Density-dependent nucleon-nucleon interaction from Urbana UIX three-nucleon force

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Abstract. A density-dependent two-nucleon potential has been derived within the formalism of correlated basis function. The effects of three-particle interactions has been included by integrating out the degrees of freedom of the third nucleon. The potential can be easily employed in nuclear matter calculations. It yields results in agreement with those obtained from the underlying three-body potential. Using the density dependent potential, we have carried out a study of the effects of three-nucleon interactions in symmetric nuclear matter within the Auxiliary Field Diffusion Monte Carlo (AFDMC) computational scheme.

1. Introduction

The Auxiliary Field Diffusion Monte Carlo (AFDMC) \cite{1} calculations of Ref. \cite{2} yield a binding energy of symmetric nuclear matter (SNM) lower than that predicted by both Fermi-Hyper-Netted-Chain (FHNC) and Brueckner-Hartree-Fock (BHF) calculations \cite{3}. This result suggested that the underbinding of SNM resulting from FHNC calculation with the UIX three body potential \cite{4} may not be avoided with a better choice of the variational wave function.

More refined three body potentials, while providing a quantitative account of the energies of the ground and low-lying excited states of nuclei with $A \leq 12$ \cite{5}, give very different results in pure neutron matter (PNM) \cite{6}.

Lagaris and Pandharipande \cite{7} and Friedman and Pandharipande \cite{8} proposed an effective, density-dependent, two-nucleon potential (TNI), suitable to take into account three- and many-nucleon forces. However, they adopted a purely phenomenological procedure, as the parameters of the density dependent functions appearing in this potential were determined through a fit of the saturation density, the binding energy per nucleon and the compressibility of symmetric nuclear matter (SNM). In general, fitting the potential to results obtained within many-body techniques makes the potential itself affected by the approximations implied in the many body calculations.

Our potential model \cite{9} improves on the TNI model, since it has been derived from a realistic microscopic three-nucleon force, the UIX potential, which provides a good description of the
properties of light nuclei. It has to be mentioned that, as recently shown [10], the UIX does not reproduce the nd scattering length and the vector polarization observables $A_y$ and $iT_{11}$. However, our approach can be applied to more refined three nucleon interactions, like the chiral NNLOL described in Ref. [10].

To obtain the density dependent potential, the average on the degrees of freedom of the third particle has been carried out using a formalism suitable to account for the full complexity of nuclear dynamics. Our results show that, in doing such reduction, of great importance is the proper inclusion of both dynamical and statistical NN correlations, whose effects on many nuclear observables have been found to be large [11, 12].

We have used Correlated Basis Function (CBF) theory and the Fantoni-Rosati (FR) cluster expansion formalism[13] to perform the calculation of the lowest order (linear-in-density) terms of the effective potential, arising from the irreducible three-nucleon interactions based on the UIX potential.

The effective potential has been implemented in the AFDMC computational scheme to obtain the equation of state (EoS) of SNM. A similar calculation using the UIX potential is not yet feasible, due to the complexities arising from the commutator term. In addition, the density-dependent potential can be used to include the effects of three-nucleon interactions in the calculation of the nucleon-nucleon scattering cross section in the nuclear medium. The knowledge of this quantity is required to obtain a number of nuclear matter properties of astrophysical interest, ranging from the transport coefficients to the neutrino emission rates [14, 15].

2. Two- and three- nucleon forces
The Argonne $v_8$ [16] two-body potential model is given by

$$\hat{v}_{ij} = \sum_{p=1}^{8} v^p(r_{ij}) O^p_{ij}$$

where

$$O^p_{ij} = (1, \sigma_{ij}, S_{ij}, \mathbf{L}_{ij} \cdot \mathbf{S}_{ij}) \otimes (1, \tau_{ij}) .$$

In the above equation, $\sigma_{ij} = \sigma_i \cdot \sigma_j$ and $\tau_{ij} = \tau_i \cdot \tau_j$, where $\sigma_i$ and $\tau_i$ are Pauli matrices acting on the spin or isospin of the $i$-th particle, while $S_{ij}$ is the tensor operator, $\mathbf{L}_{ij}$ is the relative angular momentum and $\mathbf{S}_{ij}$ is the total spin of the pair.

We used the so called Argonne $v'_8$ and Argonne $v'_6$ potentials, which are not simple truncations of the Argonne $v_{18}$ potential [17], but rather reprojections [18] obtained by refitting the scattering data and the deuteron binding energies. In all light nuclei and nuclear matter calculations the results obtained with the $v'_8$ are very close to those obtained with the full $v_{18}$.

Using a nuclear Hamiltonian including only two-nucleon interactions leads to the underbinding of light nuclei and overestimating the equilibrium density of nuclear matter. Hence, the contribution of three-nucleon interactions must be taken into account, by adding to the Hamiltonian the corresponding potential. One of the most widely used is Urbana IX (UIX) [4] potential, consisting of two terms: the Fujita and Miyazawa [19] attractive two-pion exchange interaction $V^{2\pi}$ and the purely phenomenological repulsive term $V^R$

$$\hat{V}^{2\pi} = A^{2\pi} \sum_{cyclic} \left( \{ \hat{X}_{ij}, \hat{X}_{jk} \} \{ \tau_{ij}, \tau_{jk} \} + \frac{1}{4} \{ \hat{X}_{ij}, \hat{X}_{jk} \} [\tau_{ij}, \tau_{jk}] \right)$$

$$V^R = U_0 \sum_{cyclic} T^2(m_\pi r_{ij}) T^2(m_\pi r_{jk}) .$$

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The spin structure of $V^{2\pi}$ is given by

$$\hat{X}_{ij} = Y(m_\pi r)\sigma_{ij} + T(m_\pi r)S_{ij}.$$  (4)

The radial functions read

$$Y(x) = \frac{e^{-x}}{x} \xi_Y(x), \quad T(x) = \left(1 + \frac{3}{x} + \frac{3}{x^2}\right)Y(x)\xi_T(x)$$  (5)

where $\xi_Y(x) = \xi_T(x) = 1 - \exp(-cx^2)$ are short-range cutoff functions. The parameters $A_{2\pi}$ and $U_0$ are varied to fit the observed binding energies of $^3$H and $^4$He and to reproduce the empirical nuclear matter saturation density, while the cutoff parameter is kept fixed at $c = 2.1$ fm$^{-2}$.

### 3. Correlated Basis Function theory and Cluster Expansion technique

In the correlated basis theories of Fermi liquids [20, 21], the expectation value of the two-body potential can be written in the form

$$\langle \hat{v} \rangle = \frac{1}{2} \rho \sum_p \int d\vec{r}_1 d\vec{r}_2 v_{12}^p g_{12}^p,$$  (6)

where

$$g_{12}^p = \frac{A(A-1)\text{Tr}_{12} \int dX \Phi_0^\dagger \Phi_0^\dagger \Phi_0^\dagger \Phi_0}{\rho^2 \int dX \Phi_0^\dagger \Phi_0},$$  (7)

are the operatorial components of the two-body distribution function. The uncorrelated wavefunction $\Phi_0$ in nuclear matter is conveniently chosen to be a Slater determinant of plane waves.

The structure of the correlation operator $\hat{F}$ reflects the complexity of the Argonne $v'_6$ nucleon-nucleon potential [22]:

$$F = S \prod_{j>i=1}^A F_{ij} \quad \text{with} \quad \hat{F}_{ij} = \sum_{p=1}^6 f^p(r_{ij})\tilde{O}_{ij}^p.$$  (8)

The radial functions $f^p(r_{ij})$, appearing in the definition of the correlation operator are determined by the minimization of the energy expectation value $E_V = \langle \Psi_0 | H | \Psi_0 \rangle$, which provides an upper bound to the true ground state energy $E_0$. As explained in detail in [9], the calculation of $E_V$ in CBF theories is carried out by i) expanding $\langle \Psi_0 | H | \Psi_0 \rangle$ in powers of dynamical correlations $h(r_{ij}) = f^0(r_{ij})^2 - 1$, $2f^0(r_{ij})f^p(r_{ij})$, $f^{p>1}(r_{ij})f^{q>1}(r_{ij})$ that vanish in uncorrelated matter and ii) summing up the main series of the resulting cluster terms by solving a set of coupled integral equations. To accomplish the first of these two steps, an extension of the FR cluster expansion [13], that is able to treat scalar correlations only, was introduced in [22] to deal with spin–isospin dependent correlation operators, like those of Eq. (8).

The cluster terms are most conveniently represented by diagrams [22] consisting of dots (vertices) connected by different kinds of correlation lines. In particular $h_{ij}$ is usually represented by a dashed line, $2f^0_{ij}f^0_{ij}$ by a single wavy line, and $f^{p>1}_{ij}f^{q>1}_{ij}$ by a doubly wavy line. In addition to the dynamical correlation lines, there are also statistical correlation lines, represented by solid oriented lines forming closed loops that never touch each other. They arise from the Slater determinant and are associated with the Slater function $\ell_{ij}$

$$\ell(k_{FR_{ij}}) = 3 \left[ \frac{\sin(k_{FR_{ij}}) - k_{FR_{ij}} \cos(k_{FR_{ij}})}{(k_{FR_{ij}})^3} \right].$$  (9)
Open dots of the diagrams represent the active (or interacting) particles (1 and 2), while black dots are associated with passive particles, i.e. those in the medium. Integration over the coordinates of the passive particles leads to the appearance of a factor $\rho$. It can be shown that all the diagrams contributing to $E_V$, hence to $\langle \hat{v} \rangle$, are linked. Those built with scalar passive bonds only, with the only exception of the so called elementary diagrams, can be summed up in closed form by solving the FHNC equations [13].

On the other hand, diagrams having one or more passive operatorial bonds are calculated at leading order only. This implies that at most two operatorial passive bonds can be attached to any internal point, thus only diagrams with Single Operator Chain (SOC) are considered. Such an approximation is justified by the observation that operatorial correlations are much weaker than the scalar ones.

4. Density dependent potential
As for the two-body potential $v_8$, it is possible to write the UIX three body potential in the following way

$$V_{123} \equiv \sum_p V^p_{123} O^p_{123}.$$  \hspace{1cm} (10)

The expectation value of $\hat{V}_{123}$ then reads

$$\frac{\langle V \rangle}{A} = \frac{1}{3!} \rho^2 \sum P \int dr_{12}dr_{13} V^p_{123} g^p_{123},$$  \hspace{1cm} (11)

with

$$g^p_{123} = \frac{A!}{(A-3)!} \frac{\int dx_4 \ldots dx_A \Phi^0_F O^p_{123} F \Phi^0}{\rho^3 \int dX \Phi^0_F \Phi^* F \Phi^0}.$$  \hspace{1cm} (12)

We require that the expectation values of $V_{123}$ and of $v_{12}(\rho)$ be the same, implying in turn

$$\sum_p \frac{P}{3} \int d\vec{r}_3 V^p_{123} g^p_{123} = \sum_p v^p_{12}(\rho) g^p_{12}.$$  \hspace{1cm} (13)

A diagrammatic representation of the above equation, which should be regarded as the definition of the $v_{12}(\rho)$, is shown in figure 1. The graph on the left-hand side represents the three-body potential times the three-body correlation function, integrated over the coordinates of particle 3. Correlation and exchange lines are schematically depicted with a line having a bubble in the middle, while the thick solid lines represent the three-body potential. The diagram in the right-hand side represents the density-dependent two-body potential, dressed with the two-body distribution function. Obviously, $v^p_{12}$ has to include not only the three-body potential, but also the effects of correlation and exchange lines.
In the construction of the density dependent interaction we have found that the most relevant diagrams are those depicted in figure 2 that indeed include both statistical and dynamical correlations. To simplify the pictures, the three-body potential acting on particles 1, 2 and 3 is not explicitly depicted. In order to include higher order cluster terms, we have replaced the scalar correlation line $f_{ij}^2$ with the Next to Leading Order (NLO) approximation to the bosonic two-body correlation function:

$$f_{ij}^2 \to g_{NLO}^{bose}(r_{ij}) = f_{ij}^2(1 + \rho \int d\vec{r}_3 h_{13} h_{23}).$$  (14)

The inclusion of both statistical and dynamical correlations plays a fundamental role in the determination of a realistic density dependent interaction, as can be seen from figure 3, where different density dependent potentials contributions are plotted and compared with the full calculation with the genuine three-nucleon interaction. In particular $v_{12}^{(I)}(\rho)$, obtained by averaging over the third particle without statistical and dynamical correlations is far from the solid curve of UIX. When only statistical correlations are considered, the relative potential, $v_{12}^{(II)}(\rho)$, comes closer to the UIX curve. Only when the full set of diagrams of figure 2 is included in the calculation

$$v_{12}^{(III)}(\rho) = \frac{\rho}{3} \int dx_3 V_{123} \left[ g_{NLO}^{bose}(r_{13}) g_{NLO}^{bose}(r_{23}) (1 - 2P_{13}f_{13}^2) + 4g_{NLO}^{bose}(r_{13}) f_{c}(r_{13}) \hat{f}(r_{23}) \right],$$  (15)

Figure 2. Diagrams contributing to the density-dependent potential. The dashed lines with diamonds represent the first order approximation to $g_{NLO}^{bose}(r_{ij})$, discussed in the text.

Figure 3. (Color online) Contributions of the density-dependent potential to the energy per particle of SNM (a) and PNM (b), compared to the expectation value of the three-body potential UIX: $\langle V_{123} \rangle / A$. 

Potential energy per nucleon (MeV)
where \( \hat{f}(r_{23}) \) denotes the sum of non central correlations, the contribution of the density dependent potential comes very close to that of the original UIX. A proper treatment of the statistical correlation is crucial for the construction of a density dependent potential to be treated as an additive term to the standard two-body potential. Some care is needed for the density dependent potential to reproduce the exchange loop involving particles 1, 2 and 3 with the appropriate symmetry factor (see [9] for a thorough discussion).

5. Numerical results: PNM and SNM equation of state

As explained in [9], the correlation functions are determined by minimizing the two-body cluster term of the energy expectation value for a set of parameters \( d_c, d_t, \beta_p \) and \( \alpha_p \). The energy expectation value \( E_V \) calculated in full FHNC/SOC approximation has then to be minimized with respect to the variation of these parameters. To this aim a Simulated annealing [23] optimization algorithm has been implemented [9].

Both SOC approximation and the fact that elementary diagrams are neglected may lead to a violation of the variational principle. To keep this effect under control, an additional constraint on the kinetic energy with respect to the optimization performed in [24] has been considered.

![Figure 4](image-url)

**Figure 4.** (Color online) Energy per particle for PNM, obtained using the density-dependent potential added to the Argonne \( v'_8 \) (a) and to Argonne \( v'_6 \) (b) potentials. The energies are compared to those obtained from the genuine three-body potential and from the two-body potentials alone.

We carried out AFDMC simulations [1] for PNM with \( A = 66 \) and SNM with \( A = 28 \) nucleons in a periodic box. The finite-size errors in PNM simulations have been investigated in [25] by comparing the Twist Averaged Boundary Conditions (TABC) with the Periodic Box Condition (PBC). The kinetic energy of 66 fermions approaches the thermodynamic limit very well, so that the energies of 66 neutrons computed using either TABC or PBC turn out to be almost the same. We can infer that the finite-size errors in the present AFDMC calculations for PNM do not exceed 2% of the asymptotic value of the energy. The finite-size effects of SNM calculations can be estimated from the difference of the energies of PNM obtained with 14 neutrons and the TABC asymptotic value, which is of the order of 7%.

FHNC/SOC and Monte Carlo calculations provide very close results, as shown in figures 4 and 5, to be compared with those of Ref. [2] where the agreement between FHNC and Monte Carlo methods were not nearly as good.

The EoS of PNM, displayed in figure 4, obtained with the three-body potential UIX and using the density-dependent two-body potential are very close to each other. In the same figure,
for the sake of comparison, we also report the results of calculations carried out including the two-body potential only.

Despite cluster contributions proportional to $\rho^2$ have been neglected in the density dependent potential, with the exception of the line with diamonds of figure 2, even at high densities the EoS relative to $v_{12}(\rho)$ remain close to those obtained with UIX. Probably, in this case a compensation among second and higher order terms takes place.

![Figure 5.](a) FHNC: $v'_6 + v_{12} (\rho)$ (b) FHNC: $v'_8 + v_{12} (\rho)$

The density-dependent potential has been also employed in AFDMC calculations. As can be seen in figure 4, the triangles representing the results of this calculation are very close, when not superimposed, to the circles corresponding to the UIX three-body potential AFDMC results.

For what concern the EoS of symmetric nuclear matter, see figure 5, at densities lower than $\rho = 0.32 \text{ fm}^{-3}$, the curves resulting from UIX and the density-dependent potential are very close to one other, while for $\rho > 0.32 \text{ fm}^{-3}$ a gap between them appears.

Table 1. Values for the saturation densities, the binding energy per particle, and the compressibility of SNM relative to the EoS of figure 5.

|             | FHNC/SOC $v'_6 + v_{12} (\rho)$ | $v'_8 + v_{12} (\rho)$ | $v'_6 + v_{123} (\rho)$ | $v'_8 + v(\rho)$ | AFDMC $v'_6 + v(\rho)$ | $v'_8 + v(\rho)$ |
|-------------|----------------------------------|------------------------|-------------------------|------------------|------------------------|------------------|
| $\rho_0$ (fm$^{-3}$) | 0.17                             | 0.16                   | 0.16                    | 0.15             | 0.17                   | 0.15             |
| $E_0$ (MeV)   | -11.3                            | -11.2                  | -10.3                   | -10.3            | -10.9                  | -10.3            |
| $K$ (MeV)     | 205                              | 192                    | 189                     | 198              | 201                    | 201              |

In table 1, the saturation density $\rho_0$, the binding energy per particle $E(\rho_0)$ and the compressibility $K = \partial E(\rho)/\partial \rho$ for all the EoS of figure 5 are listed. It is remarkable that the values obtained with density dependent potential are very close to those resulting from the genuine UIX three body potential. While the saturation density is well reproduced, this being not surprising since the constant $U_0$ of the UIX potential has been fitted to this value, the values of the compressibility are slightly lower than the experimental ones. AFDMC calculations lower the variational results for $E_0$ by 0.6 MeV only, showing that the binding energy overestimation of the FHNC/SOC calculations performed with UIX potential can not be solved with a better choice of the variational wavefunction.
6. Conclusions
We have developed a novel scheme, suitable to obtain an effective density-dependent NN potential taking into account the effects of three-nucleon interactions. Our approach is fully consistent with the treatment of correlations underlying the FHNC and AFDMC approaches.

The PNM and SNM equation of state resulting from the density-dependent potential turn out to be very close to those obtained with the UIX three-body potential. In this context, a critical role is played by the treatment of both dynamical and statistical correlations. This is a distinctive feature of our approach, as compared to different reduction schemes based on effective interactions [26, 27].

A AFDMC calculation of the equation of state of SNM consistently including the effects of three nucleon forces has been carried out for the first time. The results of this calculation show that the $v'_6+\text{UIX}$ hamiltonian, or equivalently the one including the effective potential, fails to reproduce the empirical data. The discrepancy has most likely to be ascribed either to deficiencies of the UIX model or to the effect of interactions involving more than three nucleons.

As a further development, we are studying the dependence on the specific model of three-nucleon force [28], as well as the inclusion of four- and many-nucleon interactions, whose effects are expected to be critical for the determination of the properties of high density neutron star matter.

As a final remark, we note that our density dependent potential could be easily employed in many-body approaches other than those based on the CBF formalism or quantum Monte Carlo simulations, such as the G-matrix and self-consistent Green function theories [29, 30, 31].

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