Quantum Monte Carlo study of inhomogeneous neutron matter

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Abstract. We present an ab-initio study of neutron drops. We use Quantum Monte Carlo techniques to calculate the energy up to 54 neutrons in different external potentials, and we compare the results with Skyrme forces. We also calculate the rms radii and radial densities, and we find that a re-adjustment of the gradient term in Skyrme is needed in order to reproduce the properties of these systems given by the ab-initio calculation. By using the ab-initio results for neutron drops for close- and open-shell configurations, we suggest how to improve Skyrme forces when dealing with systems with large isospin-asymmetries like neutron-rich nuclei.

1. Introduction
Even though neutron drops, neutrons confined by an external field, cannot be studied experimentally, they have nevertheless generated intense theoretical interests for a couple of reasons. Neutron drops provide a very simple model of neutron-rich nuclei, in which the core is modeled as an external potential acting on valence neutrons. For example in Refs. [1, 2, 3] neutron-rich oxygen isotopes have been modeled by neutron drops, and in Ref. [4] the same model has been used to study calcium isotopes. Second, they provide a description of inhomogeneous neutron matter that can be used as data for calibrating model energy density functionals in several conditions [5, 6, 7].

On the experimental side, new facilities plan to study the physics of neutron-rich nuclei close to the drip line. The study of these systems is typically done by using methods based on energy density functionals and effective forces. These forces, like Skyrme or Gogny, are fitted to reproduce properties of nuclear and neutron matter combined with experimental energies of nuclei close to stability. By constraining the density functionals to the equation of state of nuclear and neutron matter, the symmetry energy is implicitly included in these forces. However, the terms dependent on derivatives of proton and neutron densities are constrained only to nuclei close to stability, and then for very small isospin-asymmetries.

The use of these functionals to study nuclei close to the neutron drip line requires then an important extrapolation to large isospin-asymmetries. This extrapolation is even more drastic when the Skyrme forces are used to study the properties of the neutron star crust, where the matter is made by extremely neutron-rich nuclei surrounded by a sea of neutrons. For these reasons, ab-initio calculations of these systems starting from accurate nuclear Hamiltonians are important to constrain density functionals.

Important advances have been made in building very accurate Hamiltonians describing nuclear systems. The modern nucleon-nucleon forces fit scattering data with high accuracy [8, 9],
and also the three-body forces are quite well constrained to reproduce properties of light nuclei [10, 11].

Furthermore, in recent years new techniques have been developed to solve for the ground-state of nuclear systems. In this paper we present results obtained with Quantum Monte Carlo techniques, i.e. the well known Green’s Function Monte Carlo (GFMC) and Auxiliary Field Diffusion Monte Carlo (AFDMC). The two different algorithms use different variational wave functions, but they show similar accuracy in calculating the energy of pure neutron systems.

2. Nuclear Hamiltonian and Quantum Monte Carlo method

In our model, neutrons are non-relativistic point-like particles interacting via two- and three-body forces, that are confined by an external potential:

\[
H = \sum_{i=1}^{A} \frac{p_i^2}{2m} + \sum_i V_{\text{ext}}(r_i) + \sum_{i<j} v_{ij} + \sum_{i<j<k} v_{ijk},
\]

(1)

where the external potential is a Harmonic Oscillator (HO)

\[
V_{HO}(r) = \frac{1}{2} m \omega^2 r^2,
\]

(2)

or a Woods-Saxon

\[
V_{WS}(r) = -V_0 \frac{1}{1 + \exp[(r - r_0)/a]}.
\]

(3)

In our study the parameters are \(V_0 = -35.5\) MeV, \(r_0 = 3\) fm and \(a = 1.1\) fm, and \(\hbar^2/m = 41.44\) MeV-fm\(^2\). The parametrization of the WS has been chosen to reproduce the properties of oxygen isotopes [1]; for the HO we consider different values of \(\hbar \omega\) to change the density of the system.

The two body-potential that we use is the Argonne AV8’ [12], a simplified form of the Argonne AV18 [8]. Although simpler to use in QMC calculations, the AV8’ provides almost the same accuracy as AV18 in fitting NN scattering data.

In this work we use the UIX three-body force, that has been originally proposed in combination with the Argonne AV18 and AV8’ [13]. Although it slightly underbinds the energy of light nuclei, it has been extensively used to study the equation of state of nuclear and neutron matter [14, 15], and gives an equation of state of pure neutron matter stiff enough to support a neutron star with mass higher than 2 solar masses [14, 15, 16, 17], as suggested by recent observations [18, 19]. The Illinois three-body forces have been introduced to improve the description of both ground- and excited-states of light nuclei with \(A \leq 8\) [10], and produce energy spectra for \(A\) up to 12 in excellent agreement with data [11], but produce an unphysical overbinding in pure neutron systems [20]. In neutron drops their contribution is also attractive [21].

We solve the many-body ground-state using both the Green’s Function Monte Carlo (GFMC) and Auxiliary Field Diffusion Monte Carlo (AFDMC) [22]. The main idea of QMC methods is to evolve a many-body wave function in imaginary-time:

\[
\Psi(\tau) = \exp[-H\tau]\Psi_v,
\]

(4)

where \(\Psi_v\) is a variational ansatz, and \(H\) is the Hamiltonian of the system. In the limit of \(\tau \to \infty\), \(\Psi\) approaches the ground-state of \(H\). The evolution in imaginary-time is performed by sampling configurations of the system using Monte Carlo techniques, and expectation values are evaluated over the sampled configurations. For more details see for example Refs. [15, 23].

The GFMC has proven to be extremely accurate to study properties of light nuclei. The variational wave function includes all the possible spin/isospin states of the nucleons and provides
a good variational ansatz to start the projection in imaginary time. The exponential growth of
the number of spin-isospin states with particle number limits the calculation to $^{12}$C [2] nuclei or
drops with 16 neutrons. The AFDMC method does not include all the spin/isospin states in the
wave function. The latter are instead sampled using the Hubbard-Stratonovich transformation.
Using the AFDMC the calculation can be extended up to many neutrons, making the simulation
of large neutron drops with up to several tens of neutrons possible. In the case of homogeneous
systems the AFDMC has been used to simulate up to 114 neutrons [15], and up to 70 neutrons
in an external well.

3. Results
The energy of neutron drops confined by $V_{HO}$ is shown in Fig. 1 for two different frequencies of
the external potential. The red points are the results obtained using the AFDMC method, and
the blue ones using the GFMC. The two solid lines are the results given by using the original
Skyrme SLY4 force [24], and a modified version. The energy is in units of the Thomas-Fermi
energy, that is proportional to $\omega N^{4/3}$, to see the extrapolation to the thermodynamic limit. The
two QMC methods agree within 1% for the $\hbar\omega = 10$ MeV trap, and the difference increases
up to 4% for $\hbar\omega = 5$ MeV. The reason is that pairing correlations are included in the GFMC
wave function, but not in the AFDMC. At low densities neutrons are superfluids, and pairing
correlations are quite important to include for open-shell configurations.

The difference between QMC and Skyrme at closed shells is mainly due to two effects, the
bulk contribution and the gradient term. Skyrme forces typically give an EOS of pure neutron
matter at densities lower than saturation that is more repulsive than microscopic calculations.
The equation of state of pure neutron matter is shown in Fig. 2 where we compare the AFDMC
results from Refs. [15, 25, 29], the GFMC calculation of Gezerlis and Carlson [26, 27], and the
equation of state given by SLY4. We make the reasonable assumption that Skyrme’s bulk term
cannot explain the difference between QMC and Skyrme energy in neutron drops. Then, since

![Figure 1. The energy of neutrons in a HO well with $\hbar\omega = 10$ MeV (upper panel) and 5 MeV
(lower panel) in units of $\omega N^{4/3}$. The red dots are the results given by AFDMC, blue squares
are from GFMC, and the black line is the result obtained using the Skyrme SLY4. The violet
line is the adjusted SLY4 where the strength of the gradient, pairing, and spin-orbit terms have
been changed. The figure is taken from Ref. [5].](image-url)
Figure 2. The equation of state of pure neutron matter. The AFDMC results [15, 25] are compared to the QMC results at low densities of Gezerlis and Carlson [26, 27], and to the equation of state of Skyrme SLY4 [28].

the pairing and the spin-orbit terms are expected to be very weak with respect to the gradient term for closed shell configurations, we can use the energy at N=8, 20 and 40 to re-adjust the gradient term of Skyrme. The energy of neutron drops with N near closed shells can be used to adjust the spin-orbit strength because for these configurations the pairing is not important. Finally, by comparing the energy of half-filled shells, we can tune the pairing term.

The ab-initio results can also be used as a benchmark for new Skyrme forces. For example, in a recent paper Kortelainen et al. have derived new Skyrme forces, and they have produced neutron drops results as a prediction to be compared with QMC calculations. The comparison is shown in Fig. 3. The agreement between the two new Skyrme forces with QMC results is remarkable considering that the ab-initio energies of neutron drops have not been included as a constraint in producing the UNEDF0 and UNEDF1 forces.

The use of a Woods-Saxon external potential $V_{WS}$ is also interesting because it reflects properties of nuclei. It saturates for some number of neutrons, providing a good test of the surface effects of Skyrme. The comparison between the original and the adjusted Skyrme SLY4 with QMC for neutrons in a WS potential is shown in Fig. 4. We stress that the SLY4 has been adjusted only to reproduce the results in the HO well, and is used here to calculate the neutron drop energies as a prediction. Also in this case the agreement with the QMC calculations is quite good.

Finally, we studied the effect of the adjusted Skyrme to the structure of neutron drops by calculating the rms radii and density. The rms radius $\sqrt{\langle r^2 \rangle}$ is shown in Fig. 5 where we compare the GFMC estimate with the original and adjusted SLY4 for different numbers of neutrons in the WS well (lower panel), and similarly with HO well for two different frequencies (upper panel). In Fig. 6 we show the radial density calculated with GFMC and Skyrme in the case of HO well for a closed shell configuration with N=8 (upper panel), and for an open-shell with N=14 (lower panel). Both for rms radii and for the density the adjusted Skyrme agree much better with QMC calculations, showing that the energy constraint is sufficient to improve
**Figure 3.** Energy of neutrons in a HO well. The AFDMC results are compared with those given by UNEDF0 and UNEDF1 calculations. Figure taken from Ref. [30] with courtesy of W. Nazarewicz.

**Figure 4.** The energy per particle of neutrons in a WS well obtained using AFDMC (blue squares) and GFMC (red points) compared to Skyrme. Figure taken from Ref. [5].
the accuracy of density functionals, whose predictive power for other operators would drastically improve.

4. Conclusions
We performed ab-initio calculations of neutrons confined in external potentials. These systems are interesting to study because they provide a simple model to describe neutron-rich nuclei, and can be used to constrain energy density functionals. By using QMC techniques, we calculated the energy of neutrons in different potentials. We have considered both harmonic oscillator and Woods-Saxon wells in order to provide results for systems with rather different geometries. We paid particular attention to the comparison between the GFMC and AFDMC methods. The agreement is quite good, at the few percent level for the total energy of the system.

We compared the energies of neutron drops given by QMC methods with those given by different Skyrme forces, and found that the latter always produce lower energies. Since the density of these systems is lower than nuclear densities, we suggest that the overbinding is mainly due to the (neutron) gradient term. We found that, in order to reproduce our ab-initio results, the pairing and the spin-orbit terms of Skyrme need to be changed too.

We have used AFDMC to calculate the energy of drops up to 54 neutrons, and GFMC to calculate the rms radii and radial densities of drops up to 16 neutrons. After re-adjustment of the SLY4 parametrization, the Skyrme force provides much better agreement for these quantities.

Acknowledgments
The author would like to thank J. Carlson and R. Schiavilla for critical comments on the manuscript, and W. Nazarewicz for the useful discussions and for the permission to show Fig. 3. This work is supported by DOE Grants No. DE-FC02-07ER41457 (UNEDF SciDAC) and No. DE-AC52-06NA25396, and by the LANL LDRD program. Computer time was made available by
Figure 6. The radial density of neutron drops for different configurations calculated using GFMC (squares) compared with the original (dashed lines) and the adjusted SLY4 (solid lines) in a HO well. In the top panel we show the results of a closed shell configuration with N=8, and in the bottom panel the density of an open shell with N=14. The two colors are for different HO frequencies. Figure taken from Ref. [5].

Los Alamos Open Supercomputing, and by the National Energy Research Scientific Computing Center (NERSC).

References
[1] Chang S Y, Morales J, Jr, Pandharipande V R, Ravenhall D G, Carlson J, Pieper S C, Wiringa R B and Schmidt K E 2004 Nucl. Phys. A 746 215
[2] Pieper S C 2005 Nuclear Physics A 751 516
[3] Gandolfi S, Pederiva F, Fantoni S and Schmidt K E 2006 Phys. Rev. C 73 044304
[4] Gandolfi S, Pederiva F and A Beccara S 2008 European Physical Journal A 35 207–211
[5] Gandolfi S, Carlson J and Pieper S C 2011 Phys. Rev. Lett. 106 012501
[6] Bogner S, Furnstahl R, Hergert H, Kortelainen M, Maris P et al. 2011 Phys. Rev. C 84 044306
[7] Drut J E and Platter L 2011 Phys. Rev. C 84(1) 014318
[8] Wiringa R B, Stoks V G J and Schiavilla R 1995 Phys. Rev. C 51 38
[9] Entem D R and Machleidt R 2003 Phys. Rev. C 68 041001
[10] Pieper S C, Pandharipande V R, Wiringa R B and Carlson J 2001 Phys. Rev. C 64 014001
[11] Pieper S C 2008 AIP Conf. Proc. 1011 143
[12] Wiringa R B and Pieper S C 2002 Phys. Rev. Lett. 89 182501
[13] Pudliner B S, Pandharipande V R, Carlson J and Wiringa R B 1995 Phys. Rev. Lett. 74 4396
[14] Akmal A, Pandharipande V R and Ravenhall D G 1998 Phys. Rev. C 58 1804
[15] Gandolfi S, Illarionov A Y, Schmidt K E, Pederiva F and Fantoni S 2009 Phys. Rev. C 79 054005
[16] Gandolfi S, Illarionov A Y, Fantoni S, Miller J, Pederiva F and Schmidt K 2010 Mon. Not. R. Astron. Soc. 404 L35
[17] Gandolfi S, Carlson J and Reddy S 2012 Phys. Rev. C 85 032801
[18] Demorest P B, Pennucci T, Ransom S M, Roberts M S E and Hessels J W T 2010 Nature 467 1081
[19] Steiner A W and Gandolfi S 2012 Phys. Rev. Lett. 108 081102
[20] Sarsa A, Fantoni S, Schmidt K E and Pederiva F 2003 Phys. Rev. C 68 024308
[21] Carlson J, Gandolfi S, Maris P, Pieper S C and Vary J P In preparation
[22] Schmidt K E and Fantoni S 1999 Phys. Lett. B 446 99
[23] Pudliner B S, Pandharipande V R, Carlson J, Pieper S C and Wiringa R B 1997 Phys. Rev. C 56 1720
[24] Chabanat E, Bonche P, Haensel P, Meyer J and Schaeffer R 1995 Phys. Scr. T56 231
[25] Gandolfi S, Illarionov A Y, Fantoni S, Pederiva F and Schmidt K E 2008 Phys. Rev. Lett. 101(13) 132501
[26] Gezerlis A and Carlson J 2008 Phys. Rev. C 77 032801
[27] Gezerlis A and Carlson J 2010 Phys. Rev. C 81(2) 025803
[28] Chabanat E, Bonche P, Haensel P, Meyer J and Schaeffer R 1998 Nuclear Physics A 635 231–256
[29] Gandolfi S, Illarionov A Y, Pederiva F, Schmidt K E and Fantoni S 2009 Phys. Rev. C 80(4) 045802
[30] Kortelainen M, McDonnell J, Nazarewicz W, Reinhard P G, Sarich J, Schunck N, Stoitsov M V and Wild S M 2012 Phys. Rev. C 85(2) 024304