Band structure effects for dripped neutrons
in neutron star crust

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Received 27 May 2004; received in revised form 23 August 2004; accepted 17 September 2004

Available online 8 October 2004

Abstract

The outer layers of a neutron star are supposed to be formed of a solid Coulomb lattice of neutron rich nuclei. At densities above neutron drip density (about one thousandth of nuclear saturation density), this lattice is immersed in a neutron fluid. Bragg scattering of those dripped neutrons by the nuclei which has been usually neglected is investigated, within a simple mean field model with Bloch type boundary conditions. The main purpose of this work is to provide some estimates for the entrainment coefficients, as required for hydrodynamical two fluid simulations of neutron star crust [nucl-th/0402057, astro-ph/0408083], which relate the momentum of one fluid to the particle currents of the other two fluids [Sov. Phys. JETP 42 (1976) 164]. The implications for the equilibrium neutron star crust structure are also briefly discussed.

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1. Introduction

In a typical neutron star (with mass about one and a half that of the sun), a solid phase of neutron rich nuclei, hence forming the crust, is expected to occur from the surface down to about \( \approx 1 \) km in depth where the pressure and density are so high that the nuclei may be strongly deformed and adopt “exotic” shapes [4]. Inside the crust, above the neutron drip density \( \rho_{\text{drip}} \approx 10^{11} \text{ g cm}^{-3} \), free neutrons are found to coexist with the nuclei. Below
the crust, the nuclei merge into a uniform mixture of nucleons and perhaps of some other mesons.

Bragg scattering of the dripped neutrons by the nuclei, which is analogous to the scattering of valence electrons by ions in ordinary solids, has been usually neglected whereas it may be of quantitative importance for evaluating the processes involved in phenomena such as pulsar glitches. This issue has been pointed out by a few authors [5–7] and band effects, namely the opening of band gaps in the single particle energy spectrum, have been actually estimated in a simplified one-dimensional case [8].

In a proceeding work [1,2] we have introduced a macroscopic effective mass $m_\text{\$}$ relevant for hydrodynamical simulations, such that the momentum of the neutron superfluid is given by the mean neutron velocity times this effective mass in the crust rest frame. In a (super)fluid mixture, in general the momentum of one species is a linear combination of the particle currents of the other components. This is known as the entrainment effect [3]. We have shown how to obtain these entrainment coefficients in a two fluid neutron star model from the effective mass $m_\text{\$}$. Our analysis is complementary to the two fluid description of neutron star core, based on the Fermi liquid theory (see Borumand et al. [10] and references therein), whose relativistic generalisation has been recently performed by Comer et al. [11] via a $\sigma-\omega$ mean field model.

Particle transport in the inner crust is induced by displacements of the whole superfluid and it can thus be seen [9] that it is therefore not very sensitive to the superfluid energy gap unlike thermal properties such as the specific heat. The main effect of the pairing interaction is actually to lead to a smearing of the sharp neutron Fermi surface. Since at the densities of interest, the pairing gap is typically much smaller than the Fermi energy, corrections of the effective mass due to pairing correlations are expected to be small. We shall therefore neglect as a first approximation pairing interaction (as in the works of Borumand et al. [10] and Comer et al. [11]).

Since the pioneering Hartree–Fock calculations of Negele and Vautherin [12], more or less arbitrary boundary conditions have usually been applied. In particular the so-called “Wigner–Seitz approximation”, whereby nuclei are treated as if isolated, has been widely used. Whereas this approximation would be appropriate for describing nucleons that are clustered around lattice nodes, it is more questionable for the dripped neutrons which are delocalized over the crust. This means that one has necessarily to consider the whole medium and apply global boundary conditions. In the case of a periodic lattice of nuclei, as a result of the well-known Floquet–Bloch theorem [13], the problem can still be reduced to the so-called Wigner–Seitz cell (not to be confused with the W–S sphere) namely some “elementary” polyhedron surrounding a nuclei whose shape is completely determined by the lattice structure, supplemented with Bloch type boundary conditions. The single particle energy spectrum is composed of a set of analytic functions $E_\alpha(k)$ (in the following we shall use brackets for functional dependence in order to avoid confusions with mere multiplication) of the momentum $k$ (related to the translational symmetry), each sheet in momentum space being specified by a band index $\alpha$ (associated with rotational symmetry).

Recently Magierski et al. [6] have suggested that shell effects arising from “unbound” neutrons may be important for neutron star crust structure and have carried out Skyrme–Hartree–Fock calculations in a cubic box with periodic boundary conditions. However, such a choice of boundary conditions does not properly account for Bragg scattering since
the ensuing single particle states are only those associated with vanishing Bloch momentum. Besides the Wigner–Seitz cell, which is taken as a cube there, is cubic only in a simple cubic lattice.

In order to prepare the way to more realistic calculations, we shall briefly review band theory and some key solid state physics techniques which may not be familiar to the reader. We shall then discuss some results for dripped neutrons in the inner crust, described within a mean field model.

2. Inner layer of the crust

2.1. Mean field model

In the absence of any previous calculation of neutron band structure except for a simple 1D model [8], even at the simplest level of approximation, we shall adopt the single particle model of Oyamatsu and Yamada [8], to estimate the effective mass of the dripped neutrons in the bottom layers of the inner crust. Previously, we only considered slab shaped and rod like nuclei for simplicity [1]. The effective mass was found to be very close to the bare mass. The reason was that the region we were focusing on was nearly homogeneous. In the present work, we shall extend the same analysis for lower densities where the crust is assumed to be composed of spherical nuclei arranged in a body centered cubic lattice. Simple cubic and face centered cubic structures will also be considered for comparison. We shall also report results for the spherical and cylinder hole phases near the crust–core transition layer.

In the model suggested by Oyamatsu et al. [8], both bound and dripped neutrons are assumed to be described as independent particles moving in a “background” phenomenological mean field. The neutron single particle states thus obey the following Schrödinger equation

\[ -\frac{\hbar^2}{2m} \Delta \varphi + V \varphi = E \varphi. \quad (1) \]

We further assume that nuclei are located at the nodes of an infinite crystal lattice. The single particle periodic potential \( V \) is thus constructed from the potential \( U \) of Oyamatsu and Yamada given in the Wigner–Seitz sphere by the following procedure:

\[ V(r) = \sum_T U(r - T), \quad (2) \]

where the sum goes over all lattice sites, i.e., \( T = \sum_{\ell=1}^3 \ell^a e_a \), where \( \ell^a \) are integers and \( e_a \) basis lattice vectors. The potential \( U \) was defined by Oyamatsu and Yamada as (Eq. (2.5) of Ref. [8])

\[ U(r) = \left( \frac{1}{\sqrt{\pi \kappa}} \right)^3 \int d^3 r' U_0(r') e^{-|r-r'|^2/\kappa^2}, \quad (3) \]

\[ U_0(r) = \delta U \frac{\delta}{\delta n_n}. \quad (4) \]
Fig. 1. Single particle central potential (left) and spin–orbit coupling potential (right) in the W–S sphere for the lowest baryon density considered $n_b = 0.03 \text{ fm}^{-3}$ from model I of Ref. [8].

where $v$ is a potential energy density functional of homogeneous nuclear matter. The local neutron and proton densities were obtained from a zero temperature Thomas–Fermi calculation in the W–S approximation. The energy density functional was parametrised such as to reproduce the properties of terrestrial nuclei on the $\beta$ stability line and the equation of state of Friedman and Pandharipande for both pure neutron matter and symmetric nuclear matter. The parameter $\kappa$ (as well as the spin–orbit coupling parameters, see Section 4 below) was adjusted so as to give reasonable values for the single particle energies of $^{208}$Pb (for further details, see [8,31]). We have used the parameter sets of model I from Ref. [8]. We have neglected the spin–orbit coupling introduced by Oyamatsu and Yamada since it is smaller by an order of magnitude compared to the central potential as shown on Fig. 1.

In order to take into account Bragg scattering, single particle states have to satisfy the Floquet–Bloch theorem

$$\phi_k(r + T) = e^{i k \cdot T} \phi_k(r).$$

As a consequence, the Schrödinger Eq. (1) can be solved within any primitive cell [13], among which the Wigner–Seitz one possesses the full symmetry of the lattice, with boundary conditions given by (5). The Wigner–Seitz (W–S) cell is defined as the set of points that are closer to a given lattice node than to any other. Examples are shown for cubic crystals on Fig. 2. Except for simple cubic lattice, the W–S cell is a complicated polyhedron. In the work of Oyamatsu and Yamada, the lattice spacing was defined as $a_{OY} = V_{cell}^{1/3}$, where $V_{cell}$ is the volume of the W–S sphere. In the present work, we have defined the lattice spacing $a$ for cubic lattices (cube length) such that the volume of the W–S polyhedron is equal to that of the sphere.

2.2. Single particle energy spectrum

For each given $k$, Eq. (1) with boundary conditions (5) will admit a discrete set of eigenvalues $E_\alpha$ (indexed by a Greek letter) referred as bands. Single particle energies will be ordered such that $E_1 < E_2 < \cdots$ which guarantees that for a given band, say $\alpha$, the energy $E_\alpha(k)$ will be continuous (besides it is analytic except for high symmetry points). Unlike the W–S approximation, the single particle energy spectrum in the Bloch case has a much more detailed structure.
It can be shown that the single particle energy is periodic in a “reciprocal” lattice [13] whose nodes are given by $\mathbf{K} = \sum_{a=1}^{3} \ell_a \mathbf{I}^a$, where $\ell_a$ are integers and the dual basis is defined by the dot products

$$\mathbf{I}^a \cdot \mathbf{e}_b = 2\pi \delta_{ab}.$$ (6)

This means in particular that Eq. (1) has only to be solved for each momentum $\mathbf{k}$, within a domain known as the first Brillouin zone (BZ), i.e., the W–S cell of the reciprocal lattice. By considering rotational symmetry, it can be shown that all the relevant information about the spectrum is contained within some “irreducible” domain of the BZ [14]. The solutions of Magierski et al. [6] are only those associated with the center of the first Brillouin zone, namely $\mathbf{k} = 0$, provided the lattice is simple cubic (for body centered or face centered cubic structures, their solutions are not of the Bloch type).

We have solved the Schrödinger equation (1) with Bloch type boundary conditions (5) variationally by a finite plane wave expansion of the single particle wavefunctions, namely

$$\varphi_{\mathbf{k}}(\mathbf{r}) = \frac{1}{\sqrt{V_{\text{cell}}}} \sum_{\mathbf{K}} \tilde{\varphi}_{\mathbf{k}}(\mathbf{K}) e^{i(\mathbf{k} + \mathbf{K}) \cdot \mathbf{r}},$$ (7)

with an energy cutoff $\varepsilon_{\text{cutoff}}$ such that

$$\frac{\hbar^2 (\mathbf{k} + \mathbf{K})^2}{2m} < \varepsilon_{\text{cutoff}}.$$ (8)

The nuclear structures that we have considered are illustrated on Fig. 3.

2.3. Effective mass

The zero temperature many body ground state is merely obtained by filling all the single particle states whose energy $\varepsilon_{\alpha}[\mathbf{k}]$ is lower than the Fermi energy $\mu$. For each total neutron density $n_\alpha$, the Fermi energy $\mu$ is thus determined via the integral over the first Brillouin zone (BZ)

$$n_\alpha = \frac{g}{(2\pi)^3} \sum_{\mathbf{K} \in \text{BZ}} \int d^3k \delta \left\{ \mu - \varepsilon_{\alpha}[\mathbf{k}] \right\}.$$ (9)
where we have explicitly introduced the Fermi distribution by the Heaviside unit step distribution $\vartheta \{x\} = 1$ if $x > 1$ and zero otherwise, allowing each momentum state to be $g$-fold degenerate (here $g = 2$ to account for the spin degeneracy). In particular, the ground state will be characterized by the Fermi surface bounding the occupied part of momentum space. In general the Fermi surface will consist of disconnected pieces, each piece being given by $\mathcal{E}_\alpha \{\mathbf{k}\} = \mu$.

Introducing the dripped or free neutron density $n$, the effective mass (from which the entrainment coefficients can be obtained [2]) is defined [1] by the simple formula

$$m_* = \frac{n}{\mathcal{K}}. \tag{10}$$

where the mobility scalar $\mathcal{K}$ is given by (summation over energy bands is implicit)

$$\mathcal{K} = \frac{1}{3} \left( \frac{g}{2\pi} \right)^3 \oint_F v \, dS_F, \tag{11}$$

and the group velocity is defined by the usual expression

$$v = \frac{1}{\hbar} \nabla_{\mathbf{k}} \mathcal{E}. \tag{12}$$

This mobility scalar is related to the electric conductivity $\sigma$ by the formula $\sigma = e^2 \tau \mathcal{K}$ in which $e$ and $\tau$ are the electric charge per particle and some characteristic relaxation time, respectively. In terms of the effective mass, this formula reduces to the well-known expression [13] $\sigma = ne^2 \tau / m_*$. 

In the following sections, we have defined the dripped neutron states to be those states whose energy is positive $\mathcal{E}_\alpha \{\mathbf{k}\} > 0$ (the energy origin being taken as the largest possible value of the potential).
2.4. Wigner–Seitz approximation

The W–S cell of body or face centered (unlike simple) cubic lattice is a nearly spherical polyhedron, this is the origin of the so-called W–S approximation [15], according to which the W–S cell is taken as a sphere and more or less arbitrary boundary conditions are applied on the single particle wave functions. For instance, the prescription given by Negele and Vautherin [12] requires that on the sphere radius, radial wave functions with even orbital angular momentum quantum number $l$ vanish and derivatives of the radial wave functions with odd $l$ vanish. Besides the density needs to be averaged in the vicinity of the cell edge in order to prevent the occurrence of unphysical density fluctuations. Within the current framework, the Floquet–Bloch theorem (5) ensures that the local total neutron density defined by

$$n_\alpha(r) = \frac{g}{(2\pi)^3} \sum_\alpha \int_{BZ} d^3 k |\psi_\alpha(k)|^2 \delta \left\{ \mu - E_\alpha(k) \right\}$$ (13)

possesses the lattice symmetry, the single particle wavefunctions being normalized as

$$\frac{1}{V_{\text{cell}}} \int_{\text{cell}} d^3 r |\psi_\alpha(k)|^2 = 1.$$ (14)

This means in particular that the density gradient $\nabla n_\alpha(T)$ will vanish whenever the crystal is left invariant under space inversion, which means that the density will be nearly constant (to first order) inside nuclei. In cases for which $r$ belongs to a symmetry plane, the density gradient will have no components orthogonal to this plane. This means that the density profile will be essentially flat (to first order) around the cell boundary, along directions perpendicular to the W–S faces whenever the face is parallel to any symmetry plane. This is true for the simple cubic and face centered cubic W–S cells. This is also true for the body centered cubic W–S cell excluding the hexagonal faces.

3. Numerical methods

3.1. Brillouin zone integrations

Brillouin zone integrations are involved in many solid state physics calculations, as, for instance, in the particle density (13). Analytic expressions for the integrand are usually unknown (except for a few academic cases), therefore one has to rely on numerical schemes based on discrete summations. Since the computation of energy bands is usually the most time consuming part of solid state calculations, several techniques have been developed in order to compute these integrals with the smallest number of terms. One of the most successful methods, which was pioneered by Baldereshi [16] and subsequently improved by Chadi–Cohen [17], is based on a weighted sum over a set of special symmetry points. The main idea relies on the fact that a function $f(k)$ which has the full symmetry of the reciprocal lattice, can be expanded over symmetrized plane waves (in the following we
shall consider only cubic structures, for which there is no glide plane or screw symmetry axis \([32]\))

\[
\tilde{f}_0 = \frac{1}{V_{BZ}} \int_{BZ} f(\mathbf{k}) \, d^3k = \sum_{j=1}^{N} w(\mathbf{k}_j) f(\mathbf{k}_j) + R_N, 
\]

where \(P\) is any rotation of the lattice and \(|P|\) their number (lattice vectors are ordered as \(|T_{s_1}| \geq |T_{s_2}|\) whenever \(s_1 > s_2\)). The integral of this function over the first Brillouin zone is rewritten as a sum over a set of unequivalent points \(\mathbf{k}_j\) with weight \(w(\mathbf{k}_j)\):

\[
\tilde{f}_0 = \frac{1}{V_{BZ}} \int_{BZ} f(\mathbf{k}) \, d^3k = \sum_{j=1}^{N} w(\mathbf{k}_j) f(\mathbf{k}_j) + R_N, 
\]

where the residual \(R_N\) is given by

\[
R_N = -\sum_{s>0} \sum_{j=1}^{N} w(\mathbf{k}_j) \frac{1}{|P|} \sum_{P} e^{iP \cdot T_s \cdot k}. 
\]

By choosing the points \(\mathbf{k}_j\) such that all plane waves are exactly zero for all \(s\) up to some largest \(s_{\text{max}}\), the residual will contain only terms with lattice vectors \(|T| \geq |T_{s_{\text{max}}}|\) for which the contribution will be vanishingly small provided the function \(f(\mathbf{k})\) is sufficiently smooth. Eventually the integral is approximated by

\[
\frac{1}{V_{BZ}} \int_{BZ} f(\mathbf{k}) \, d^3k \simeq \sum_{j=1}^{N} w(\mathbf{k}_j) f(\mathbf{k}_j). 
\]

The determination of these special points \(\mathbf{k}_j\) and their associated weights \(w(\mathbf{k}_j)\) was rather complicated in the first scheme involving a recursion process. Several authors have provided formulae (see [18] and references therein).

The convergence of the discrete sum with respect to the number of special points is exponential for smooth integrands and an error of less than one per cent can be attained within a few points only. Unfortunately the presence of a sharp Fermi surface (especially in any metallic solid) introduces discontinuities in the integrands, such as in Eq. (13), which spoil the convergence. This method is thereby most satisfactory for insulating materials and semiconductors. Nevertheless, accurate results can still be obtained with a small number of special points by "smearing" the Fermi surface [19]. Since the volume of the first Brillouin zone and of the Wigner–Seitz cell are related by

\[
V_{BZ} = \frac{(2\pi)^3}{V_{\text{cell}}}, 
\]

the number of special points for a given precision is smaller for larger cells. We have tested the special point method by calculating the integral

\[
\frac{1}{V_{BZ}} \int_{BZ} k^2 \, d^3k, 
\]
which is equal to 1/4, 3/8, 19/32 in units of $(2\pi/a)^2$ for simple cubic, body centered cubic and face centered cubic lattice, respectively. The convergence is illustrated on Fig. 4.

We have also shown on Fig. 5 how the smearing procedure of Methfessel et al. [19] could improve the convergence in the extreme case of one single special point, by computing the total neutron density $n_n$ as a function of the Fermi energy $\mu$ defined in Eq. (9) in the Shockley “empty” lattice test [20] for which we have

$$n_n = \frac{1}{3\pi^2} \left( \frac{\sqrt{2m\mu}}{\hbar} \right)^3. \quad (22)$$
3.2. Fermi energy

The determination of the Fermi energy is numerically expensive as it requires the calculation of all occupied single particle states (unlike the effective mass which depends only on the highest occupied states on the Fermi surface). The computational cost is significant since the number of levels is of the order of several hundreds up to nearly one thousand. Nonetheless a useful estimate of the Fermi energy with an accuracy of a few percents can still be easily obtained as follows. First, among all occupied states, one should distinguish between core states whose wavefunction is localised in the neighborhood of nuclei and valence or conduction states whose wavefunction extends over all space. The core states are easily obtained by finding the bound states of one isolated nucleus. Having found the number of such states, let us say \( N_{\text{core}} \), Eq. (9) can be written as

\[
n_n = \frac{g}{(2\pi)^3} V_{\text{BZ}} N_{\text{core}} + \frac{g}{(2\pi)^3} \sum \int_{\text{BZ}} d^3 k \vartheta \{ \mu - E_\alpha[\mathbf{k}] \} \vartheta \{ E_\alpha[\mathbf{k}] - E_{\text{core}} \}, \tag{23}
\]

where \( E_{\text{core}} \) is the energy of the highest core state. The determination of the Fermi energy is essentially a state counting procedure which becomes less and less sensitive to the detailed structure of the energy spectrum as the number of particles is increased. Consequently assuming that the valence states can be roughly described as non-interacting particles, the previous equation thus leads to the following estimate:

\[
\mu \simeq E_{\text{core}} + \frac{\hbar^2}{2m} \left( n_n - n_{\text{core}} \right) \frac{6\pi^2}{g}^{2/3}, \tag{24}
\]

where from (20) we have \( n_{\text{core}} = g N_{\text{core}}/V_{\text{cell}} \).

For instance, for the baryon density \( n_b = 0.03 \text{ fm}^{-3} \), we found for the core states: three \( s \) states, two \( p \) states, two \( d \) states, one \( f \) state and one \( g \) state. The total number of core states is thus equal to \( N_{\text{core}} = 35 \) and the Fermi energy is approximately given by \( \mu \simeq 18.02 \text{ MeV} \), which is about 5\% larger than the value \( \mu = 17.16 \text{ MeV} \), obtained by direct integrations of (9) via special points. The value of the Fermi energy was also found to be completely unsensitive to the lattice structure, which is due to the fact that Eq. (9) does not explicitly depend on the precise values of energy bands but only on their number in a given energy range.

3.3. Fermi surface integrations

Calculations of the effective mass (10) require a fine mesh sampling of the Fermi surface. One of the most accurate methods for performing Fermi surface integrations is based on the Gilat–Raubenheimer (GR) scheme [21]. The integration over the Fermi surface is first reduced by symmetry to the irreducible wedge of the first Brillouin zone which is partitioned into microcells. The integral is thereby discretized as a sum over the microcells with suitable weight factors to account for the fact that microcells may overfill each wedge. The Fermi surface is then approximated by a plane inside the microcells which it intersects, so that the integration within each cell can be calculated analytically [22].

We have implemented the original linear extrapolation method with cubic cells [23]. The energy is linearly extrapolated from the values of the energy and velocity at the center
of the cell. This scheme therefore requires the energy bands as well as their gradients, whose evaluation with plane waves is straightforward via the Hellmann–Feynman theorem [24], namely
\[ v = \frac{\hbar k}{m} + \sum_{K} \frac{\hbar K}{m} |\tilde{\phi}_k(K)|^2, \] (25)
with the normalisation of the wavefunction given by
\[ \sum_{K} |\tilde{\phi}_k(K)|^2 = 1. \] (26)

The GR method seems to be not so popular as the tetrahedron method in solid state physics, in which the irreducible domain is divided into tetrahedra and the energy is linearly interpolated from its values at the four corners of each tetrahedron [25,26]. One of the main reasons is that the calculation of the velocity may be computationally expensive with more elaborate basis functions than plane waves. Nevertheless the linear extrapolation of the energy within the GR method, which is a first order Taylor expansion around the cell center, is a better approximation than an interpolation. Since in the present case the integrand in Eq. (11) involves the energy gradient, the GR method seems much more appropriate. Besides interpolation faces the band crossing problem: the ordering of the energy bands with increasing energy ensures that the energy is continuous for a given band but does not prevent discontinuities in the energy gradient from occurring. Unlike extrapolation, interpolation will yield an incorrectly small gradient resulting in a loss of convergence (see [27–29] for a discussion about systematic errors).

In order to test this scheme, we have employed the “cubium” toy model, i.e., a simple cubic crystal with one single s band whose momentum dispersion in Cartesian coordinates is given by \( E\{k\} = - (\cos k_x + \cos k_y + \cos k_z) \). Analytic expressions for the density of single particle states at energy \( E \), defined by
\[ \mathcal{N}(E) = \frac{1}{(2\pi)^3} \frac{g}{\hbar^2} \oint dS E v \] (27)
are available in terms of the elliptic integral of the first kind [30]. As can be seen on Fig. 6 \((g = 2)\) the GR method yields results in very good agreement with the analytic expressions, in particular in the vicinity of the critical points associated with \( v = 0 \) (producing the “kinks” in the density of states).

4. Results

We have considered the three types of cubic lattices, namely simple cubic, face centered cubic and body centered cubic for spherical nuclei and bubbles (i.e., periodic lattice of “holes” containing dripped neutrons in uniform nuclear matter). The Fermi surface integrations have been carried out with of the order of \( \sim 10^3 \) cubic cells, for which the relative deviation was a few percents as shown on Figs. 7 and 8.

The effective mass \( m_* \) shown on Fig. 10 has been found to be greatly enhanced due to Bragg scattering, reaching \( m_* \sim 15m \) at a total baryon density \( n_b = 0.03 \, \text{fm}^{-3} \). Deeper
Fig. 6. Density of states of the cubium model computed with the GR method and compared with the analytic expressions given by Morita et al. [30].

Fig. 7. Convergence of the effective neutron mass $m_\star/m$ with respect to the number of GR cells, for baryon density $n_0 = 0.055 \text{ fm}^{-3}$ with nuclei in a body centered cubic lattice.
Fig. 8. Convergence of the Fermi surface area $S/S_{gas}$ with respect to the number of GR cells, for baryon density $n_b = 0.055 \text{ fm}^{-3}$ with nuclei in a body centered cubic lattice.

inside the crust, the renormalization is smaller, tending to $m_* \sim m$ at very high density as the single particle potential tends to a constant. We have also shown on Fig. 11 the transverse effective neutron mass in the “exotic” phases [1] including the “antispaghetti” layer, where non-spherical nuclei occur (only the velocity components respectively perpendicular to the slabs or to the rods, for the “lasagna” or “(anti)spaghetti” phases are included in Eq. (11) for the mobility scalar; the other velocity components have the same expressions as those of a non-interacting gas and therefore the corresponding effective mass coincide with the ordinary mass). The dependence on the lattice structure is rather weak. Numerical results are summarized in Appendix A. We have also carried out the calculations with the estimate of the Fermi energy suggested in Section 3.2. While this is usually a good approximation for bulk quantities such as the conduction neutron density, the errors are much larger for quantities depending only on states at the Fermi level. In particular, the resulting effective mass is about 20% lower at baryon density $n_b = 0.03 \text{ fm}^{-3}$, having value $m_*/m \simeq 14$, where the relative error in the conduction neutron density is less than one percent.

In order to understand the origin of this effective mass enhancement, we have compared the Fermi surface area $S_F$ with its expression obtained from the assumption that the dripped neutrons form an ideal Fermi gas of density $n$, which is given by

$$S_{gas} = 4\pi g (6\pi^2 n/g)^{2/3}. \quad (28)$$

We have found that the Fermi surface area is strongly reduced in the outer layers compared to that of the sphere, whereas it tends to its uniform expression (28) near the crust–core interface as illustrated on Fig. 9. It is first to be remarked that the “enclosed” Fermi volume depends only on the density, as is easily seen on Eq. (9) and therefore does not depend on the particular shape of the Fermi surface. It is well known that among all the closed surfaces of a given volume, the sphere has the minimum surface area. In the present case,
however, the opening of numerous band gaps in the single particle energy spectrum leads to a Fermi surface composed of disjoint open pieces. By band gaps, we mean avoided crossings of energy band sheets in $k$ space. This is the reason why the Fermi surface area is found to be smaller than that of the sphere in the low density layers of the crust where the gaps take their largest values.
It is quite instructive to evaluate the mean Fermi velocity $\bar{v}_F$ defined by

$$\bar{v}_F = \frac{1}{S_F} \int_F v \, dS_F. \quad (29)$$

It is readily seen that it is expressible, in terms of the Fermi surface area $S_{gas}$ and Fermi velocity $v_{gas}$ of the non-interacting neutron gas, as

$$\bar{v}_F v_{gas} = \frac{S_{gas}}{S_F} \frac{m}{m^*}. \quad (30)$$

For instance, at the lowest density we considered, namely $n_b = 0.03$ fm$^{-3}$, the mean Fermi velocity is about one third the Fermi velocity of the gas. The ratio increases to one half at $n_b = 0.055$ fm$^{-3}$ and tends to one as the density approaches the crust–core transition density. This is a clear and intuitive manifestation of Bragg scattering.

The spin–orbit coupling, as we have seen, is very small compared to the central potential in the bottom layers we have considered (see Fig. 1). However, such a term would probably not be negligible in the lower density regions of neutron star crust. The presence of this term in the Schrödinger equation (1) preserves the lattice symmetry which implies that single particles wave functions can still be expressed as Bloch states. However, the spin–orbit coupling breaks the spin symmetry and therefore raise some degeneracy. As a result, this will open new gaps in the energy spectrum, in the vicinity of which the group velocity will be vanishingly small. Consequently, it can be seen from (11) and (10) that the resulting effective masses would be expected to be increased by the spin–orbit coupling since the density would not be much affected. This conclusion is confirmed by a numerical calculation in the “spaghetti” phase. The computations are quite involved since the size of the secular (complex) matrices is doubled. We have taken into account the spin–orbit coupling given by Oyamatsu and Yamada (Eq. (2.12) of Ref. [8]), which is given in the
W–S approximate cell (a cylinder in this case directed along the $z$ axis) by an expression of the form

$$V_{LS}(r) = \frac{1}{r} \left( \lambda_1 \frac{dn_b}{dr} - \lambda_2 \frac{d}{dr} (n_n - n_p) \right) \frac{1}{2} l_z \sigma_z,$$

(31)

where $r$ is the distance to the cylinder axis, $l_z$ the angular momentum component along the cylinder axis and $\sigma_z$ the diagonal Pauli spin matrix. The parameters $\lambda_1$ and $\lambda_2$ were adjusted so as to give the correct sequence of single particles states of $^{208}$Pb. We have used the values of model I from Ref. [8] namely $\lambda_1 = 175.8$ MeV fm$^5$ and $\lambda_2 = 16.39$ MeV fm$^5$.

We have carried out the calculation at the total baryon density $n_b = 0.06$ fm$^{-3}$ for the hexagonal lattice. Since the densities varies smoothly, the spin–orbit coupling has a small effect, yet the opening of band gaps can be clearly observed as shown on Fig. 13 (see also Fig. 14). The transverse effective mass is found to be slightly larger with than without spin–orbit coupling as expected, respectively, $m_{\perp}^*/m = 1.37$ and $m_{\perp}^*/m = 1.35$ while the Fermi surface area is smaller, respectively, $S_F/S_{\text{gas}} = 0.88$ compared to $S_F/S_{\text{gas}} = 0.89$.

5. Conclusion

We have considered the band effects in neutron star crust induced by Bragg scattering of dripped neutrons upon the solid lattice of nuclei. This is the analog in a nuclear context, of a well-known phenomenon in ordinary solid state physics whereby valence electrons are being scattered by ions leading notably to an understanding of semiconductor band gaps. Based on a mean field model with Bloch type boundary conditions, we have computed the effective dripped neutron macroscopic mass $m_\star$ necessary for evaluating the entrainment coefficients in hydrodynamical descriptions of neutron stars [1,2]. We found that this effective mass is increased by several hundred percents compared to the ordinary neutron mass in the middle layers of the inner crust but becomes negligible near the crust–core boundary where nuclei merge into a uniform mixture of neutrons, electrons and protons (and perhaps of some other mesons). We have also
Fig. 13. Neutron band structure, with (thin lines) and without (thick lines) along high symmetry directions shown on Fig. 12 for an hexagonal lattice of spaghetti (rod) like nuclei at baryon density $n_b = 0.055$ fm$^{-3}$.

Fig. 14. Zoom of the region in the box of Fig. 13. Band splittings induced by spin–orbit coupling can be clearly seen.

found that the lattice structure plays a minor role. However, the differences are significant between 3D crystals and the 1D or 2D liquid crystal configurations that were previously studied within the same model (respectively, slab shaped and rod like nuclei).

The present results also suggest that the widely used Wigner–Seitz approximation, whereby Bragg scattering is neglected, may be questionable for the study of neutron star crust. In particular, it is to be remarked that the very large values of the effective mass $m_*$ that we found in the middle layers tend to indicate that dripped neutron shell (actually band) effects may have significant consequences for the neutron star crust equilibrium.
structure compared to that obtained in the W–S approximation. It has been recently shown that the composition of the inner crust of neutron star is quite sensitive to the proton shell effects [33]. The unbound neutron shell effects have been investigated by Magierski and Heenen [6] who suggested that such quantum effects play an important role in the determination of the neutron star crust equilibrium structure since the energy difference between different nuclear shapes, lattice structures and compositions is typically small. Their analysis was based on a semiclassical approach. They have recently carried out Skyrme–Hartree–Fock calculations [34] but with ordinary periodic boundary conditions. The implementation of Bloch type boundary conditions within more realistic models, especially taking into account neutron pairing as outlined recently [9], should therefore be further investigated.

Acknowledgements

The author would like to thank B. Carter, P. Haensel and D.G. Yakovlev for instructive conversations.

Appendix A. Numerical results

Effective dripped neutron mass $m_\ast$ for spherical nuclei and bubbles in a body centered cubic lattice:

<table>
<thead>
<tr>
<th>$n_b$ (fm$^{-3}$)</th>
<th>$n_n$ (fm$^{-3}$)</th>
<th>$n/n_n$</th>
<th>$m_\ast/m$</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.03</td>
<td>0.029</td>
<td>0.94</td>
<td>15.4</td>
</tr>
<tr>
<td>0.055</td>
<td>0.053</td>
<td>0.95</td>
<td>5.5</td>
</tr>
<tr>
<td>0.0855</td>
<td>0.0826</td>
<td>0.98</td>
<td>1.07</td>
</tr>
</tbody>
</table>

Transverse effective dripped neutron mass for “pasta” phases: “spaghettis” (hexagonal lattice), “lasagna” and cylinder holes (hexagonal lattice):

<table>
<thead>
<tr>
<th>$n_b$ (fm$^{-3}$)</th>
<th>$n_n$ (fm$^{-3}$)</th>
<th>$n/n_n$</th>
<th>$m_\ast^\perp/m$</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.06</td>
<td>0.0581</td>
<td>0.9553</td>
<td>1.347</td>
</tr>
<tr>
<td>0.065</td>
<td>0.063</td>
<td>0.9578</td>
<td>1.242</td>
</tr>
<tr>
<td>0.07</td>
<td>0.0678</td>
<td>0.9606</td>
<td>1.174</td>
</tr>
<tr>
<td>0.074</td>
<td>0.0716</td>
<td>0.9633</td>
<td>1.124</td>
</tr>
<tr>
<td>0.076</td>
<td>0.0735</td>
<td>0.9634</td>
<td>1.031</td>
</tr>
<tr>
<td>0.0775</td>
<td>0.0749</td>
<td>0.9646</td>
<td>1.029</td>
</tr>
<tr>
<td>0.08</td>
<td>0.0773</td>
<td>0.9666</td>
<td>1.025</td>
</tr>
<tr>
<td>0.082</td>
<td>0.0792</td>
<td>0.9687</td>
<td>1.020</td>
</tr>
<tr>
<td>0.083</td>
<td>0.0802</td>
<td>0.9734</td>
<td>1.010</td>
</tr>
<tr>
<td>0.084</td>
<td>0.0811</td>
<td>0.9746</td>
<td>1.003</td>
</tr>
<tr>
<td>0.085</td>
<td>0.0821</td>
<td>0.9762</td>
<td>1.001</td>
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References