pb we have to simplify the approach compared to a fully consistent treatment. The first simplification consists in the use of a phenomenological three-body interaction, which allows for an efficient computation of matrix elements but violates the consistency discussed above. The second simplification consists in the use of Hartree-Fock plus many-body perturbation theory for the approximate solution of the many-body problem. Despite of these simplifications, we will obtain valuable information on the interplay between realistic two-body and phenomenological three-body interactions and on how well the experimental systematics of ground-state energies and charge radii can be reproduced. Furthermore, these studies prepare the ground for calculations with consistently transformed twoplus three-nucleon interactions.

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After a brief reminder of the basic concepts of the unitary correlation operator method and the similarity renormalization group we introduce the phenomenological three-body interaction and calculate the matrix elements in the harmonic oscillator basis in the second section. In the third section, we discuss the inclusion of the three-body interaction in Hartree-Fock and many-body perturbation theory and discuss the systematics of ground-state energies and charge rms radii across the whole nuclear mass range and its dependence on the two- and three-nucleon interaction.

II. FORMALISM

A. Unitary correlation operator method and similarity renormalization group

The unitary correlation operator method and the similarity renormalization group provide two conceptually different but physically related approaches for the construction of soft phase-shift equivalent interactions.

The similarity renormalization group $[7-11]$ aims at the prediagonalization of the Hamiltonian for a given basis by means of a unitary transformation implemented through the renormalization-group flow equation:

$$
\frac{dH_{\alpha}}{d\alpha} = [\eta_{\alpha}, H_{\alpha}], \qquad (1)
$$

where α is the flow parameter and H_{α} the evolved Hamiltonian, with $H_0 = H$ being the initial or "bare" Hamiltonian. The anti-Hermitian generator η_{α} defines the specifics of the flow evolution, e.g., the representation with respect to which the Hamiltonian should become diagonal or block-diagonal. Various choices for this generator have been investigated [\[11\]](#page-8-0), and we restrict ourselves to the simple generator [\[7,9\]](#page-8-0)

$$
\eta_{\alpha} = [T_{\text{int}}, H_{\alpha}] \tag{2}
$$

with $T_{\text{int}} = T - T_{\text{cm}}$ being the intrinsic kinetic energy, which leads to a prediagonalization of the Hamiltonian with respect to the eigenbasis of the kinetic energy or momentum operator. Once the generator is fixed, the Hamiltonian and all operators of interest can be evolved easily using a matrix representation of the flow equation (1) .

In *A*-body space the evolution generates up to *A*-body operators even if the initial Hamiltonian contains only up to two- or three-body operators. For reasons of practicability one has to truncate the evolution at some low particle number typically this is done by solving the evolution equation in a matrix representation in two- or three-body space. For the moment we restrict ourselves to transformations in two-body space, i.e., we will discard any induced three-body interactions.

The aim of the unitary correlation operator method [\[10,12](#page-8-0)[–14,16\]](#page-9-0) is to explicitly treat short-range correlations induced by the nuclear interaction via a static unitary transformation. This transformation can either be used to correlate the many-body states or to similarity transform operators of interest, e.g., the Hamiltonian

$$
\tilde{H} = C^{\dagger} H C,\tag{3}
$$

using the correlation operator *C*. The dominant short-range correlations are induced by the strong short-range repulsion

and the tensor part of the nuclear interaction. Therefore, the correlation operator is written as a product of two unitary operators, C_r for the central correlations and C_Ω for the tensor correlations. We choose an explicit form of the correlation operators:

$$
C = C_{\Omega}C_r = \exp\left(-i\sum_{j < k} g_{\Omega,jk}\right) \exp\left(-i\sum_{j < k} g_{r,jk}\right) \tag{4}
$$

with the following ansatz for the Hermitian generators *gr* and *g*:

$$
g_r = \frac{1}{2} [q_r s(r) + s(r) q_r],
$$

\n
$$
g_{\Omega} = \frac{3}{2} [(\sigma_1 \cdot r) (\sigma_2 \cdot q_{\Omega}) + (\sigma_1 \cdot q_{\Omega}) (\sigma_2 \cdot r)],
$$
\n(5)

where $q_r = \frac{1}{2}(\frac{r}{r} \cdot q + q \cdot \frac{r}{r}), q_{\Omega} = q - \frac{r}{r} \cdot q_r,$ and $q =$
¹[n = n] The strengths and radial dependencies of the two $\frac{1}{2}$ [$\mathbf{p}_1 - \mathbf{p}_2$]. The strengths and radial dependencies of the two transformations are governed by the correlation functions $s(r)$ and $\vartheta(r)$ for the central and tensor correlations, respectively. One can obtain these functions via an energy minimization in the two-body system [\[16\]](#page-9-0). Recently, we have also employed the SRG as a tool for the determination of the UCOM correlation functions $s(r)$ and $\vartheta(r)$ as discussed in Refs. [\[9,10\]](#page-8-0). Here, we will use these SRG-optimized UCOM correlation functions only.

Though the SRG and UCOM transformations have a different formal background, they address the same physics of short-range correlations. A first connection becomes clear at the level of the generators [\[8\]](#page-8-0)—the SRG generator (2) in two-body space at $\alpha = 0$ reveals the same operator structures that appear in the UCOM generators (5). At the level of matrix elements, both the SRG and UCOM transformations lead to a suppression of the off-diagonal momentum-space matrix elements and an enhancement of the low-momentum matrix elements as discussed in detail in Ref. [\[10\]](#page-8-0).

In the following, we employ both transformations to generate one-parameter families of phase-shift equivalent two-body interactions starting from a specific initial *NN* interaction, the Argonne V18 (AV18) in our case. For the SRG transformation the flow parameter α directly spans this family of two-body interactions. We will study two versions of the SRG transformation, one where the flow equations are solved for all partial waves and one where only the partial waves containing relative *S* waves, i.e., the ${}^{1}S_{0}$ and the coupled ${}^{3}S_{1}$ - ${}^{3}D_{1}$ partial waves, are transformed. The latter is motivated by the fact that short-range correlations affect the *S*-wave channels most, because for all higher orbital angular momenta the relative wave functions are suppressed by the centrifugal barrier at short distances. We use the label "SRG" for the fully transformed interactions and "S-SRG" for the *S*-wave-only transformations. For the UCOM transformation we use correlation functions determined from SRG-evolved two-body wave functions as discussed in Refs. [\[9,10\]](#page-8-0), thus the flow parameter α also spans a family of different UCOMtransformed interactions. Note that the standard formulation of UCOM only uses different transformations for the different (*S,T*) channels. We thus use the SRG-evolved wave functions for the lowest partial waves for each (S, T) channel to define the correlation functions, leading to a transformed

interaction labeled "UCOM(SRG)." Analogously to the S-SRG transformation, we can also use an *S*-wave-only UCOM transformation, denoted "S-UCOM(SRG)," which acts only in the ¹S₀ and the coupled ${}^{3}S_{1}$ - ${}^{3}D_{1}$ partial waves.

So far, we have assumed that both transformations are evaluated in two-body space, leading to a transformed interaction containing two-body terms only. A consistent first-principles treatment requires the transformation to be performed in *A*-body space, leading to a hierarchy of induced interactions up to the *A*-body level, as mentioned earlier. The most advanced attempts along these lines use the full SRG evolution at the three-body level to construct a consistently transformed twoplus three-nucleon interaction [\[5\]](#page-8-0). The use of those twoplus three-body interactions in many-body calculations is very demanding and presently limited to rather small model spaces.

Therefore, we follow a more pragmatic path in this work. We evaluate the unitary transformations at the two-body level and mimic the three-body contributions (genuine plus induced) through a simple phenomenological three-body interaction. By using a simplified three-nucleon (3*N*) interaction, e.g., a regularized contact or a Gaussian interaction, the calculation of the three-body matrix elements becomes formally and computationally much less demanding. This allows us to study the impact of 3*N* interactions on various nuclear structure observables for nuclei and model spaces beyond the domain accessible with realistic 3*N* interactions. Furthermore, we can develop and benchmark approximate treatments of the threebody contributions and establish the technical framework to include 3*N* interaction into different many-body methods.

The parameters of the phenomenological 3*N* interactions will be adjusted depending on the flow parameter *α* of the transformed two-nucleon (*NN*) interaction. For a wide range of α parameters the transformed two-body interaction alone produces an overbinding compared to the experimental ground-state energy. This is illustrated in Fig. 1 for the groundstate energy of ⁴He as function of α obtained in converged no-core shell-model calculations for the UCOM(SRG)-, the

FIG. 1. (Color online) Binding energy of ⁴He as function of the flow parameter α obtained from a converged no-core shellmodel calculation using the UCOM(SRG)-transformed $(•)$, the S-UCOM(SRG)-transformed (\blacklozenge) , the SRG-transformed (\blacktriangle) , or the S-SRG-transformed (\blacksquare) AV18 potential. The horizontal lines indicate the experimental binding energy (—) and the exact groundstate energy for the bare AV18 two-body interaction (---) [\[17\]](#page-9-0).

S-UCOM(SRG)-, the SRG-, and the S-SRG-transformed AV18 interaction. Thus the additional phenomenological interaction, which mimics the net effect of the genuine and the induced 3*N* interaction, has to be repulsive in order to lead to a 4He binding energy consistent with experiment. Note that the phenomenological three-body forces that are used in connection with the bare AV18 interaction are generally attractive. Thus the induced 3*N* interaction resulting from the unitary transformation of the *NN* interaction alone has to be repulsive and sufficiently strong to create an overall repulsive three-body contribution.

B. Three-body contact interaction

The simplest choice for a phenomenological 3*N* interaction is a spin-isopin-independent contact interaction

$$
V_{3N} = C_{3N} \, \delta^{(3)}(x_1 - x_2) \, \delta^{(3)}(x_1 - x_3) \tag{6}
$$

with variable strength C_{3N} . Despite its simplicity it allows us to study the impact of a 3*N* interaction on bulk observables like ground-state energies or charge radii. Obviously, this simplistic choice offers substantial computational advantages.

For evaluating the matrix elements of a realistic 3*N* interaction for the use in configuration-space Hartree-Fock or no-core shell-model type calculations one typically adopts a two-step procedure: First the matrix elements are evaluated in a Jacobi-coordinate basis for the relative motion in the three-nucleon system. Then, through a sequence of Talmi-Moshinski transformations and angular-momentum recouplings, the matrix elements are transformed into the *m* scheme to perform the many-body calculation. Both steps are nontrivial and computationally demanding, thus limiting the model-space sizes for which those matrix elements can be handled.

In contrast, the matrix elements of the contact interaction can be directly evaluated in the m-scheme in a straightforward manner. We first consider the matrix elements of the 3*N* contact interaction with respect to the spatial part of three-particle product states in the harmonic oscillator basis

$$
\langle n_1 l_1 m_{l_1}, n_2 l_2 m_{l_2}, n_3 l_3 m_{l_3} | V_{3N} | n_4 l_4 m_{l_4}, n_5 l_5 m_{l_5}, n_6 l_6 m_{l_6} \rangle. \tag{7}
$$

The spin and isospin quantum numbers and the antisymmetrization will be included subsequently. We can insert a unit operator in position representation using Cartesian coordinates and directly evaluate the Kronecker deltas. This leaves us with a single integration over a single-particle coordinate, which can be rewritten in spherical coordinates. Introducing the position representation of the harmonic oscillator single-particle states, $\phi_{nlm_l}(\mathbf{x}) = R_{nl}(x)Y_{lm_l}(\Omega)$, with radial wave functions $R_{nl}(x)$ and spherical harmonics $Y_{lm}(\Omega)$, we obtain:

$$
\langle n_1 l_1 m_{l_1}, n_2 l_2 m_{l_2}, n_3 l_3 m_{l_3} | V_{3N} | n_4 l_4 m_{l_4}, n_5 l_5 m_{l_5}, n_6 l_6 m_{l_6} \rangle
$$

= $C_{3N} \int dx x^2 R_{n_1 l_1}(x) R_{n_2 l_2}(x) R_{n_3 l_3}(x)$
 $\times R_{n_4 l_4}(x) R_{n_5 l_5}(x) R_{n_6 l_6}(x)$
 $\times \int d\Omega Y_{l_1 m_{l_1}}^*(\Omega) Y_{l_2 m_{l_2}}^*(\Omega) Y_{l_3 m_{l_3}}^*(\Omega)$
 $\times Y_{l_4 m_{l_4}}(\Omega) Y_{l_5 m_{l_5}}(\Omega) Y_{l_6 m_{l_6}}(\Omega).$ (8)

The integral over the six radial wave functions $R_{nl}(x)$ has to be calculated numerically while the integral over the six spherical harmonics $Y_{lm_l}(\Omega)$ can be evaluated analytically. The product of three spherical harmonics can be reduced to one spherical harmonic and the integral over the remaining two spherical harmonics can be solved analytically, leading to

$$
\int d\Omega Y_{l_1m_{l_1}}^*(\Omega) Y_{l_2m_{l_2}}^*(\Omega) Y_{l_3m_{l_3}}(\Omega) Y_{l_4m_{l_4}}(\Omega) Y_{l_5m_{l_5}}(\Omega) Y_{l_6m_{l_6}}(\Omega)
$$
\n
$$
= \frac{1}{16\pi^2} \hat{t_1} \hat{t_2} \hat{t_3} \hat{t_4} \hat{t_5} \hat{t_6} \sum_{\substack{L_1L_2L_3 \\ M_{L_1}M_{L_2}M_{L_3}}} \frac{1}{2L_2 + 1}
$$
\n
$$
\times c \begin{pmatrix} l_1 & l_2 & | & L_1 \\ 0 & 0 & | & 0 \end{pmatrix} c \begin{pmatrix} L_1 & l_3 & | & L_2 \\ 0 & 0 & | & 0 \end{pmatrix}
$$
\n
$$
\times c \begin{pmatrix} l_4 & l_5 & | & L_3 \\ 0 & 0 & | & 0 \end{pmatrix} c \begin{pmatrix} L_3 & l_6 & | & L_2 \\ 0 & 0 & | & 0 \end{pmatrix}
$$
\n
$$
\times c \begin{pmatrix} l_1 & l_2 & | & L_1 \\ m_{l_1} & m_{l_2} & | & M_{l_1} \end{pmatrix} c \begin{pmatrix} L_1 & l_3 & | & L_2 \\ M_{L_1} & m_{l_3} & | & M_{L_2} \end{pmatrix}
$$
\n
$$
\times c \begin{pmatrix} l_4 & l_5 & | & L_3 \\ m_{l_4} & m_{l_5} & | & M_{l_3} \end{pmatrix} c \begin{pmatrix} L_3 & l_6 & | & L_2 \\ M_{L_3} & m_{l_6} & | & M_{L_2} \end{pmatrix}
$$
\n
$$
(9)
$$

with $\hat{l} = \sqrt{2l+1}$ and $c(\frac{l_1}{m_{l_1}} \frac{l_2}{m_{l_2}} | \frac{l}{M_L})$ being Clebsch-Gordan coefficients.

We precompute and store those angular integrals as well as the radial integrals in [\(8\)](#page-2-0). The inclusion of the spin and isospin quantum numbers, the coupling of the single-particle orbital angular momenta and the spins, and the antisymmetrization are then done on the fly during the many-body calculation. This makes calculations in large model spaces feasible.

For applications beyond the mean-field level a regularization of the contact interaction is inevitable. However, the regularization should preserve the simplicity of the matrixelement calculation, which rules out momentum-space cutoffs and such. Hence, we introduce an energy cut-off parameter *e*3*^N* , which defines an upper bound for total oscillator energy of the three-particle state, $(2n_1 + l_1) + (2n_2 + l_2) + (2n_3 + l_1)$ l_3) \leq e_{3N} . The implementation of this cutoff is trivial and it preserves all computational advantages of the contact interaction.

III. MANY-BODY CALCULATIONS

We adopt the 3*N* contact interaction together with unitarily transformed *NN* interactions for the study of the systematics of nuclear ground-state energies and charge radii throughout the whole mass range from 4 He to 208 Pb using Hartree-Fock and many-body perturbation theory.

A. Hartree-Fock Approximation

We have employed the Hartree-Fock (HF) approximation as a first indicator for the gross systematics of binding energies and charge radii obtained with unitarily transformed two-body interactions in Refs. $[10,15]$ $[10,15]$ already. In order to assess the impact of 3*N* contact interactions we extend our HF framework in a first step.

All calculations are based on the translationally invariant Hamiltonian

$$
H_{\text{int}} = T_{\text{int}} + V_{NN} + V_{3N} = H_{\text{int}}^{(2)} + V_{3N} \tag{10}
$$

with V_{NN} being the UCOM- or SRG-transformed *NN* interaction and $T_{\text{int}} = T - T_{\text{cm}}$ the intrinsic kinetic energy. This Hamiltonian includes all charge-dependent and electromagnetic terms of the transformed AV18 potential as well as the phenomenological three-body force. For the intrinsic kinetic energy we use the pure two-body form of Ref. [\[15\]](#page-9-0).

The HF equations are formulated in a harmonic oscillator basis representation, i.e., the single-particle states are expanded in the harmonic oscillator states:

$$
|vljmm_t\rangle = \sum_n C_n^{(vljm_t)} |nljm m_t\rangle, \qquad (11)
$$

where $|nljmm_t\rangle$ denotes the harmonic oscillator eigenstates with radial quantum number *n*, orbital angular momentum *l*, total angular momentum *j* with projection *m*, and isospin projection quantum number m_t . Since we only consider closedshell nuclei in the following, the expansion coefficients are independent of *m*. The HF equations can now be written as

$$
\sum_{\bar{n}} h_{n\bar{n}}^{(ljm_t)} C_{\bar{n}}^{(vljm_t)} = \varepsilon^{(vljm_t)} C_n^{(vljm_t)}
$$
(12)

with the single-particle energies $\varepsilon^{(vljm_t)}$. The matrix elements of the single-particle HF Hamiltonian

$$
h_{n\bar{n}}^{(ljm_t)} = \sum_{l'j'm'_t} \sum_{n'\bar{n}'} \langle nljm_t, n'l'j'm'_t |
$$

\n
$$
\times H_{int}^{(2)} |\bar{n}ljm_t, \bar{n}'l'j'm'_t\rangle \varrho_{\bar{n}'n'}^{(l'j'm'_t)}
$$

\n
$$
+ \frac{1}{2} \sum_{l'j'm'_t \atop l''j'm''_t \bar{n}'n''} \sum_{n'\bar{n}''} \langle nljm_t, n'l'j'm'_t, n''l''j''m''_t |
$$

\n
$$
\times V_{3N} |\bar{n}ljm_t, \bar{n}'l'j'm'_t, \bar{n}''l''j''m''_t\rangle \varrho_{\bar{n}'n'}^{(l'j'm'_t)} \varrho_{\bar{n}''n''}^{(l''j''m'')} (13)
$$

are obtained by contractions of the antisymmetrized matrix elements of the two-body part of the Hamiltonian $H_{\text{int}}^{(2)}$ and the three-body interaction V_{3N} with the one-body density matrix given by

$$
\varrho_{\bar{n}n}^{(ljm_t)} = \sum_{v} O^{(vljm_t)} C_{\bar{n}}^{(vljm_t)^*} C_n^{(vljm_t)}
$$
(14)

with $O^{(\nu l j m_t)}$ being the number of occupied magnetic sublevels which is $2j + 1$ for closed-shell nuclei.

In the following the HF approach is applied to selected closed-shell nuclei from 4 He to 208 Pb. The HF equations are solved iteratively until full self-consistency is reached. The model space is truncated at a given major oscillator quantum number $e = 2n + l \le e_{\text{max}}$, where $e_{\text{max}} = 10$ is sufficient to obtain converged ground-state energies and radii at the HF level. The oscillator parameter is chosen for each nucleus separately such that the experimental charge radius is reproduced by a shell-model Slater determinant built from harmonic oscillator single-particle states.

FIG. 2. (Color online) Ground-state energies per nucleon and charge radii of selected closed-shell nuclei resulting from HF calculations based on pure two-body interactions for $e_{\text{max}} = 10$: UCOM(SRG) with $\alpha = 0.16$ fm⁴ (^{\bullet}), S-UCOM(SRG) with $\alpha =$ 0.16 fm⁴ (■), SRG with *α* = 0.10 fm⁴ (◆), S-SRG with *α* = 0.10 fm⁴ (\triangle) . The bars indicate the experimental values [\[18,19\]](#page-9-0).

As a first illustration of the behavior of unitarily transformed two-body interactions Fig. 2 summarizes the ground-state energies per nucleon and the charge radii obtained at the HF level for nuclei up to 208Pb. We adopt four different twobody interactions—UCOM(SRG), S-UCOM(SRG), SRG, and S-SRG—with flow parameters relevant for the later calculations including the 3*N* contact interaction. We observe that the general trend of the binding energies and charge radii is similar for the UCOM(SRG), the S-UCOM(SRG), and the S-SRG interactions. All three interactions produce binding energies that are within 2 MeV per nucleon of the experimental values for the whole mass range. By including correlations beyond HF, e.g., through many-body perturbation theory, all interactions would lead to an overbinding compared to experiment. At the same time the charge radii are underestimated for all but the lightest isotopes. Those systematic deviations can be remedied by a repulsive 3*N* interaction, as it will be included in the next step.

The SRG-transformed interaction exhibits a vastly different behavior. The binding energies per nucleon increase rapidly with mass number, leading to an completely unphysical overbinding already at the HF level for intermediate and heavy nuclei. At the same time the charge radii are even smaller than the ones obtained with the other transformed interactions. The origin of this difference in systematics as compared to the UCOM- and the S-SRG-transformed interactions seems to lie in the treatment of higher partial waves. Only the standard SRG transformations rigorously prediagonalize the higher partial waves, all other transformations do not affect those channels at all (S-SRG and S-UCOM) or transform them in a way not specifically optimized for them (UCOM). The overbinding produced by the SRG interaction as compared to the corresponding S-SRG interaction has to be compensated by stronger induced three- and many-body interactions, which

FIG. 3. (Color online) Ground-state energies per nucleon and charge radii of selected closed-shell nuclei resulting from HF calculations for the pure two-body interaction S-UCOM(SRG) for $e_{\text{max}} = 10$ and different flow parameters: $\alpha = 0.04 \text{ fm}^4$ (\bullet), $\alpha =$ 0.12 fm^4 (\blacksquare), $\alpha = 0.16 \text{ fm}^4$ (\blacklozenge). The bars indicate the experimental values [\[18,19\]](#page-9-0).

seem to be influenced sizably by contributions from higherpartial waves.

For the construction of phenomenological 3*N* interactions the UCOM, S-UCOM, and S-SRG interactions provide a better starting point, because the corrections to the binding energy and radius systematics are much smaller. Therefore, we will focus on those interactions in the following.

Before including the 3*N* contact interaction explicitly, we analyze the dependence of the HF results obtained with the transformed two-body interactions on the flow parameter *α*. In Fig. 3 the binding energies and charge radii for the S-UCOM(SRG) interactions with $\alpha = 0.04$ fm⁴, 0.12 fm⁴, and 0.16 fm⁴ are shown. For the smallest flow parameter $\alpha =$ 0*.*04 fm⁴ the ground-state energies reproduce the systematics of the experimental values up to a constant shift. The missing binding energy can be explained by beyond-HF correlations that can be recovered, e.g., by perturbation theory. This flow parameter would be used for calculations based on the pure *NN* interaction, as they are discussed in detail in Refs. [\[9,10](#page-8-0)[,15\]](#page-9-0).

When increasing the flow parameter entering into the construction of the S-UCOM(SRG) interaction to $\alpha = 0.12$ fm⁴ or 0*.*16 fm⁴ the ground-state energy at the HF level decreases substantially. For most nuclei the binding energy per nucleon more than doubles when going from $\alpha = 0.04$ fm⁴ to 0.16 fm⁴. For heavier nuclei the increase is larger, thus leading to a tilt of the ground-state energy systematics with respect to the experimental behavior. Unlike the energies, the charge radii exhibit a very weak α dependence as shown in the lower panel of Fig. 3. For all flow parameters considered here, the radii are somewhat underestimated. The situation is very similar for the UCOM(SRG) and the S-SRG interactions.

This general phenomenology of ground-state energies and charge radii obtained from unitarily transformed interactions at larger flow parameters illustrates that the purely repulsive

FIG. 4. (Color online) Binding energies per nucleon and charge radii of selected closed-shell nuclei resulting from HF calculations based on the S-UCOM(SRG) interaction for $\alpha = 0.16$ fm⁴, $e_{\text{max}} = 10$, $e_{3N} = 20$ and different strengths of the three-body interaction: $C_{3N} = 1600 \text{ MeV fm}^6$ (\bullet), $C_{3N} = 2200 \text{ MeV fm}^6$ (\bullet), $C_{3N} = 2800 \,\text{MeV fm}^6$ (\blacklozenge). The bars indicate the experimental values [\[18,19\]](#page-9-0).

phenomenological 3*N* interaction can be used to improve energies and radii. Since the radii are insensitive to the flow parameter in a certain regime, we can fix the strength of the 3*N* interaction such that the systematics of the charge radii is in good agreement with experiment. The flow parameter can then be chosen to provide an optimal description of the ground-state energies in a beyond-HF calculation.

The impact of 3*N* contact interactions with different strength parameters C_{3N} is illustrated in Fig. 4 using the S-UCOM(SRG) interaction for $\alpha = 0.16$ fm⁴. As compared to the HF calculation with the pure two-body interaction, the binding energies are reduced significantly and the charge radii are increased as a result of the purely repulsive 3*N* interaction. It is remarkable that the charge radii are in excellent agreement with experiment for the whole mass range from ⁴He to ²⁰⁸Pb when using a 3*N* interaction with strength parameters in the range $C_{3N} = 2200$ to 2800 MeV fm⁶. For the same values C_{3N} the ground-state energy systematics at the HF does again resemble the experimental systematics up to a constant shift, i.e., the tilt of the energy curve toward an overbinding for heavier nuclei is cured as well. The missing binding energy of 3 to 4 MeV per nucleon at the HF level can be recovered by including correlations beyond HF, as will be discussed in the next section.

In addition to ground-state energies and radii the HF approximation provides us with an estimate for the singleparticle energies that can be used to diagnose the various $NN + 3N$ interactions. Examples for the single particle spectra obtained with the various unitarily transformed interactions for 40 Ca and 90 Zr are shown in Figs. 5 and 6, respectively. We use the UCOM(SRG) and the S-UCOM(SRG) interactions for $\alpha = 0.16$ fm⁴ and the S-SRG interaction for $\alpha = 0.10$ fm⁴ each supplemented with a 3*N* contact interaction with strength parameter adjusted to provide a good overall description of

FIG. 5. (Color online) Single-particle spectra of 40 Ca for different interactions: (1) UCOM(SRG) with $\alpha = 0.16$ fm⁴, $C_{3N} = 1600 \text{ MeV fm}^6$, (2) S-UCOM(SRG) with $\alpha = 0.16 \text{ fm}^4$, $C_{3N} = 2200 \text{ MeV fm}^6$, (3) S-SRG with $\alpha = 0.10 \text{ fm}^4$, $C_{3N} =$ 2000 MeV fm⁶. Three-body cut-off set to $e_{3N} = 20$ for all calculations. Occupied states are indicated by solid lines, unoccupied states by dashed lines. Experimental data taken from Ref. [\[20\]](#page-9-0).

the charge radii at the HF level, i.e., $C_{3N} = 1600 \,\text{MeV fm}^6$ for UCOM(SRG), $C_{3N} = 2200 \,\text{MeV fm}^6$ for S-UCOM(SRG), and $C_{3N} = 2000 \,\text{MeV fm}^6$ for S-SRG. We note that the standard SRG interaction for $\alpha = 0.10$ fm⁴ would require $3N$ strengths C_{3N} in the range of 4000 to 5000 MeV fm⁶ to generate realistic radii.

The gross structure of the single-particle spectra obtained with the S-UCOM(SRG) + $3N$ and the S-SRG + $3N$ interactions agrees rather well with the single-particle energies extracted from experiment. The quality of the agreement is

FIG. 6. (Color online) Single-particle spectra of $90Zr$ for the same interactions used in Fig. 5. Experimental data taken from Refs. [\[21,22\]](#page-9-0).

comparable with other mean-field type calculations and some of the characteristic deviations, e.g., the overestimation of the gaps at the Fermi energy, are expected to be remedied by the inclusion of beyond HF corrections. Other important quantities, e.g., the splittings between spin-orbit partner states, are reproduced rather well at the HF level already.

The picture is different for the UCOM(SRG) + 3*N* interaction. In particular for the single-particle spectrum of ^{90}Zr shown in Fig. [6](#page-5-0) and for all heavier nuclei we observe a collapse of the spin-orbit splittings. Since this problem does not appear in the corresponding S-UCOM(SRG) calculation, it has to be caused by the UCOM transformation of the higher partial waves. The problem is also absent in UCOM(SRG) interactions for smaller flow parameters α , e.g., for the UCOM(SRG) interaction at $\alpha = 0.04$ fm⁴ that was used in Fig. [3.](#page-4-0) Thus the long-range character of the tensor correlation functions as they appear for larger α (cf. Refs. [\[9,10\]](#page-8-0)) acting on the higher partial wave leads to this unphysical behavior. We will, therefore, restrict ourselves for the following discussion to the S-UCOM(SRG) and S-SRG interactions.

B. Many-body perturbation theory

A simple means to estimate the impact of correlations beyond the HF approximation is many-body perturbation theory (MBPT). In particular low-order MBPT corrections to the energy [\[23–26\]](#page-9-0) can be computed quite efficiently for the whole mass range up to ²⁰⁸Pb. We have used second- and thirdorder MBPT to investigate various two-body Hamiltonians and the importance and systematics of correlations beyond HF in Refs. [\[10,](#page-8-0)[15\]](#page-9-0). One should be aware, however, that low-order MBPT can only provide an estimate for the exact ground-state energies and that the order-by-order convergence is not guaranteed, as we have shown in Ref. [\[27\]](#page-9-0) using a harmonic oscillator single-particle basis.

Because of its computational simplicity we adopt secondorder MBPT as a guideline for the effect of beyond-HF correlations on the energy in the presence of a phenomenological 3*N* interaction. The second-order energy correction to the HF ground-state energy for the intrinsic Hamiltonian [\(10\)](#page-3-0) including the 3*N* interaction reads

$$
E^{(2)} = \frac{1}{4} \sum_{hh'}^{<\varepsilon_F} \sum_{pp'}^{<\varepsilon_F} \frac{\left| \langle hh'| H_{int}^{(2)} | pp' \rangle + \sum_{\tilde{h}}^{<\varepsilon_F} \langle hh' \bar{h} | V_{3N} | pp' \bar{h} \rangle \right|^2}{\varepsilon_h + \varepsilon_{h'} - \varepsilon_p - \varepsilon_{p'}} + \frac{1}{36} \sum_{hh'h''}^{<\varepsilon_F} \sum_{pp'p''}^{>\varepsilon_F} \frac{\left| \langle hh'h'' | V_{3N} | pp'p'' \rangle \right|^2}{\varepsilon_h + \varepsilon_{h'} + \varepsilon_{h''} - \varepsilon_p - \varepsilon_{p'}} , \tag{15}
$$

where h, h', \ldots , denote the HF single-particle states (11) below the Fermi energy ϵ_F (hole states) and p, p', \ldots , the corresponding HF single-particle states above the Fermi energy (particle states). All two- and three-body matrix elements appearing here are understood to be antisymmetrized matrix elements.

Already the structure of the second-order energy correction (15) is interesting. Obviously, if we set all matrix elements of the 3*N* interaction to zero we recover the well-known

form of the second-order correction for a pure two-body Hamiltonian. The inclusion of the 3*N* interaction affects this expression in two ways: (i) The matrix elements of the two-body Hamiltonian are modified by an effective or in-medium two-body term that results from the three-body matrix elements by a contraction of the third single-particle index. (ii) An additional pure three-body term involving three particle and three hole indices appears.

To separate the effect of these two contributions we study three variants of the second-order energy correction: MBPT(2B) includes only the contribution of the two-body Hamiltonian, i.e., the first matrix element in Eq. (15). $MBPT(2B + 3Bpphh)$ includes the in-medium two-body contribution generated by the 3*N* interaction, i.e., the complete first term in Eq. (15) . Finally, MBPT $(2B + 3B)$ includes all terms of Eq. (15) .

The ground-state energies of 4 He, 16 O, and 40 Ca obtained with MBPT(2B), MBPT(2B + 3Bpphh), and MBPT(2B + 3B) on top of the HF result for the S-UCOM(SRG) interaction are shown in Fig. 7 as function of the cut-off parameter e_{3N} . Note that the calculations are done in a model space with $e_{\text{max}} = 10$ and, therefore, all energies will become independent of the three-body cutoff once $e_{3N} \ge 3e_{\text{max}} = 30$. For the HF calculations presented so far we had fixed this cutoff to $e_{3N} = 20$, which was sufficiently large to guarantee that the HF energies were practically independent of this cutoff for all nuclei. As soon as we include the second-order perturbative correction we cannot expect the results to be independent of *e*3*^N* , because the sums over particle states above the Fermi energy directly probe high-lying matrix elements. Eventually we will have to fix e_{3N} to a certain value as part of the definition of the phenomenological 3*N* interaction. For the study of the different contributions a cutoff variation nevertheless provides a useful diagnostic tool. As seen in Fig. 7, the HF energies are practically independent of *e*3*^N* as mentioned earlier. When including the second-order correction due to the two-body Hamiltonian, MBPT(2B), the ground-state energies are lowered by about 1 MeV per nucleon for ⁴He and by about 2.5 MeV per nucleon for ${}^{16}O$ and ${}^{40}Ca$. The MBPT(2B) energies are sensitive to e_{3N} only indirectly via high-lying HF singleparticle states; therefore, the dependence is marginal. For $MBPT(2B + 3Bpphh)$ and $MBPT(2B + 3B)$ the cutoff directly

FIG. 7. (Color online) Contributions to the ground-state energy resulting from MBPT based on the S-UCOM(SRG) interaction for $\alpha = 0.16 \text{ fm}^4$, $e_{\text{max}} = 10$, $C_{3N} = 2200 \text{ MeV fm}^6$ as function of the cut-off parameter e_{3N} for HF (\bullet), HF + MBPT(2B) (\blacksquare), $HF + MBPT(2B + 3Bpphh)$ (\blacklozenge), and $HF + MBPT(2B + 3B)$ (\blacktriangle).

affects the perturbative correction via the three-body matrix elements and the *e*3*^N* dependence becomes more pronounced. Generally, the step from MBPT(2B) to MBPT(2B + 3Bpphh) can modify the ground-state energy in either direction, whereas the change from MBPT($2B + 3B$ pphh) to MBPT($2B + 3B$) always results in a lowering of the ground-state energy, as evident from Eq. [\(15\)](#page-6-0).

For ⁴He we obtain a significantly lower ground-state energy when fully including the three-body terms. For the heavier nuclei the ground-state energy is increased at small e_{3N} and remains almost unchanged for larger *e*3*^N* . Generally, the change in the ground-state energy per nucleon when going from MBPT(2B) to MBPT(2B $+$ 3B) for fixed and sufficiently large e_{3N} decreases with increasing particle number. Beyond 40 Ca the impact of the three-body terms to the second-order energy correction is smaller than other uncertainties of the calculation, e.g., the degree of convergence with respect to the model space. Therefore, we will limit ourselves to the MBPT(2B) corrections and will continue using $e_{3N} = 20$ in the following. One should keep in mind, however, that for nuclei below 40 Ca and in particular for 4 He the full second-order correction MBPT $(2B + 3B)$ leads to a lower ground-state energy than MBPT(2B) and thus to a much better agreement with experiment.

As discussed earlier, we can fix the strength of the threebody interaction based on the systematics of the charge radii and use the flow parameter entering into the two-body interaction to control the binding-energy systematics. In Fig. 8 we illustrate the influence of α on the energies obtained in MBPT(2B) using the S-UCOM(SRG) interaction with $\alpha = 0.12$ fm⁴ and 0.16 fm⁴ and a 3*N* contact interaction with $C_{3N} = 2200 \text{ MeV fm}^6$. Whereas the charge radii are practically identical for both values of α , the HF and the MBPT(2B) ground-state energies are systematically lower for the larger flow parameters. The difference is smaller for

FIG. 8. (Color online) Binding energies per nucleon and charge radii of selected closed-shell nuclei resulting from HF calculations (filled symbols) and MBPT (open symbols) based on the S-UCOM(SRG) interaction for $e_{\text{max}} = 10$, $C_{3N} = 2200 \text{ MeV fm}^6$, $e_{3N} = 20$ and different flow parameters: $\alpha = 0.12$ fm⁴ (\bullet), $\alpha =$ 0.16 fm⁴ (\blacksquare). The bars indicate the experimental values [\[18,19\]](#page-9-0).

FIG. 9. (Color online) Binding energies per nucleon and charge radii of selected closed-shell nuclei resulting from HF calculations (filled symbols) and MBPT (open symbols) based on the S-UCOM(SRG) interaction for $\alpha = 0.16$ fm⁴, $C_{3N} = 2200$ MeV fm⁶, $e_{3N} = 20$ and different basis sizes: $e_{\text{max}} = 10$ (\bullet); $e_{\text{max}} = 12$, $l_{\text{max}} =$ 10 (\blacksquare); $e_{\text{max}} = 14$, $l_{\text{max}} = 10$ (\blacktriangle). The bars indicate the experimental values [\[18,19\]](#page-9-0).

the MBPT(2B) energies than for the HF energies, as to be expected. The unitary transformation for larger *α* accounts for more of the correlations explicitly, thus the ground-state energy at the HF level is lower and the gain due to the inclusion of residual correlations through MBPT(2B) is smaller. Thus we can still use the α dependence to control the binding-energy systematics.

So far we have used a model space with $e_{\text{max}} = 10$ for all calculations. This is absolutely sufficient to obtain converged HF results, but it is not enough to obtain converged HF + MBPT energies. The convergence behavior of the $HF + MBPT(2B)$ energies is illustrated in Fig. 9 for the S-UCOM(SRG) interaction with $\alpha = 0.16$ fm⁴ and in Fig. [10](#page-8-0) for the S-SRG interaction with $\alpha = 0.10$ fm⁴ using $e_{\text{max}} =$ 10*,* 12*,* and 14. The HF energies and charge radii are fully converged in all cases but not the $HF + MBPT(2B)$ energies. Although the largest model space includes 15 major oscillator shells, this is still not sufficient to obtain convergence of the MBPT(2B) contribution for heavier nuclei. The change of energy per nucleon between successive model-space sizes increases with increasing nucleon number.

The slow convergence is partly due to the use of *S*-wave only UCOM and SRG transformations for the construction of the two-nucleon interactions. For all higher-partial waves the transformed interactions are thus identical to the initial interaction. Since those partial waves become increasingly important for heavier nuclei, the deterioration of the convergence is unsurprising. The use of a softer all-channel transformed interactions, e.g., the standard SRG interaction discussed in Fig. [2,](#page-4-0) would help with the convergence. However, for those interactions a simple contact force is not sufficient to provide reasonable energy and radius systematics.

Despite the nonoptimal convergence, the results presented in Figs. 9 and [10](#page-8-0) show that after inclusion of beyond-HF

FIG. 10. (Color online) Binding energies per nucleon and charge radii of selected closed-shell nuclei resulting from HF calculations (filled symbols) and MBPT (open symbols) based on the S-SRG interaction for $\alpha = 0.10 \text{ fm}^4$, $C_{3N} = 2000 \text{ MeV fm}^6$, $e_{3N} = 20$ and different basis sizes: $e_{\text{max}} = 10$ (\bullet); $e_{\text{max}} = 12$, $l_{\text{max}} = 10$ (\bullet); $e_{\text{max}} = 14$, $l_{\text{max}} = 10$ (\triangle). The bars indicate the experimental values [\[18,19\]](#page-9-0).

correlations also the systematics of the ground-state energy is reproduced quite well. Given the additional gain in binding energy that is expected until convergence with respect to model-space size and the additional binding resulting from the three-body contributions to the second-order correction for light isotopes the energies are in good systematic agreement with experiment. For the results presented in Fig. 10 an exponential extrapolation $e_{\text{max}} \rightarrow \infty$ typically leads to binding energies per nucleon that are within a few tenths on an MeV of the experimental data. The charge radii follow the experimental results very closely already at the HF level. Perturbative corrections to the radii, as studied in Ref. [\[16\]](#page-9-0), are very small and will not affect the general agreement.

IV. CONCLUSIONS AND OUTLOOK

We have investigated the systematics of binding energies and charge radii for closed-shell nuclei from 4 He to 208 Pb starting from unitarily transformed realistic *NN* interactions supplemented by phenomenological 3*N* forces. We have shown that already a simplistic 3*N* contact interaction is

sufficient to cure the systematic deviations from experiment that the UCOM- and SRG-transformed two-body interactions exhibit. By supplementing an S-UCOM or S-SRGtransformed interaction for sufficiently large flow parameters *α* with a repulsive 3*N* contact interaction we were able to reproduce the experimental charge radius and ground-state energy systematics simultaneously. Only for cases where the two-body interaction exhibits a pathological energy systematics a contact force is clearly not sufficient to arrive at a reasonable behavior.

In a next step, we can apply these interactions in a variety of many-body schemes and study the elementary effects of 3*N* interactions on nuclear observables. We will employ the phenomenological 3*N* interactions in exact no-core shellmodel calculations for the spectroscopy of light nuclei and in approximate many-body approaches such as RPA for the study of the collective response of heavier nuclei. The fundamental advantage of 3*N* contact interactions is that the computation of three-body matrix elements itself is not a limiting factor and thus a larger range of nuclei and observables can be explored.

A major goal, however, it to go beyond phenomenological 3*N* interactions for supplementing a unitarily transformed *NN* interaction toward a consistent two- plus three-body interaction resulting from a combined unitary transformation of an initial two- plus three-body interaction. This can be done, e.g., using an SRG evolution of the chiral *NN* plus 3*N* interaction. A study of the ground-state energy and radius systematics for those interactions, even in a simple framework like $HF + MBPT$, will provide crucial information on the quality of the presently available chiral interactions for nuclear structure studies beyond the light isotopes and it will serve as a test-case for the SRG evolution in the *NN* plus 3*N* sector. Based on our developments for phenomenological 3*N* interactions discussed here, we will perform a similar analysis with fully realistic *NN* plus 3*N* interactions next.

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