Abstract. Extrapolations in Euclidean time form a central part of Nuclear Lattice Effective Field Theory (NLEFT) calculations using the Projection Monte Carlo method, as the sign problem in many cases prevents simulations at large Euclidean time. We review the next-to-next-to-leading order NLEFT results for the alpha nuclei up to $^{28}$Si, with emphasis on the Euclidean time extrapolations, their expected accuracy and potential pitfalls. We also discuss possible avenues for improving the reliability of Euclidean time extrapolations in NLEFT.

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1. Introduction

Several ab initio methods are currently being applied to the study of nuclear structure. These include coupled-cluster expansions [1], the no-core shell model [2, 3], the in-medium similarity renormalization group [4], self-consistent Green’s functions [5], Green’s function Monte Carlo [6], and Auxiliary-Field Diffusion Monte Carlo [7].

Much of the recent progress in ab initio nuclear structure calculations is due to “soft” chiral nuclear EFT interactions. The lattice formulation of chiral nuclear EFT is described in Ref. [8], and a review of lattice EFT methods can be found in Ref. [9]. A comprehensive overview of chiral nuclear EFT is available in Refs. [10, 11]. This framework, known as Nuclear Lattice Effective Field Theory (NLEFT), has been used to calculate the ground states of alpha nuclei from $^4$He to $^{28}$Si, as well as to describe the structure of the Hoyle state [12-13] and the dependence of the triple-alpha process on the fundamental parameters of nature [14].

NLEFT is an ab initio method where chiral nuclear EFT is combined with Auxiliary-Field Quantum Monte Carlo (AFQMC) lattice calculations. This Monte Carlo approach differs from other ab initio methods in that it does not require truncated basis expansions, many-body perturbation theory, or any constraint on the nuclear wave function. While our NLEFT results are thus unbiased Monte Carlo calculations, the AFQMC approach nevertheless relies on Euclidean time projection of Projection Monte Carlo (PMC) data in order to compute the properties of the low-lying states of light and medium-mass nuclei. One of the largest sources of computational uncertainty is then due to the appearance of complex sign oscillations or the “sign problem” which limits the extent of Euclidean time available for direct PMC calculations. While the sign problem is greatly suppressed by the soft interaction employed in NLEFT, it still represents a significant obstacle to practical PMC calculations, especially in cases where the number of protons is not equal to the number of neutrons.

In this paper, we focus on the question of the accuracy and reliability of the Euclidean time extrapolations. We start in Section 2 by reviewing the PMC formalism, along with the methodology for extrapolating to infinite Euclidean time in Section 3. Next, we provide in Section 4 an overview of the NLEFT results for the alpha nuclei ranging from $^4$He to $^{28}$Si, as obtained from Euclidean time extrapolations of PMC data corresponding to the lattice action described in Refs. [12-13, 15]. In Section 5 we consider the effects of statistical and systematical errors on the accuracy of the extrapolation method, and we conclude in Section 6 by a discussion of future improvements and refinements to our extrapolation methods.

2. Projection Monte Carlo formalism

Our NLEFT calculations are, as in chiral nuclear EFT, organized in powers of a soft scale $Q$, which is associated with factors of momenta and the pion mass. The contributions of $O(Q^0)$ to the nuclear Hamiltonian are referred to as leading order (LO), the $O(Q^2)$ terms are of next-to-leading order (NLO), and the $O(Q^3)$ terms are of next-to-next-to-leading order (NNLO), at which point our present calculations are truncated. The LO lattice Hamiltonian includes a significant part of the NLO and higher-order contributions, as smeared contact interactions $^{8, 16, 17}$ are used. It should also be noted that since we are using a low-momentum power counting scheme, no additional two-nucleon contributions arise at NNLO beyond the terms already appearing at NLO, as these can be absorbed into redefinitions of the NLO couplings. A full discussion of the interactions used in the reported results can be found in Ref. [15].
In the present NLEFT calculations, we have used a periodic $L = 6$ cube and a lattice
spacing of $a = 1.97$ fm, which translates into a cube length of $La = 11.82$ fm. Our initial
wave functions, $|\Psi_A^{init}\rangle$, are Slater-determinant states composed of delocalized standing waves
in the periodic cube with $A$ nucleons. Localized alpha-cluster trial states have also been used
for studies of $^{12}$C and $^{16}$O [13, 18]. These provide not only a consistency check on the
Euclidean time extrapolation, but also an opportunity to assess the spatial structure of the
nuclei. For simplicity, we describe our calculations using the language of continuous time
evolution, even though our AFQMC calculations use transfer matrices with a temporal lattice
spacing of $a_t = 1.32$ fm [9]. The Euclidean projection time is given by $t = N_t a_t$, where $N_t$
denotes the number of Euclidean time slices.

We start the Euclidean time projection by means of a “low-energy filter” based upon
Wigner’s SU(4) symmetry, where the spin and isospin degrees of freedom of the nucleon are
all equivalent as four components of an SU(4) multiplet. The SU(4) symmetric Hamiltonian
is of the form

$$ H_{SU(4)} = H_{\text{free}} + \frac{1}{2} C_{SU(4)} \sum_{\vec{n},\vec{n}'} \rho(\vec{n}) f(\vec{n} - \vec{n}') \rho(\vec{n}') , $$

where $f(\vec{n} - \vec{n}')$ is a Gaussian smearing function with its width set by the average effective
range of the two $S$-wave interaction channels, and $\rho$ is the total nucleon density. Application
of the exponential of $H_{SU(4)}$ gives

$$ |\Psi_A(t')\rangle \equiv \exp(-H_{SU(4)} t') |\Psi_A^{init}\rangle , $$

referred to as a “trial state”. This part of the calculation is computationally straightforward,
as it only requires a single auxiliary field. Most significantly, it does not generate any sign
oscillations in the Monte Carlo calculation.

Next, we use the full LO Hamiltonian $H_{LO}$ to evolve the trial state for a time $t$, and
construct the Euclidean-time projection amplitude

$$ Z_A(t) \equiv \langle \Psi_A(t') | \exp(-H_{LO} t) | \Psi_A(t') \rangle , $$

from which we compute the “transient energy”

$$ E_A(t) = -\partial [\ln Z_A(t)] / \partial t , $$

by means of a numerical finite difference. Hence, if the lowest eigenstate of $H_{LO}$ that
possesses a non-vanishing overlap with the trial state $|\Psi_A(t')\rangle$ is denoted $|\Psi_A,0\rangle$, the energy
$E_{A,0}$ of $|\Psi_A,0\rangle$ is obtained as the $t \rightarrow \infty$ limit of $E_A(t)$. Sign oscillations in the Monte Carlo
calculation set the main limitation on the number of Euclidean time steps for which Eq. (4) can
be evaluated. The coupling $C_{SU(4)}$ is a free parameter which can be used either to optimize
the convergence of the Euclidean time evolution, or to provide additional constraints for the
extrapolation $t \rightarrow \infty$.

Higher-order corrections to $E_{A,0}$ are evaluated using perturbation theory. We compute
expectation values using

$$ Z_A^\mathcal{O}(t) \equiv \langle \Psi_A(t') | \exp(-H_{LO} t/2) \mathcal{O} \exp(-H_{LO} t/2) | \Psi_A(t') \rangle , $$

for any operator $\mathcal{O}$. Given the ratio

$$ X_A^\mathcal{O}(t) = Z_A^\mathcal{O}(t) / Z_A(t) , $$

the expectation value of $\mathcal{O}$ for the desired state $|\Psi_A,0\rangle$ is again obtained in the $t \rightarrow \infty$ limit
according to

$$ X_{A,0}^\mathcal{O} \equiv \langle \Psi_A,0 | \mathcal{O} | \Psi_A,0 \rangle = \lim_{t \rightarrow \infty} X_A^\mathcal{O}(t) , $$
which gives the corrections to $E_{A,0}$ induced by the NLO and NNLO contributions, including the effects of strong and electromagnetic isospin symmetry breaking.

The closer the trial state $|\Psi_A(t')\rangle$ is to $|\Psi_{A,0}\rangle$, the less the required projection time $t$. The trial state can be optimized by adjusting both the SU(4) projection time $t'$ and the strength of the coupling $C_{SU(4)}$ of $H_{SU(4)}$. As shown in Section 2, the accuracy and reliability of the extrapolation $t \to \infty$ is greatly improved by simultaneously incorporating data from multiple trial states that differ in the choice of $C_{SU(4)}$. This approach enables a “triangulation” of the asymptotic behavior as the common limit of several different functions of $t$.

3. Extrapolation in Euclidean time

In most cases, reaching the limit $t \to \infty$ requires an extrapolation from finite values of $t$. Given the limited extent of the data, there are uncertainties in this extrapolation. The behavior of $Z_A(t)$ and $Z^O_A(t)$ at large $t$ is controlled by the low-energy spectrum of $H_{LO}$. Let $|E\rangle$ label the eigenstates of $H_{LO}$ with energy $E$, and let $\rho_A(E)$ denote the density of states for a system of $A$ nucleons. For simplicity, we omit additional labels needed to distinguish degenerate states. We can then express $Z_A(t)$ and $Z^O_A(t)$ in terms of their spectral representations,

$$Z_A(t) = \int dE \, \rho_A(E) \left| \langle E|\Psi_A(t')\rangle \right|^2 \exp(-Et),$$

$$Z^O_A(t) = \int dE \, dE' \, \rho_A(E) \rho_A(E') \exp(-(E + E')t/2),$$

$$\times \langle \Psi_A(t')|E\rangle \langle E|O|E'\rangle \langle E'|\Psi_A(t')\rangle.$$  \hspace{1cm} (9)

The spectral representations of $E_A(t)$ and $X^O_A(t)$ are then obtained by using Eq. (4) and Eq. (6), respectively. We can approximate these to arbitrary accuracy over any finite range of $t$ by taking $\rho_A(E)$ to be a sum of energy delta functions,

$$\rho_A(E) \approx \sum_{k=0}^{k_{\text{max}}} c_{A,k} \delta(E - E_{A,k}).$$ \hspace{1cm} (10)

Let us now consider the extrapolation formulas for the LO energy and the higher order perturbative corrections. From Eqs. (9) and (10), we find for the LO energy

$$E_{A}^j(N_t) = E_{A,0} + \sum_{k=1}^{k_{\text{max}}} |c_{A,j,k}| \exp \left( -\frac{\Delta_{A,k} N_t}{\Lambda_t} \right),$$

where $t = N_t/\Lambda_t$ with $\Lambda_t = 150$ MeV, corresponding to $a_t = 1.32$ fm. The energy gaps are defined as $\Delta_{A,k} \equiv E_{A,k} - E_{A,0}$, and the index $j$ denotes a specific choice of $t'$ and $C_{SU(4)}$ in the trial wave function $|\Psi_A(t')\rangle$. We take $\Delta_{A,k}^{k+1} > \Delta_{A,k}$, and $k_{\text{max}} = 3$ for $^4$He ($A = 4$) and $k_{\text{max}} = 2$ for $A \geq 8$. For the operator matrix elements that make up the perturbative NLO and NNLO corrections, we find

$$X_{A}^O,N_t = X_{A,0}^O + \sum_{k=1}^{k_{\text{max}}} x_{A,j,k} \exp \left( -\frac{\Delta_{A,k} N_t}{2\Lambda_t} \right),$$

where the dominant contributions are taken to be due to transition amplitudes involving the ground state and excited states. In order for this to be a good approximation, it is necessary that the overlap between our trial state and the ground state not be small compared to the overlap with the low-lying excited states. It should be noted that the coefficients $x_{A,j,k}$ can be positive as well as negative, which gives us the possibility of “triangulating” the asymptotic
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and all NLO and NNLO matrix elements in NLEFT is performed for each value of $t$ and below is possible. However, we are helped by the fact that the rate of convergence is twice that of Eq. (12). In order to determine $\Delta_{A,t}$ is monotonically decreasing and thus no triangulation of the asymptotic value from above and below is possible. However, we are helped by the fact that the rate of convergence is twice that of Eq. (14). In order to determine $E_{A,0}$ and $X_{A,0}^{\gamma}$, a correlated $\chi^2$ fit to the LO energy and all NLO and NNLO matrix elements in NLEFT is performed for each value of $A$. This procedure also determines the coefficients $c_{A,j,k}, \Delta_{A,j,k}$, and $X_{A,0}^{\gamma}$. We find that using 2 to 6 distinct trial states for each $A$ allows for a significantly more accurate and stable determination of $E_{A,0}$ and $X_{A,0}^{\gamma}$ than would be possible with a single trial state. Note that the energy gaps

Figure 1. Extrapolation of NLEFT results for $^3$He with $k_{\text{max}} = 3$. The definitions of the observables are given in the main text. The LO energy is $E_{\text{LO}} = -28.87(6)$ MeV, and the result at NNLO including effective 4N interactions (as reported in Ref. [19]) is $E_{\text{NNLO+4N}} = -28.93(7)$ MeV. The empirical binding energy is $-28.30$ MeV. For each trial state, the value of $C_{\text{SU}(4)}$ is given in units of the (spatial) lattice spacing.
fuclidean Time Extrapolation in Lattice Effective Field Theory /\[\text{Fig. 1 and 2, respectively.}

Further, our results for the light nuclei \(A, k\) are confronted with actual PMC data. First, we show our data for the light nuclei \(\text{He}^4\) and \(\text{Be}^8\) in Figs. 1 and 2, respectively. Our results for the light nuclei \(\text{He}^4\) and \(\text{Be}^8\) are shown in Figs. 1 and 2, respectively. Further, our results for \(\text{C}^{12}\) are given in Fig. 3 for \(\text{O}^{16}\) in Fig. 4.

\[\Delta A, k\] in the extrapolation functions are taken to be independent of the trial wave function \(j\), which gives an additional consistency criterion. We find that a simultaneous description using Eqs. (11) and (12) accounts for all of the PMC data we have obtained for different \(\langle \Psi_A (\ell') \rangle\).

4. Analysis of Projection Monte Carlo data

We shall next elaborate on how the extrapolation methods of Section 3 perform when confronted with actual PMC data. First, we show our data for the light nuclei \(\text{He}^4\) and \(\text{Be}^8\) in Figs. 1 and 2, respectively. Further, our results for \(\text{C}^{12}\) are given in Fig. 3 for \(\text{O}^{16}\) in Fig. 4.
for $^{20}\text{Ne}$ in Fig. 3 for $^{24}\text{Mg}$ in Fig. 4 and for $^{28}\text{Si}$ in Fig. 7. The curves show a correlated $\chi^2$ fit for all trial states with a given $A$, using the same spectral density $\rho_A(E)$. The upper row in each figure shows the LO energy, the total isospin-symmetric 2NF correction (NLO), the electromagnetic and isospin-breaking corrections (EMIB) and the total 3NF correction. The remaining panels show the matrix elements $X_A^C(t)$ that form part of the NLO and 3NF terms.

In the second and third rows of Fig. 1 through Fig. 4 the operators $\partial E_A/\partial C_i$ give the contributions of the NLO contact interactions. The interactions that involve $C_0$, $C_{12}$, $C_{Q2}$, $C_{P2Q2}$, $C_{S2Q2}$, $C_{S2FQ2}$, $C_{Q2}^2$ and $C_{P2Q2}^2$ are defined in Eqs. (18) to (23) of Ref. [17]. Similarly, the interactions involving $C_{S2Q2SO(3)B}$ and $C_{P2Q2SO(3)B}$ are given in Eqs. (55),...
and (56) of Ref. [17], and \(\Delta E_A(\Delta x_c)\) denotes the energy shift due the \(O(a^2)\)-improved pion-nucleon coupling in Eq. (57) of Ref. [17]. The operators \(\partial E_A/\partial D_i\) give the individual contributions to the 3NF correction, which are defined in Eqs. (37) to (41) of Ref. [17].

The error estimates given in parentheses in the captions of Fig. 1 through Fig. 7 have been obtained from a \(\chi^2\) minimization using the PMC errors for each datapoint as weight factors. This procedure could be affected by autocorrelations in Monte Carlo time, and secondly by the fact that the observables (except for the LO energy) are formed out of a ratio of amplitudes for which the PMC error is known separately, according to Eq. (6). Instead of simply adding these PMC errors in quadrature, a more realistic error estimate could be obtained by means of.
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Figure 5. Extrapolation of NLEFT results for $^{20}$Ne with $k_{\text{max}} = 2$. The definitions of the observables are given in the main text. The LO energy is $E_{\text{LO}} = -199.7(9)$ MeV, and the result at NNLO including effective 4N interactions (as reported in Ref. [19]) is $E_{\text{NNLO+4N}} = -165.9(9)$ MeV. The empirical binding energy is $-160.64$ MeV. For each trial state, the value of $C_{\text{SU(4)}}$ is given in units of the (spatial) lattice spacing.

A resampling algorithm such as the Jackknife or the Bootstrap (for a pedagogical introduction, see e.g. Ref. [20]). In order to decrease the effects of any residual autocorrelations in the PMC data, the Jackknife method can be combined with “blocking” of the data, whereby adjacent (in Monte Carlo time) samples are combined into blocks of increasing size until the variance of the sample converges as a function of the block size (see e.g. Ref. [21]).

In our PMC production runs, we do not use $k_{\text{max}} = 1$ (corresponding to a single energy gap) as this would in most cases lead to large values of $\chi^2$ as well as inconsistent results for different choices of trial states (see Table I). Extrapolations with $k_{\text{max}} = 2$ account very well for the medium-mass nuclei, which appear to be highly compact objects for which
the contamination from low-lying excited states is small. For $^4$He, we find that $k_{\text{max}} = 3$ is required in order to account for all the PMC data for all observables and all trial states. However, while extrapolations with $k_{\text{max}} = 3$ may provide a more accurate description over a larger range of trial states, such fits are also much more difficult to constrain adequately, due to the much larger number of adjustable parameters involved. The extent and resolution of our PMC data in Euclidean time also limits, in most cases, the number of resolvable energy gaps to $k_{\text{max}} = 2$. 

Figure 6. Extrapolation of NLEFT results for $^{24}$Mg with $k_{\text{max}} = 2$. The definitions of the observables are given in the main text. The LO energy is $E_{\text{LO}} = -253(2)$ MeV, and the result at NNLO including effective 4N interactions (as reported in Ref. [19]) is $E_{\text{NNLO+4N}} = -198(2)$ MeV. The empirical binding energy is $-198.26$ MeV. For each trial state, the value of $C_{\text{SU}(4)}$ is given in units of the (spatial) lattice spacing.
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Figure 7. Extrapolation of NLEFT results for $^{28}$Si with $k_{\text{max}} = 2$. The definitions of the observables are given in the main text. The LO energy is $E_{\text{LO}} = -330(3)$ MeV, and the result at NNLO including effective 4N interactions (as reported in Ref. [19]) is $E_{\text{NNLO+4N}} = -233(3)$ MeV. The empirical binding energy is $-230.54$ MeV. For each trial state, the value of $C_{\text{SU}(4)}$ is given in units of the (spatial) lattice spacing.

5. Statistical and systematic errors

Our extrapolation procedure is examined in detail for the case of $^{12}$C in Table 1. Our main fit, labeled “fit 5”, is also shown in Fig. 3 and consists of a simultaneous fit to five trial states that differ in the value of $C_{\text{SU}(4)}$. We observe that $\chi^2 \approx 0.68$, which indicates that the error bars of the individual MC data points are likely to be overestimated. This could plausibly happen as the observables are formed from the ratio of two amplitudes, the errors of which are at present simply added in quadrature. A full-fledged jackknife error analysis may yield a more realistic result. The uncertainties shown in parentheses correspond to the variances reported by the $\chi^2$.
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Table 1. Uncertainty analysis of the Euclidean time extrapolation for $^{12}$C with $k_{\text{max}} = 2$. The values of $C_{SU(4)}$ (in MeV$^{-1}$) for each trial state shown in Fig. 3 are “1” = $-8.0 \times 10^5$, “2” = $-1.2 \times 10^4$, “9” = $-7.5 \times 10^5$, “10” = $-6.5 \times 10^3$, and “11” = $-7.0 \times 10^5$. The quantities shown (in MeV) are: The LO non-perturbative 2NF result, followed by the perturbative higher-order and isospin-breaking corrections as described in the main text. The fit labeled “5” (shown in Fig. 3) is a correlated extrapolation using all trial states. The fits labeled “4a” - “4e” check the consistency of fit “5” under the removal of a single trial state from the full analysis. We also show the (poorly constrained) extrapolations “1a” - “1e” where each trial state is treated separately. Note that this only allows for an analysis with $k_{\text{max}} = 1$. The one-standard deviation error estimates (given in parentheses) are obtained from a Marquardt-Levenberg minimization of $\chi^2$ (per d.o.f.) with the Monte Carlo error estimates used as weights.

| Fit | Trial states | LO (2NF) | NLO (2NF) | EMIB (2NF) | NNLO (3NF) | $\chi^2$ |
|-----|--------------|-----------|-----------|------------|------------|---------|
| 5   | 1, 2, 9, 10, 11 | -96.9(2)  | 10.48(3)  | 7.76(1)   | -14.80(6)  | 0.68    |
| 4a  | 1, 2, 9, 10  | -96.8(2)  | 10.46(4)  | 7.76(1)   | -14.84(6)  | 0.71    |
| 4b  | 1, 2, 9, 11  | -97.0(2)  | 10.45(3)  | 7.76(1)   | -14.85(6)  | 0.72    |
| 4c  | 1, 2, 10, 11 | -96.8(2)  | 10.46(4)  | 7.76(2)   | -14.80(6)  | 0.72    |
| 4d  | 1, 9, 10, 11 | -97.3(2)  | 10.54(5)  | 7.72(2)   | -14.61(6)  | 0.57    |
| 4e  | 2, 9, 10, 11 | -96.9(2)  | 10.44(4)  | 7.75(2)   | -14.94(7)  | 0.69    |
| 1a  | 1            | -95.0(2)  | 10.17(2)  | 7.79(1)   | -13.93(4)  | 1.83    |
| 1b  | 2            | -94.4(2)  | 10.55(2)  | 7.98(1)   | -14.46(5)  | 3.25    |
| 1c  | 9            | -93.5(2)  | 10.03(2)  | 7.72(1)   | -13.60(4)  | 1.14    |
| 1d  | 10           | -94.0(9)  | 9.10(9)   | 7.32(3)   | -13.73(16) | 2.36    |
| 1e  | 11           | -92.1(2)  | 9.83(2)   | 7.58(1)   | -13.34(4)  | 1.28    |

minimization. In order to assess the accuracy of these error estimates as well as the stability of the central values, we have generated a number of fits where each one of the five trial states has been excluded in turn, in the spirit of the Jackknife method [20]. These fits are denoted “4a” through “4e” in Table 1. Evidently, these agree closely with the full analysis, the largest discrepancy being due to the exclusion of trial state “2”, which generates the largest shift in the extrapolated values as well as a significant reduction in $\chi^2$. As is evident from Fig. 3, trial state “2” (denoted by blue open squares) is furthest away from the “triangulation point”, and may therefore not be completely described by an extrapolation with $k_{\text{max}} = 1$. An even more stable result might be obtained by replacing that trial state in the analysis with one which is closer to the triangulation point.

For comparison, we also show in Table 1 the results of independent, uncorrelated fits to each of the five trial states for $^{12}$C. These extrapolations do not benefit from the consistency requirements of the multi-trial state extrapolations, and furthermore these can only be taken to $k_{\text{max}} = 1$, as the extent of the data in Euclidean time is too short to constrain more than one energy gap. We observe that such extrapolations are clearly much less reliable, and suffer from several pronounced issues. One is the clear tendency for “spurious early convergence”, which is due to the lack of enforced independence on the value of $C_{SU(4)}$. We also observe that the extrapolated values as well as the $\chi^2$ fluctuate significantly between extrapolations to different trial states. Again, trial state “2” appears to be the most problematic, although we also find that fits with a smaller $\chi^2$ are no more reliable than those with a larger value. Neither do the variances produced by the $\chi^2$ minimization properly describe the uncertainties.

Also, as shown in Table 1, the total uncertainty is dominated by that of the LO contribution, which does not consist of a ratio of amplitudes according to Eq. 6. This relatively large extrapolation error is due to the appearance of the absolute values of the
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Figure 8. Euclidean time extrapolation for the deuteron with $k_{\text{max}} = 1$ in a periodic $L = 3$ cube with lattice spacing $a = 1.97$ fm, for the quantities given in Table 2. The PMC data points at small Euclidean times (indicated by open squares) have been excluded from the analysis in order to increase the stability of the results. The PMC data correspond to trial state "1" with $C_{\text{SU}(4)} = -8.0 \times 10^2$ MeV$^{-2}$. For full details and definitions, see Ref. [14].

coefficients $c_{A_{j,k}}$ in Eq. (11), which prevents a triangulation of the LO contribution. Nevertheless, the Jackknife method could be used to evaluate the effect of autocorrelations between consecutive auxiliary-field configurations in Monte Carlo time on the LO result. We find that the elimination of such autocorrelations from the start, by allowing for sufficient decorrelation time between consecutive measurements, does not present any difficulties. For the NLO and NNLO operators, more consistent and reliable error estimates for the individual
data points could clearly be obtained by Jackknife resampling of the Monte Carlo data. While such an analysis would not alter any of the conclusions concerning the stability and consistency of the Euclidean time extrapolations, we have investigated the likely outcome of a full Jackknife resampling of our data by performing multiple independent Monte Carlo runs for $^{12}$C with $N_t = 12$. For such runs, we obtain LO energies for $N_t = 11.0, 11.5$ and $12.5$ by means of a numerical finite difference. Similarly, for the higher-order corrections we compute matrix elements for $N_t = 11.5$ and $12.5$. As expected, we find that the uncertainties of the LO energies are accurately given by the Monte Carlo errors, whereas those of the individual NLO matrix elements appear overestimated by a factor of $\approx 2$ due to cancellations between the numerator and denominator of Eq. (6). This result is consistent with the values of $\chi^2 < 1$ obtained from the full extrapolation.

As the extent of our PMC data in Euclidean time is relatively short, we discuss next the expected reliability of our extrapolated results. Fortunately, the number of Euclidean time steps $N_t$ available for the extrapolation does not decrease drastically with the number of nucleons $A$. At this time, our method has been successfully applied to the spectrum, structure and electromagnetic properties of $^{12}$C in Refs. [13, 14], and to those of $^{16}$O in Ref. [15], where consistency between delocalized plane-wave and alpha-cluster trial wave functions was established. In our “triangulation” method, the extrapolation is very strongly constrained by the requirement that all observables, for all trial states, should be described by the same exponential dependence on the Euclidean projection time $t$. Rapid convergence in $t$ then translates into a small sensitivity to $C_{SU(4)}$ at large values of $t$, which helps to guard against “spurious early convergence”, where a smaller energy gap is overlooked.

Nevertheless, in the absence of consistency conditions on the extrapolations, we find that our method is accurate even for $k_{\text{max}} = 1$ when the leading energy gap is very large, such as for the deuteron in a periodic $L = 3$ cube with lattice spacing $a = 1.97$ fm (see Ref. [14]). In that case, the extrapolated results can be directly compared with Lanczos diagonalization, as shown in Table 1 and Fig. 8. In spite of this impressive agreement, we still need to consider the possibility that a sufficiently small energy gap can be missed in the extrapolation due to the limited extent and resolution of the PMC data in Euclidean time. Of all the results presented here, those for $^{8}$Be are likely to be most affected by the limited Euclidean projection time.
as the convergence to the ground state is clearly the slowest and the data do not allow for extrapolations beyond \( k_{\text{max}} = 2 \), unlike the case of \(^4\text{He}\), where up to three energy gaps could be constrained by the PMC data.

6. Summary and outlook

We have presented an overview of the techniques and the analysis used for Euclidean time extrapolations in NLEFT. The core issue is that, due to computational constraints, one must extract asymptotic values from a limited region in Euclidean projection time. In order to reduce the uncertainty of the extrapolation, we perform multi-exponential fits as prescribed by spectral decomposition for the asymptotic behavior of the projection amplitudes. We find that the fitted asymptotic values are greatly stabilized by using multiple initial states and observables. As examples of this analysis, we have shown energies at LO, NLO, and NNLO, as well as individual operator expectation values for the alpha nuclei up to \( A = 28 \).

While these results are promising, the methods we have presented leave much room for further improvement. In particular, the current analysis is not adequate for the case when our trial states have only a small overlap with the ground state. In such cases, the raw PMC data will be far from their asymptotic values and it is unlikely that combining several different sets of data will provide any significant improvement. Another unfavorable situation that can arise is when the sign oscillations are severe and the quality of the PMC data degrades very quickly with projection time.

In these more difficult scenarios, one can improve on the situation by performing coupled multi-channel projections that evolve more than one initial state simultaneously in Euclidean time. This is the same approach which is being used in a technique called the “adiabatic projection method” used for describing scattering states of clusters \([22, 23]\). As shown in Ref. \([23]\), when projecting \( n \) states the exponential convergence of the ground state energy in projection time is given by the excitation energy of the \( n^{th} \) excited state rather than by that of the first excited state. This has the potential to help significantly in circumventing the limited extent of the Euclidean time propagation.

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