Progress on the nuclear many-body problem has been hindered for decades because nucleon-nucleon (NN) potentials that reproduce elastic-scattering phase shifts typically exhibit strong short-range repulsion as well as a strong tensor force. This leads to strongly correlated many-body wave functions and highly nonperturbative few- and many-body systems. But recent work shows how a cutoff on relative momentum can be imposed and evolved to lower values using renormalization group (RG) methods, thus eliminating the troublesome high-momentum modes [1,2]. The evolved NN potentials are energy-independent and preserve two-nucleon observables for relative momenta up to the cutoff. Such potentials, known generically as \( V_{\text{low } \mathbf{k}} \), are more perturbative and generate much less correlated wave functions [2–7], vastly simplifying the many-body problem. However, a full RG evolution of essential few-body potentials has not yet been achieved.

An alternative path to decoupling high-momentum from low-momentum physics is the similarity renormalization group (SRG), which is based on unitary transformations that suppress off-diagonal matrix elements, forcing the Hamiltonian toward a band-diagonal form. A simple SRG transformation applied to nucleon-nucleon interactions leads to greatly improved convergence properties while preserving observables hindered for decades because nucleon-nucleon (NN) potentials

\[ H_{\text{rel}} = U(s)H U^{-1}(s) \equiv H_{\text{rel}} + V_{s}, \]

Choosing \( \eta(s) \) specifies the transformation. Here we make perhaps the simplest choice [10].

\[ \eta(s) = [T_{\text{rel}}, H_1], \]

which gives the flow equation,

\[ \frac{dH_s}{ds} = [[T_{\text{rel}}, H_1], H_1]. \]

Other choices will be studied elsewhere [14]. For any given partial wave in the space of relative momentum NN states, Eq. (5) means that the potential in momentum space evolves as (with normalization so that \( 1 = \frac{1}{\pi} \int_{0}^{\infty} |q|^2 dq |q| \) and in units where \( \hbar^2/M = 1 \)),

\[ \frac{dV_s(k, k')}{ds} = - (k^2 - k'^2)^2 V_s(k, k') + \frac{2}{\pi} \int_{0}^{\infty} q^2 dq \left( k^2 + k'^2 - 2q^2 \right) \times V_s(k, q) V_s(q, k'). \]

(The additional matrix structure of \( V_s \) in coupled channels such as \( ^3S_1 - ^3D_1 \) is implicit.) For matrix elements far from the diagonal, the first term on the right side of Eq. (6) evidently dominates and exponentially suppresses these elements as \( s \) increases. The parameter \( \lambda \equiv s^{-1/4} \) provides a measure of the spread of off-diagonal strength. Although further analytic analysis is possible, we instead turn to a numerical demonstration that the flow toward the diagonal is a general
result. By discretizing the relative momentum space on a grid of Gaussian integration points, we obtain a simple (but nonlinear) system of first-order coupled differential equations, with the boundary condition that $V_s(k, k')$ at the initial $s$ (or $\lambda$) is equal to the initial potential.

The evolution of the Hamiltonian according to Eq. (6) as $s$ increases (or $\lambda$ decreases) is illustrated in Fig. 1, using two initial chiral EFT potentials [12,13]. On the left in Fig. 1 is $^1S_0$ starting from the harder (600-MeV cutoff) potential from Ref. [12], which has significant strength near the high-momentum diagonal, and on the right is the $S$-wave part of the $^3S_1-^3D_1$ coupled channel starting from one of the potentials from Ref. [13], which has more far off-diagonal strength initially and comparatively weaker higher-momentum strength on the diagonal. The initial momentum-space potential differs significantly among interactions that are phase equivalent up to the $NN$ inelastic threshold, but these examples show characteristic features of the evolution in $\lambda$. In particular, we see a systematic suppression of off-diagonal strength, as anticipated, with the width of the diagonal scaling as $\lambda^2$. The same behavior is observed when evolving from conventional high-precision $NN$ potentials, such as Argonne $v_{18}$, or other (softer) chiral potentials [14,15].

Because the SRG transformation is unitary, observables are unchanged at all energies, up to numerical errors. This is shown by Fig. 2, in which phase shifts for the two chiral EFT potentials are plotted, including the values at high energies where they are not constrained by data (above $E_{lab} = 300$ MeV). For a given potential, there is no visible variation with $\lambda$. Similarly, the binding energy and asymptotic normalizations for the deuteron are independent of $\lambda$ [15].

As $\lambda$ is lowered, different initial potentials flow to similar forms at low momentum while remaining distinct at higher momentum. The low-momentum parts also become similar to $V_{low k}$ potentials. These observations are illustrated in Fig. 3 for two particular slices of the potentials from Fig. 1. They will be explored in much greater detail in Ref. [14].

We can quantify the perturbativeness of the potential as we evolve to lower $\lambda$ by using the eigenvalue analysis introduced long ago by Weinberg [16] and recently applied in an analysis of $V_{low k}$ potentials [6]. Consider the operator Born series for
The improvement in convergence is similar to that found with smoothly regulated $V_{\text{low }k}$ potentials evolved from chiral N$^3$LO potentials [7]. At finite density, analogous effects led to perturbative behavior in nuclear matter for $V_{\text{low }k}$ potentials [5]. Results for the $G$-matrix support a similar conclusion for the SRG potentials (see the Web site in Ref. [14] for pictures).

In Fig. 6, the calculations for different $\lambda$ converge to different values for the binding energy of the triton. This reflects the contributions of the omitted (and evolving) three-body interaction. The evolution with $\lambda$ of the binding energy with $NN$ interactions only, which is also the evolution of the net three-body contribution, follows a similar pattern to that seen with $V_{\text{low }k}$ [4,7]: a slow decrease as $\lambda$ decreases, reaching a minimum for $\lambda$ between 1.5 and 2 fm$^{-1}$, and then a rapid increase.

The consistent RG evolution of few-body interactions is an important unsolved problem for low-momentum potentials. In $V_{\text{low }k}$ calculations to date, an approximate evolution is made by fitting the leading chiral EFT three-body force at

\begin{equation}
\lambda \leq 0: \end{equation}

\[ T(E) = V_s + \frac{1}{E - T_{\text{rel}}} V_s + \ldots \]  

By finding the eigenvalues and eigenvectors of

\[ \frac{1}{E - T_{\text{rel}}} V_s |\Gamma_v\rangle = \eta_v(E) |\Gamma_v\rangle, \]

and then acting with $T(E)$ on the eigenvectors,

\[ T(E) |\Gamma_v\rangle = V_s |\Gamma_v\rangle(1 + \eta_v + \eta_v^2 + \ldots), \]

it follows that nonperturbative behavior at energy $E$ is signaled by one or more eigenvalues with $|\eta_v(E)| \geq 1$ [16]. (See Ref. [6] for a more detailed discussion in the context of evolving $V_{\text{low }k}$ potentials.)

It suffices for our purposes to consider a single energy (e.g., $E = 0$), and to consider only the negative eigenvalues, which are associated with the short-range repulsion. Weinberg eigenvalues at zero energy are shown as a function of $\lambda$ in Fig. 4 for the $1S_0$ channel and the $3S_1-3D_1$ coupled channel. In both channels, the large negative eigenvalues at large $\lambda$ reflect the repulsive core of the initial potentials. They rapidly evolve to small values as $\lambda$ decreases to 2 fm$^{-1}$ and below, as also observed with the corresponding $V_{\text{low }k}$ evolution [6]. However, the intermediate increase for the subleading eigenvalues in $1S_0$ is a new feature of the SRG that merits further study.

The more perturbative potentials at lower $\lambda$ induce weaker short-range correlations in few- and many-body wave functions, which in turn leads to greatly improved convergence in variational calculations. This is illustrated via calculations of the binding energy of the deuteron and triton by diagonalization in a harmonic oscillator basis, as shown in Figs. 5 and 6. For a fixed basis size, a more accurate estimate is obtained with smaller $\lambda$ or, conversely, at fixed $\lambda$ the convergence with basis size becomes more rapid. The improvement in convergence is

\begin{equation}
\text{FIG. 3. (Color online) Matrix elements of the evolved SRG potentials at } \lambda = 2 \text{ fm}^{-1} \text{ for } ^1S_0 \text{ (top, diagonal elements) and } ^3S_1 \text{ (bottom, off-diagonal elements) for the same initial potentials as in Fig. 1. Also shown is the } V_{\text{low }k} \text{ potential with a smooth (exponential) regulator for momentum cutoff } \Lambda = 2 \text{ fm}^{-1}, \text{ evolved from the two potentials (600 MeV above and 550/600 MeV below).}
\end{equation}

\begin{equation}
\text{FIG. 4. (Color online) The largest repulsive Weinberg eigenvalues as a function of } \lambda \text{ in the } ^1S_0 \text{ channel and the } ^3S_1-3D_1 \text{ coupled channel for the same initial potentials as in Fig. 1.}
\end{equation}

\begin{equation}
\text{FIG. 5. (Color online) The absolute error vs. } \lambda \text{ of the predicted deuteron binding energy from a variational calculation in a fixed-size basis of harmonic oscillators } (N_{\text{max ho}} \text{ excitations). The initial potential is from Ref. [13].}
\end{equation}
(Note: all of the potentials depend implicitly on s.) The relative kinetic energy operator $T_{rel}$ can be decomposed in three ways:

$$T_{rel} = T_{12} + T_3 = T_{13} + T_2 = T_{23} + T_1,$$

and $T_i$ commutes with $V_{jk}$,

$$[T_3, V_{12}] = [T_2, V_{13}] = [T_1, V_{23}] = 0,$$

so the commutators of $T_{rel}$ with the two-body potentials become

$$[T_{rel}, V_{12}] = [T_{12}, V_{12}],$$

and similarly for $V_{13}$ and $V_{23}$.

Because we define $T_{rel}$ to be independent of $s$, the SRG equation (5) for the three-body Hamiltonian $H_s$ simplifies to

$$\frac{dV_s}{ds} = \frac{dV_{12}}{ds} + \frac{dV_{13}}{ds} + \frac{dV_{23}}{ds} + \frac{dV_{123}}{ds} = [[T_{rel}, V_s], H_s],$$

with $V_s$ from Eq. (12). The corresponding equations for each of the two-body potentials (which are completely determined by their evolved matrix elements in the two-body systems, e.g., on a Gaussian momentum grid) are

$$\frac{dV_{12}}{ds} = [[T_{12}, V_{12}], (T_{12} + V_{12})],$$

and similarly for $V_{13}$ and $V_{23}$. After expanding Eq. (16) using Eq. (12) and the different decompositions of $T_{rel}$, it is straightforward to show using the equations for the two-body potentials that the derivatives of two-body potentials on the left side cancel precisely with terms on the right side, leaving

$$\frac{dV_{123}}{ds} = [[T_{12}, V_{12}], (T_3 + V_{13} + V_{23} + V_{123})] + [[T_{13}, V_{13}], (T_2 + V_{12} + V_{23} + V_{123})] + [[T_{23}, V_{23}], (T_1 + V_{12} + V_{13} + V_{123})] + [T_{rel}, V_{123}],$$

(18)

The importance of these cancellations is that they eliminate the “dangerous” delta functions, which make setting up the integral equations for the three-body system problematic [17]. We emphasize that the $s$ dependence of the two-body interactions on the right side of Eq. (18) is completely specified by solving the two-body problem in Eq. (6). This is in contrast to RG methods that run a cutoff on the total energy of the basis states (e.g., Lee-Suzuki and Bloch-Horowitz techniques). Such methods generate “multivalued” two-body interactions, in the sense that the RG evolution of two-body operators in $A > 2$ systems depends nonlocally on the excitation energies of the unlinked spectator particles [18,19].

Further simplifications of Eq. (18) follow from symmetrization and applying the Jacobi identity, but this form is sufficient to make clear that there are no disconnected pieces. The problem is thus reduced to the technical implementation of a momentum-space decomposition analogous to Eq. (6). Note in particular that the evolution of $V_{123}$ is carried out without ever having to solve a bound-state or scattering problem.
We have verified that this formalism does generate three-body interactions that leave eigenvalues invariant for simple model Hamiltonians, such as a two-level system of bosons. Work is in progress on proof-of-principle tests using one-dimensional many-body systems (to avoid angular momentum complications) and the three-dimensional nuclear problem will be tackled soon.

In summary, the SRG applied to nucleon-nucleon potentials works as advertised even for a simple choice of transformation, driving the Hamiltonian (in momentum space) toward the diagonal, making it more perturbative and more convergent in few-body calculations. There is much to explore, such as the nature of the decoupling of high- and low-energy physics implied by Fig. 1 (see Ref. [15] for demonstrations of decoupling and the evolution of an operator in the two-body space) and whether other choices of $\eta(s)$ instead of Eq. (4) could be more effective in making the Hamiltonian diagonal. For example, the replacement $T_{\text{rel}} \rightarrow H_d$, where $H_d$ is the diagonal part of the Hamiltonian, or some function of $T_{\text{rel}}$ are easily implemented. Most important is the consistent evolution of nuclear three-body operators [14].

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