Static Response of Neutron Matter

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We generalize the problem of strongly interacting neutron matter by adding a periodic external modulation. This allows us to study from first principles a neutron system that is extended and inhomogeneous, with connections to the physics of both neutron-star crusts and neutron-rich nuclei. We carry out fully non-perturbative microscopic Quantum Monte Carlo calculations of the energy of neutron matter at different densities, as well as different strengths and periodicities of the external potential. In order to remove systematic errors, we examine finite-size effects and the impact of the wave function ansatz. We also make contact with energy-density functional theories of nuclei and disentangle isovector gradient contributions from bulk properties. Finally, we calculate the static density-density linear response function of neutron matter and compare it with the response of other physical systems.

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Neutron matter (NM) is directly related to the properties of neutron-star crusts and cores [1]. The neutron-matter equation of state (EOS) has been intensively studied over the last few decades within *ab initio* approaches [2–7], due to its astrophysical relevance and its connections with the physics of neutron-rich nuclei. Neutron matter, though strongly interacting, is not self-bound, making it more straightforward to tackle than isospin asymmetric matter or nuclei. This has led to the use of pure neutron systems as a testbed of nuclear forces, whether phenomenological [8–12] or chiral [13–21].

While the interface between NM calculations and neutron-star properties is relatively direct (following from Einstein’s field equations), the connection with neutron-rich nuclei is more circuitous. It usually proceeds via nuclear energy-density functionals (EDFs), the only approach currently able to provide a global description of the nuclear chart [22]. EDFs contain a number of undetermined parameters, typically fit to nuclear masses and radii. EDF approaches have benefitted from including as constraints “synthetic data”, i.e., dependable *ab initio* many-body results. Most commonly, these refer to the EOS of neutron matter [23–29], though other quantities have also been used, like the neutron pairing gap [30], the energy of a neutron impurity [31, 32], or the properties of artificially confined neutron drops [33–36].

In this Letter, we introduce another constraint: the static response of neutron matter, namely the behavior of an extended neutron system in the presence of a periodic external potential. Like neutron drops or the neutron polaron, the static-response problem provides input into the physics of neutron-rich nuclei. Furthermore, the static response of neutron matter is directly equivalent to the situation prevailing in a neutron-star crust. There, unbound neutrons interact strongly with each other and with a lattice of nuclei: the unbound neutrons therefore experience the nuclei as a periodic modulation. The inhomogeneity of nucleon matter in nature has been addressed in several works, employing a variety of approximations [37–42]: this Letter opens up a corresponding *ab initio* avenue.

The physical setting can be thought of as a periodic system of neutron drops. Our investigation aims to provide insights on the relative magnitude of bulk vs surface effects. In other words, the static response of neutron matter can guide us in the creation of “neutron pasta” [43], i.e., non-spherical nuclear systems, as well as their interplay with surrounding matter. Carrying out analogous calculations at very small densities one can then envision similarly disentangling pairing effects from the bulk and surface terms.

The problem of static response for strongly correlated systems has a long history as well as direct experimental relevance, motivating us to compare and contrast nuclear phenomena with systems in other areas of physics. Specifically, there are close analogies with both liquid and plasma systems has a long history as well as direct experimental relevance, motivating us to compare and contrast nuclear phenomena with systems in other areas of physics. Specifically, there are close analogies with both liquid and plasma.

Turning to the specifics of our investigation, we start from a microscopic Hamiltonian containing a non-relativistic kinetic energy, a two-nucleon (NN) interaction (here taken to be the standard high-quality Argonne v8’ potential [49]) and a three-nucleon (NNN) interaction (specifically, the Urbana IX potential [50]). We will not focus on the details of the NN and NNN interactions in what follows. The situation can be viewed as the application of a static (i.e., not dynamic) external potential $V_{\text{ext}}$ to an unperturbed Hamiltonian. Altogether, we have:

\[
\hat{H} = -\frac{\hbar^2}{2m} \sum_i \nabla_i^2 + \sum_{i<j} V_{ij} + \sum_{i<j<k} V_{ijk} + V_{\text{ext}}
\]  

The last term is the external periodic potential: $V_{\text{ext}} = \ldots$
\[ \sum_i V_i, \text{ where } V_i = 2v_\beta \cos(q \cdot r_i). \] The periodicity of the potential is contained in \( q \). We impose periodic boundary conditions on the wave function. If the density is \( n \) and the simulation is carried out using \( N \) particles, then the box length is \( L = (N/n)^{1/3} \). In order to respect translational invariance, we ensure that an integer multiple of the potential’s period fits in the box.

We employ a family of \textit{ab initio} non-perturbative many-body approaches that have been very successful in nuclear physics [11, 15, 50], namely Quantum Monte Carlo (QMC). Specifically, we use Variational Monte Carlo (VMC), which evaluates the expectation value of the Hamiltonian using a trial/variational wave function \( |\Psi_T\rangle \), as well as Auxiliary-Field Diffusion Monte Carlo (AFDMC), which extracts the ground state by projecting out excited states in imaginary time \( \tau \): \[ |\Psi_0\rangle = \exp(-\hat{H}\tau)|\Psi_T\rangle. \] Both methods have been applied to modern nuclear interactions, which include central, spin, tensor, spin-orbit, and more complicated terms.

We used VMC and AFDMC to calculate the equation of state of neutron matter both excluding and including a modulating \( V_{\text{ext}} \) potential. The results for densities 0.02 to 0.12 fm\(^{-3} \) (roughly corresponding to the crust and the outer core) are shown in Fig. 1. Black squares show the result of using only an NN interaction for 66 particles; they match known values [51]. (Here and below AFDMC results are shown – the VMC results behave analogously [52]). The trial wave function has the form:

\[
|\Psi_T\rangle = \prod_{i<j} f(r_{ij}) \mathcal{A} \prod_i |\phi_i, s_i\rangle
\]

This is a product of a Jastrow factor (with central correlations, so in principle irrelevant) and an antisymmetrized term (a Slater determinant of single-particle orbitals). As is standard for unpolarized neutron matter, these results follow from plane-wave orbitals \( \phi_i \).

Turning on the sinusoidal potential \( V_{\text{ext}} \), one can expect that the energy will be lowered: the particles will tend to stay away from repulsive regions and form clusters in the wells of the potential. We have carried out simulations that bear out this expectation, using the same plane-wave orbitals as above, a periodicity \( q = 4\pi/L \), and a strength that is a constant fraction of the Fermi energy \( E_F \) (2\( v_q = 0.5E_F \), green diamonds in Fig. 1). Note that our one-body strength is density dependent. Also, as the density increases, for fixed \( N \) the box size \( L \) decreases; since the box length is a fixed multiple of the period, the latter decreases. This is similar to what goes on in a neutron-star crust, where as the density increases the lattice spacing decreases.

We then proceeded to optimize the trial wave function \( |\Psi_T\rangle \) in VMC: instead of using plane-wave orbitals (which are the solution to the one-body problem in the absence of an external potential), we employ Mathieu functions (the solutions to the one-body problem in the presence of a sinusoidal potential). We also carried out a second optimization procedure: instead of assuming that the Mathieu functions are eigenstates for noninteracting particles in the external potential with strength \( 2v_q \), we used a new one-body strength \( \beta \) (only in the evaluation of the wave function) as a variational parameter [44]. The final AFDMC result (using the \( \beta \) that gives the minimum VMC energy) is shown using blue circles in Fig. 1. The difference between the plane-wave and Mathieu results (up to \( \sim 1 \) MeV) underscores the importance of optimizing the wave function properly. In comparison, the effect of the \( \beta \) parameter is small, typically \( \sim 50 \) keV.

The periodically modulated EOS results in Fig. 1 were calculated using only a single one-body potential strength, \( 2v_q = 0.5E_F \). While that was useful in understanding the importance of the wave function ansatz, it may be misleading: there is, after all, nothing special about any one specific strength value. Thus, it will be enlightening to study the dependence of the EOS on the potential strength, elucidating the departure from the case of homogeneous matter. Figure 2 shows the result of precisely such an investigation using both NN and NNN interactions, with \( 2v_q \) values chosen as follows: in order to ensure that the modulated ground-state energy in VMC is statistically distinct from the homogeneous-matter energy, \( 2v_q \) needs to be not too small. At the other end, above a certain \( 2v_q \) the single-particle orbital
The class of Skyrme EDFs can be given in the isospin presentation: 

\[ E = \sum_{T=0,1} \left[ (C_T^{n,a} + C_T^{n,b} n_0^a) n_T^2 + C_T^{\Delta n} n_T \Delta n_T + C_T^{n,n} n_T n_T \right] \] (3)

along with an external potential-energy term (possibly also dependent on the isospin projection) \( \sum_T n_T V_{ext,T} \), as well as a kinetic-energy term, which for simplicity can be taken to be an isoscalar: \( h^2 \sigma_0 / 2m \). The SLy4 C values used to produce Fig. 2 are one possible (standard) choice, though several others are being explored 52.

At first sight, Fig. 2 appears to be rather uninteresting: the AFDMC results are always more repulsive than the SLy4 ones, as found in QMC studies of neutron drops 23. It is noteworthy, however, that the separation between the AFDMC and SLy4 results depends on the one-body strength. Specifically, the isovector gradient term, \( C_1^{\Delta n} n_1 \Delta n_1 \), which is absent in homogeneous neutron matter, has the effect of bringing the SLy4 results closer to the AFDMC ones as \( 2\nu_q \) is increased.

Upon closer inspection, we realize that the more repulsive nature of the AFDMC results merely tracks the unperturbed (\( \nu_q = 0 \)) relationship between AFDMC and SLy4 results (though at very large \( 2\nu_q \) the gradient term becomes strong enough to change this). This crucial observation implies that, at different densities, for many values of the one-body strength the modulated EOS under SLy4 will have the same relationship to the AFDMC EOS as in the unmodulated (\( \nu_q = 0 \)) case. This is illustrated in the inset to Fig. 2, where the homogeneous neutron matter energy versus density in AFDMC and SLy4 is plotted. The AFDMC results are more repulsive at densities above 0.06 \( \text{fm}^{-3} \), while below that density the SLy4 ones are more repulsive. We have explicitly checked that the modulated interacting neutron system behaves as described at low density, for a variety of nuclear-force inputs.

Thus, the assumption that, since Skyrme functionals are fit to ab initio results for homogeneous matter, the difference between QMC and EDF in inhomogeneous neutron systems must come from the isovector gradient term alone 35 is unwarranted. In other words, at a finite one-body potential strength the difference between QMC and EDF comes partly from the gradient term and partly from the bulk/homogeneous matter mismatch in energy. Thus, in order to separate the two contributions while also respecting the homogeneous-matter properties, one should modify the isovector gradient parameter \( C_1^{\Delta n} \) so that the new Skyrme values match not the AFDMC ones, but a new set (here more attractive than the original) that is always as far away from the AFDMC perturbed value as the difference between the unperturbed AFDMC and SLy4 energies. The result is the dashed line in Fig. 2, which required a modification to \( C_1^{\Delta n} \) from -16 to -28 MeV \( \text{fm}^5 \), using only the lowest-strength results as constraints in the new fit.

The density dependence shown in the inset to Fig. 2, taken together with our low-density findings mentioned above, suggest a corresponding need for a density dependence in the adjustment to \( C_1^{\Delta n} \). In other words, if one is to carefully disentangle bulk from gradient contributions, the isovector gradient coefficient \( C_1^{\Delta n} \) has to become density dependent (cf. the bulk term(s) \( C^n \) in Eq. (3)), in addition to necessitating a global refit of all the Skyrme parameters. (This is consistent with what is found using the density-matrix expansion 53, 54.) Another (possibly equivalent) option is to consider gradient terms of different form 47.

The calculations reported on in Figs. 1 & 2 varied the density \( n \) and one-body potential strength \( 2\nu_q \), respectively, but kept the one-body potential periodicity \( q \) fixed. We have also carried out calculations where the \( q \) is varied and thereby computed the linear density-density static response function of neutron matter.

We first establish the notation 52, 55. Let the unperturbed system be characterized by a Hamiltonian \( \hat{H}_0 \).
(in our case, the first three terms in Eq. (1)). The last term in Eq. (1) can then be viewed as the coupling of an external potential $V(r)$ to the one-body density operator $\hat{n} = \sum_i \delta(r - r_i)$: $\int d^3r \, \hat{n}(r)V(r)$, where $V(r_i) \equiv V_i$. We can now define the linear density-density static response function from the functional expansion of the modulated density $n_{\text{tot}}(r)$ in terms of $V(r)$:

$$n_{\text{tot}}(r) = n_0 + \int d^3r' \chi(r' - r)V(r') \tag{4}$$

Explicitly, this $\chi(r' - r)$ is the density-density response function of order 1, so a response function of order 3 expansion can be written down for the modulated energy function of order 1, so a response function of order 3

$$\chi V \bigg|_{r = r_i}$$

of the modulated potential, $V(r) = \sum q v_q \exp(2i \mathbf{q} \cdot \mathbf{r})$, and the Fourier transform of the response function, $\chi(q)$:

$$\frac{E_{\text{tot}}}{N} = \frac{E_0}{N} + \frac{\chi(q)}{n_0} v_q^2 + C_4 v_q^4 + \cdots \tag{5}$$

The higher-order coefficients come from the higher-order response functions. Note that there are only even powers present. It’s also worth mentioning that the response function $\chi(q)$ is related to an energy integral over the dynamic structure factor.

In producing the static-response function results we first addressed finite-size corrections. The situation is illustrated in the inset to Fig. 3, removing NN and NNN interactions, one can compute quasi-analytically

![Figure 3](image_url)

**FIG. 3.** Static-response function of neutron matter at a density of $n = 0.10 \text{fm}^{-3}$. Points follow from AFDMC results using NN+NNN interactions, as well as several one-body strengths and periodicities. The line is the Lindhard function describing the response of a non-interacting Fermi gas. Inset: finite-size dependence of a non-interacting Fermi gas in the presence of a one-body potential of fixed strength $2v_q = 0.5E_F$ and periodicity $q = 4\pi/L$.

the energy of a modulated non-interacting Fermi gas, $E_{\text{NI}}$. This is shown in units of the energy per particle in the unmodulated non-interacting Fermi gas, $E_{\text{FG}}/N = (3/5)E_F$. The answer for 66 particles shows considerably larger shell effects than are present in the free Fermi gas. The thermodynamic-limit value depends on the specific strength and periodicity of the potential being examined. We have carried out similar calculations with up to $\sim 100,000$ particles, arriving at a systematic way of connecting finite-$N$ results to the thermodynamic limit. As a check, we adjusted the SLy4 results of Fig. 2 and were able to reproduce the thermodynamic-limit SLy4 values to less than 0.5%, thereby increasing our confidence in the prescription’s validity. (Since the periodic potential is externally specified, we do not expect major interaction-dependent finite-size effects, barring the extreme case of many clusters.)

For a fixed value of $q$ we carried out calculations for various one-body strengths $(2v_q = 0.25, 0.30, 0.35, 0.50, 0.75 \times E_F$, as above) and extracted the $\chi(q)$ using Eq. (5) by fitting in powers of $v_q$ to $v_q^4$. We repeated this process for several values of the periodicity: $q = 2, 4, 6, 8, 12, 16, 20 \times \pi/L$. This corresponds to $1, 2, 3, 4, 6, 8, 10$ periods of the potential, respectively; note that even at the lowest density, this is still 2-3 times smaller than the relevant neutron-star crust scale. The result of using NN+NNN interactions in AFDMC is shown in Fig. 3 (Keeping different powers in Eq. (5) is related to how one treats the higher-strength results, but the qualitative features are robust). We reiterate that each point in this figure corresponds to an entire plot like Fig. 2. Thus, since each simulation uses a minimum of 66 strongly interacting particles, producing dependable *ab initio* results for the static-response function of neutron matter requires a not insignificant amount of computing resources. Due to the varied connections with neutron-rich matter and nuclei, this effort is worthwhile.

Seeking a physical interpretation of our microscopic results, we compared them to the Lindhard function, corresponding to a non-interacting Fermi gas [54]:

$$\chi_L = \frac{-m q_F}{2\pi^2 \hbar^2} \left[1 + q_F q \left(1 - \frac{q}{2q_F} \right)^2 \ln \frac{q + 2q_F}{q - 2q_F} \right] \tag{6}$$

where $q_F$ is the Fermi wave number. (We also used calculations like those in the inset to Fig. 3 to reproduce this expression, further confirming our prescription.) We find that at large $q/q_F$ the interacting $\chi(q)$ matches the Lindhard function: since $C_4$ is small, Eq. (5) shows that when several clusters are present the interactions have a negligible effect. In the low-$q$ region our results exhibit a suppression: the particles are constrained closer to each other, thereby accenting the repulsion. This suppression is different than in the electron gas, where at vanishing $q$ the response goes rapidly to zero [45]. We used the
random-phase approximation to calculate the low-\(q\) behavior for a model short-range interaction, finding a value between the Lindhard and Coulomb results. While this result is merely qualitative, it nicely parallels our \textit{ab initio} prediction for the response of interacting neutron matter.

In summary, we have proposed the study of periodically modulated strongly interacting neutron matter. We have tackled this problem using Variational Monte Carlo and Auxiliary-Field Diffusion Monte Carlo. We investigated the ground-state energy at different densities, paying close attention to the effect of optimizing the wave function. We then addressed the question of how the homogeneous problem turns into the inhomogeneous one, by carrying out calculations for several values of the one-body potential strength. In that process, we touched upon the effects our results have on Skyrme energy-density functionals and disentangled the contributions of gradient and bulk terms. We then went on to study several different one-body potential periodicities. After going over the basics of linear-response theory, we were able to make a prediction for the static density-density response function of neutron matter. We found that the neutron-matter response function exhibits similarities to the response of the non-interacting Fermi gas and the electron gas, but also novel behavior. This work opens up a host of \textit{ab initio} calculations for neutron-star crusts and improved descriptions of nuclei.

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