AB INITIO OPTICAL POTENTIALS AND NUCLEON SCATTERING ON MEDIUM MASS NUCLEI

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(Received December 14, 2016)

We show the first results for the elastic scattering of neutrons off oxygen and calcium isotopes obtained from ab initio optical potentials. The potential is derived using self-consistent Green’s function theory (SCGF) with the saturating chiral interaction NNLO$_{\text{sat}}$. Our calculations are compared to available scattering data and show that it is possible to reproduce low-energy scattering observables in medium mass nuclei from first principles.

DOI:10.5506/APhysPolB.48.273

1. Introduction

Recent years have seen considerable advances in the theory of optical potentials. Non-locality effects have been seen to be necessary for describing three-body processes [1, 2], the importance of both scattering and bound states in the coupling to breakup channels has been explored [3], and global dispersive optical potentials have been developed [4].

The greatest challenge remains, however, the one of describing the nuclear structure and scattering consistently from the same theory. Many-body Green’s function methods are particularly suited to attempt this for medium and large nuclei since their central quantity, the self-energy, is naturally linked to the Feshbach theory of optical potentials [5, 6]. In particular, the particle part of the self-energy is equivalent to the original formulation

* Presented at the Zakopane Conference on Nuclear Physics “Extremes of the Nuclear Landscape”, Zakopane, Poland, August 28–September 4, 2016.
of Feshbach, while its hole part describes the structure of the target [7]. Nuclear field theory is one of the first (semi phenomenological) attempts to build such a theory for atomic nuclei [8, 9] and it has been extended to nuclear transfer reactions [10, 11]. Another incarnation of Green’s function related theories is the dispersive optical model [12], which is a data driven formulation of global (local and non-local) potentials constructed as the best possible parameterization of the microscopic self-energy [4, 13].

For transfer reactions, such as \((d, p)\), it would be particularly important to have an optical potential that is deduced consistently from the same Hamiltonian used in the proton–neutron channel [2]. To do so, one needs realistic nuclear interactions and \textit{ab initio} calculations of elastic nucleon–nucleus scattering. The no-core shell model with continuum (NCSMC) has been successful to calculate scattering and transfer reactions for light targets [14–16]. On the other hand, the self consistent Green’s function (SCGF) formalism [17, 18] is better suited to derive optical potentials for medium mass targets. SCGFs have been used to calculate phase shifts [19] and to investigate analytical properties of optical potentials [20]. However, these calculations were limited to two-body forces and a direct comparison to the experiment has been hindered by the lack of realistic interactions capable to reproduce accurately nuclear radii.

Three-body interactions have been recently implemented for SCGF in [21–23]. Moreover, the introduction of the NNLO\textsubscript{sat} interaction [24] has allowed a good reproduction of nuclear saturation and, hence, of radii and binding energies across the oxygen [25] and calcium chains [26]. Although this interaction has limitations regarding the symmetry energy in neutron-rich nuclei, we are now in the position to make a meaningful comparison of first principle approaches to scattering data. Here, we perform state of the art SCGF calculations to test the quality of current \textit{ab initio} methods and of the NNLO\textsubscript{sat} Hamiltonian in predicting elastic scattering.

2. The microscopic optical potential

The irreducible self-energy, \(\Sigma^{*}(\omega)\), has the general spectral representation

\[
\Sigma^{*}_{\alpha\beta}(E) = \Sigma^{(\infty)}_{\alpha\beta} + \sum_{i,j} M^{\dagger}_{\alpha,i} \left[ \frac{1}{E - (K^{>} + C) + i\eta} \right]_{i,j} M_{j,\beta} \\
+ \sum_{r,s} N_{\alpha,r} \left[ \frac{1}{E - (K^{<} + D) - i\eta} \right]_{r,s} N^{\dagger}_{s,\beta},
\]

(1)

where \(\Sigma^{(\infty)}_{\alpha\beta}\) is the correlated and energy-independent mean field. We perform calculations with the third order algebraic diagrammatic construction
[ADC(3)] method, where the matrices $M(N)$ couple single particle states to intermediate 2p1h (2h1p) configurations, $C$ ($D$) are interaction matrices among these configurations and $K$ are their unperturbed energies.

We use a spherical harmonic oscillator basis consisting of $N_{\text{max}}+1$ oscillator shells, so the optical potential for a given partial wave $(l,j)$ is expressed in terms of the oscillator radial functions $R_{n,l}(r)$ as

$$\Sigma^{*l,j}(r, r'; E) = \sum_{n,n'} R_{n,l}(r) \Sigma^{*l,j}_{n,n'}(E) R_{n',l}(r'),$$

which is non-local and depends on energy, angular momentum and parity. We solve the corresponding scattering problem in the full one-body space (so that the kinetic energy is treated exactly, without truncations) and account for the non-locality and $l,j$ dependence of Eq. (2). For each partial wave and parity, the phase shifts $\delta(E)$ are obtained as a function of the projectile energy, from where the differential cross section is calculated. We show results for incident energies in the laboratory frame, except for Fig. 4 below.

3. Results

In the following, we consider the volume integrals of the real ($J_V$) and imaginary ($J_W$) parts of the self-energy (i.e., the optical potential):

$$J_V(E) = 4\pi \int dr r^2 \int dr' r'^2 \sum_{l,j} \Re \left\{ \Sigma^{*l,j}(r, r'; E) \right\},$$

$$J_W(E) = 4\pi \int dr r^2 \int dr' r'^2 \sum_{l,j} \Im \left\{ \Sigma^{*l,j}(r, r'; E) \right\},$$

since these are strongly constrained by experimental data [6].

Figure 1 shows the volume integrals of the neutron-\(^{16}\)O for different model space truncations. Both the part of the self-energy below the Fermi surface (which describes the structure of the target) and the resonant structures for scattering at low energy are substantially converged already for $N_{\text{max}} = 11$. The oscillations seen at higher energies ($E > 10$ MeV) are an artefact of using a discretised model space and keep changing with $N_{\text{max}}$. They can fade away for an infinite space, or by exploiting a basis with the continuum.

Figure 2 shows $J_W$ for selected closed sub-shell Ca isotopes. The gap at the Fermi surface, where $\Im \Sigma^*(E) = 0$, shifts to higher energies and eventually crosses the continuum threshold with increasing neutron number. Compared to previous calculations using the Argonne $v_{18}$ and N$^3$LO(500) interactions [20], the NNLO$\text{sat}$ predicts an increased level density in the proximity of the Fermi energy, as expected for a correct nuclear saturation.
Fig. 1. Volume integrals of the real (left) and imaginary (right) parts of the neutron-\(^{16}\)O optical potential calculated for different numbers of oscillator shells in the model space: \(N_{\text{max}} = 7\) (dotted), 11 (dashed) and 13 (solid lines). Note that \(\Im\{\Sigma^{*}(E = E_{F})\} = 0\), so \(J_{W}(E = E_{F}) = 0\), where \(E_{F}\) is the Fermi energy. Thus, the potential for particle (holes) states is above (below) the gap in the \(J_{W}\) plot.

Fig. 2. Volume integral of the imaginary part of the neutron optical potentials, \(J_{W}(E)\), for the \(^{40}\)Ca, \(^{48}\)Ca and \(^{54}\)Ca targets. Calculated at \(N_{\text{max}} = 11\).

In Fig. 3, the neutron \(s_{1/2}\) phase shifts for \(^{16}\)O is shown for \(N_{\text{max}} = 11\) and 13. The resonance at \(E \approx 5\) MeV is almost converged for these spaces. Note that this state is dominated by \(2p1h\) components and thus it can still be affected by many-body truncations. The wiggles computed for energies \(E > 8\) MeV are due to similar but very narrow resonances. Again, these are linked to the discretisation of the model space and drift when increasing the number of oscillator shells. The right panel of Fig. 3 shows the phase shifts for other representative partial waves. The \(p_{1/2}\) has a sub-threshold bound state, while there is a very narrow \(f_{7/2}\) that is also seen experimentally within 1 MeV of our calculation \([27]\). The dominant \(d_{3/2}\) resonance is converged with respect to \(N_{\text{max}}\) and it is computed at \(\approx 1.15\) MeV in
the c.m. energy, while the experimental value is 0.94 MeV. In general, we find that NNLO\textsubscript{sat} predicts the location of dominant quasiparticle and holes states with a (conservative) accuracy of $<2$ MeV for this nucleus.

Finally, Fig. 4 compares the differential cross section for the elastic scattering of neutrons off $^{40}$Ca with the experiment at 13.56 MeV c.m. energy, with $N_{\text{max}} = 11$. Minima in the cross section are reproduced reasonably well, confirming the correct prediction of matter radii, but there appears to be a general lack of absorption. This may be due to either missing doorway configurations ($3p2h$ and beyond) or to the (still crude) model space.

Even with the limitations of a (non-optimal) oscillator basis, we found that most important features of optical potentials are well-reproduced. In the long term, it will be necessary to properly account for the continuum
in calculating the self-energy and to improve the realistic nuclear interactions. Nevertheless, it is clear from the present results that reliable \textit{ab initio} calculations of optical potentials are now a goal within reach.

A.I. is supported by the Royal Society and Newton Fund through the Newton International Fellowship scheme. This work was also supported by the United Kingdom Science and Technology Facilities Council (STFC) under grant No. ST/L005816/1 and by the NSERC grant No. SAPIN-2016-00033. TRIUMF receives federal funding via a contribution agreement with the National Research Council of Canada. Calculations were performed by using HPC resources from the DiRAC Complexity system at the University of Leicester (BIS National E-infrastructure capital grant No. ST/K000373/1 and STFC grant No. ST/K0003259/1).

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