APPLICATION OF DENSITY FUNCTIONAL THEORY IN NUCLEAR STRUCTURE

By

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ABSTRACT

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The nuclear density functional theory (DFT) is a microscopic self-consistent framework suitable for describing heavy nuclei and performing large-scale studies. In this dissertation I discuss my research works on the development and application of the Skyrme nuclear-DFT framework, covering a broad range of topics, including the nucleon localization in rotating systems, the origin of reflection-asymmetric deformations, the parameter calibration for beta decays, and the development of a new coordinate-space DFT solver.

The nucleon localization function (NLF), discussed in the first part, is a useful tool for the visualization of structure information. It has been utilized to characterize clustering and shell structure. How the NLF pattern evolves in rotating systems, how it visualizes internal nuclear structure, and how it is connected with single-particle (s.p.) orbits are discussed in this dissertation. The second part deals with nuclei having reflection-asymmetric shapes, which are important candidates for the search of permanent electric dipole moments. In this dissertation, the origin of the pear-like deformation is investigated through both the multipole expansion of the energy density functional and the spectrum of canonical s.p. states. Theoretical predictions of beta-decay rates are discussed next; they are important for r-process simulations that involves nuclei whose experimental beta-decay data are unknown. To provide reliable predictions with quantified uncertainties, the $\chi^2$ optimization is performed to constrain parameters that significantly affect beta-decay transitions in proton-neutron finite-amplitude-method calculations. Besides a well calibrated functional, a reliable and efficient DFT solver is also crucial. The Hartree-Fock-Bogoliubov (HFB) method in the
coordinate space is preferred for deformed and weakly bound nuclei, as solvers based on basis expansions often have difficulty correctly describing continuum effects. A new HFB solver based on the canonical-basis HFB formalism in the three-dimensional coordinate space is developed in this dissertation. It is a well parallelized solver and has been carefully benchmarked against other established HFB solvers.
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Chapter 1

Introduction

1.1 Nuclear density functional theory (DFT)

Being a many-body quantum system, the atomic nucleus is difficult to describe with theoretical and computational models. The nuclear density functional theory (DFT) [1, 2] provides a microscopic self-consistent mean-field model suitable for studying medium- to heavy-mass nuclei and for performing global surveys over the whole nuclear landscape. In a mean-field model, every nucleon moves approximately in a mean potential generated by other nucleons, and the concept of the mean field is justified by shell effects observed experimentally (e.g., the existence of magic numbers); see Ref. [3] for a comprehensive discussion. This dissertation discusses my research on the applications of the nuclear DFT.

In theoretical and computational studies of nuclear structure, it is of great importance to choose appropriate degrees of freedom and approximations to achieve good descriptions with computational expenses under control. From this perspective, the nuclear DFT model lies between the shell model [4, 5] and microscopic-macroscopic (mic-mac) method [6, 7, 8, 9, 10]. Compared with the DFT model, the mic-mac method is computationally inexpensive but not reliable enough for extrapolations beyond experimentally accessible region, while the \textit{ab initio} approach [11] and shell model can provide beyond-mean-field descriptions but are too computationally expensive for heavy systems and large-scale surveys. Hence, the nuclear
DFT is usually the appropriate choice for such studies.

Within the nuclear-DFT framework, the energy of a system is given by the energy density functional (EDF), and there are three functional forms that are widely used – Skyrme [12, 13, 14, 15], Gogny [16, 17, 18] and relativistic [19, 20] EDFs. Inside each category, there also exist variations and different parameterizations. This dissertation focuses on the application of the Skyrme DFT; the underlying theoretical framework is presented in Chapter 2.

1.2 Nuclear deformation, rotation and nucleonic localization

Nuclear deformation comes from spontaneous symmetry breaking related to the nuclear Jahn-Teller effect [21, 22, 23, 24], a concept originally proposed in molecular physics to explain the geometric distortion of non-linear molecules.

Although the vast majority of isotopes exhibit spherical, prolate or oblate ground-state shapes, there is abundant experimental evidence supporting the existence of stable pear-like deformations which break the reflection symmetry in the intrinsic frame [25, 26]. As even-even nuclei with reflection-asymmetric shapes usually have low-energy negative-parity excitations related to octupole collective modes, they are also referred to as “octupole deformed.” Atoms with pear-shaped nuclei have also been shown to be good laboratories for searching the permanent electric dipole moment [27, 28, 29], which is strongly correlated with the nuclear octupole moment [30]. Within the nuclear-DFT framework, a global survey for reflection-asymmetric ground states of even-even nuclei has recently been conducted in Ref. [31]. Following this work, we investigate the microscopic origin of reflection-asymmetric shapes from two perspectives: the multipole expansion of the EDF and single-particle (s.p.)
spectra. An in-depth discussion on this topic is given in Chapter 4.

A number of nuclear collective modes, such as rotations and vibrations, are related to nuclear deformation. The observation of rotational bands provides abundant information about the nuclear ground-state shape and underlying shell structures, as well as the interplay between collective and s.p. degrees of freedom [32, 33, 34, 35]. This interplay results from the fact that one cannot distinguish slow and fast wave-function components in nuclear systems as is usually done in molecules. Perfect nuclear rotors are rare and the adiabatic approximation is usually inapplicable; thus, the fully self-consistent and microscopic nuclear-DFT model is a good choice for the descriptions of nuclear rotation.

Recently, the nucleon localization function (NLF) has been employed in the nuclear-DFT studies for better visualization of clusters and shell effects. It was originally introduced in electronic-DFT studies to characterize shell structure in atoms and chemical bonds in molecules [36, 37, 38, 39]. In nuclear physics, the NLF has been proved to be a useful tool for the identification and description of (i) clusters in light nuclei [40, 41, 42] and heavy-ion collisions [43], (ii) fragment formation along the fission pathway [44, 45, 46, 47, 48, 49], (iii) nuclear pasta phases in the inner crust of neutron stars [41], and (vi) shell structures of electrons and nucleons in Oganesson [50]. Compared with the particle distribution that is quite smooth in the nuclear interior, the NLF clearly displays the internal structure of a nucleus through its characteristic oscillating or cluster patterns. In Chapter 3 I discuss how the NLF characterizes the nuclear response to rotation and show why the NLF constitutes a powerful visualization tool for nuclear structure studies. Based on my work, the concept of the Pauli kinetic energy is brought up in Ref. [51], which quantifies the effect of the Pauli exclusion principle in the heavy-ion collision.
1.3 Model calibration for beta-decay calculations

How heavy elements are created in the universe is a fascinating open question [52]. Various astrophysical processes contribute to the synthesis of these elements, and both slow and rapid neutron-capture processes (s and r processes) are believed to be predominant [53, 54, 55, 56, 57]. The study of these processes requires ample astronomical observations as well as data on the properties of nuclei involved. Theoretical predictions of beta-decay rates are crucial nuclear inputs for the simulations of some astrophysical processes, especially the r process that traverses the neutron-rich region that experiments cannot currently access. The competition between neutron captures, photodissociation, and beta decays determines how the r process proceeds in different astronomical environments, and the final abundances of stable nuclei are strongly affected by the beta-decay rates of their progenitors [7, 58, 59, 60, 61, 62]. Therefore, the reliability of r-process simulations depends heavily on the quality of beta-decay predictions.

Within the nuclear-DFT framework, the finite amplitude method (FAM) is an efficient approach for the solution of the (quasiparticle) random phase approximation (QRPA) [63, 64]. Thanks to recent developments of the Skyrme proton-neutron FAM (PNFAM), it is now feasible to calculate charge-changing transitions in deformed nuclei and to conduct large-scale calculations for beta-decay rates [65, 62, 66, 67]. In the PNFAM, the time-odd isovector Skyrme couplings, isoscalar pairing strength, and effective axial-vector coupling have a strong impact on beta-decay transitions, but they are not constrained by ground-state properties of even-even nuclei and should thus be calibrated based on experimental data related to beta decays. In this work we carry out the $\chi^2$ optimization for the model calibration, following the procedure discussed in Refs. [68, 69]. After the model calibration
we will be able to provide reliable predictions with quantified uncertainties for $r$-process simulations. Details pertaining to this topic are presented in Chapter 5.

1.4 Reliable and efficient DFT solver in coordinate space

Exotic nuclei, which are far from the valley of stability and hence weakly bound, provide rich information that helps us improve nuclear models; they also play important roles in a number of astrophysical processes. Within the nuclear-DFT framework, the ground state of an even-even nucleus is solved by the Hartree-Fock (HF) + Bardeen-Cooper-Schrieffer (BCS) or Hartree-Fock-Bogoliubov (HFB) methods. The former method is simpler but can be problematic in exotic nuclei, and the full HFB scheme that treats nuclear pairing in a fully self-consistent way should instead be employed for such systems [70, 71, 72, 73, 74]. The HFB calculations of exotic nuclei, however, can be computationally expensive, as a large model space is needed for weakly bound s.p. states; these s.p. states have broad density distributions and are strongly affected by the continuum. The problem becomes even more severe for symmetry-unrestricted calculations.

There have been a number of HF+BCS and HFB solvers developed for nuclear-DFT studies, which can be divided into two categories: Some are formulated in the coordinate-space representation while others are based on basis expansions of wave functions. Table 2 in Ref. [75] gives a summary of these solvers. For exotic nuclei, the coordinate-space HFB solver is a preferred choice as basis-based solvers often have difficulty producing correct asymptotic behaviors of weakly bound s.p. orbits. We have developed a reliable and efficient Skyrme-HFB solver HFBFFT in the three-dimensional Cartesian coordinate representation with no spatial symmetry imposed. It is named after the fast-Fourier-transform (FFT) technique
used for numerical differentiation. The new solver is based on the canonical-basis HFB formalism proposed in Refs. [76, 77] and finds the HFB solution via the damped gradient method [76, 78, 79, 80, 81]. Compared with the quasiparticle basis, the canonical basis does not have an intractably huge level density as canonical states are spatially localized; the computational cost is thus under control. Furthermore, HFBFFT is well optimized and highly parallelized; it is also benchmarked against other established HFB solvers. Numerical details and benchmark results can be found in Chapter 6.

1.5 Organization of this dissertation

This dissertation is organized as follows. Chapter 2 discusses the theoretical formulation of the Skyrme-DFT approach and covers the numerical methods employed in following chapters. The study of the NLF in rotating systems can be found in Chapter 3, which is followed by the investigation about the origin of reflection-asymmetric shapes in Chapter 4. In Chapter 5 I discuss the model calibration for beta-decay calculations, and Chapter 6 describes the new solver HFBFFT. Finally, the conclusions are given in Chapter 7.
Chapter 2

Theoretical framework

The foundation of the DFT is the Hohenberg-Kohn (H-K) theorem [82] and Kohn-Sham (K-S) equation [83]. Here only the main points of the formalism are discussed, while a comprehensive discussion can be found in Ref. [84].

The H-K theorem states that for a many-body quantum system in an external field $v_{\text{ext}}(x)$, there exists a unique energy functional $E_v[\rho]$ of the density $\rho(x)$ such that

$$E_v[\rho] = F[\rho] + \sum d x v_{\text{ext}}(x) \rho(x),$$  \hspace{1cm} (2.1)

which is minimized if and only if $\rho(x)$ is the ground-state density. Here $x = (r, s, \tau)$ includes spatial, spin, and isospin coordinates. The $F[\rho]$ part is independent of the external field and thus universal. Although the H-K theorem does not tell how $F[\rho]$ should be constructed, it allows us to search for or to guess an approximate functional. The K-S theory suggests a practical approach to minimize the energy functional: It constructs an auxiliary system of non-interacting particles in the K-S potential, whose density is identical to the true ground state of interacting particles in the external field $v_{\text{ext}}(x)$. The many-body wave function of the auxiliary system is a Slater determinant, simpler to handle than the true wave function which is a superposition of Slater determinants.

The DFT framework was originally developed in the context of electronic systems. In a
nuclear system, however, a Slater determinant given by the K-S equation is not enough to account for pairing correlations, so the HFB method has been utilized, in which the nuclear system is described by a product state consisting of non-interacting quasiparticles. In the following I outline the theoretical framework of the Skyrme DFT, which has been widely employed for the global studies of various nuclear properties, such as ground-state energies, deformations, and low-lying excitations [1, 2, 85, 86].

2.1 Skyrme energy density functional

In this section we follow the notations employed in Refs. [1, 15] and define various local densities and the Skyrme EDF.

2.1.1 Local densities

Starting from a many-body wave function $|\Psi\rangle$ and a complete s.p. basis $\{\psi_1, \psi_2, \cdots \}$, one can define the one-body density matrix $\rho$ and pairing tensor $\kappa$ as

$$
\rho_{ij} = \langle \Psi | \hat{a}_j^\dagger \hat{a}_i | \Psi \rangle, \quad \kappa_{ij} = \langle \Psi | \hat{a}_j \hat{a}_i | \Psi \rangle,
$$

(2.2)

where $\hat{a}_i^\dagger$ and $\hat{a}_i$ create and annihilate, respectively, a nucleon in the s.p. state $\psi_i$. In the coordinate space, the non-local density matrix and pairing tensor can be similarly defined as

$$
\rho \left( r_s \tau, r'_s' \tau' \right) = \langle \Psi | \hat{a}_{r_s r'_s' \tau'}^\dagger \hat{a}_{r_s \tau} | \Psi \rangle, \quad \kappa \left( r_s \tau, r'_s' \tau' \right) = \langle \Psi | \hat{a}_{r_s' r_s \tau'} \hat{a}_{r_s \tau} | \Psi \rangle,
$$

(2.3)

where $\hat{a}_{r_s \tau}^\dagger$ and $\hat{a}_{r_s \tau}$ create and annihilate, respectively, a nucleon at point $r$ with spin $s = \pm \frac{1}{2}$ and isospin $\tau = \pm \frac{1}{2}$. 

8
In terms of spin and isospin components, the density matrix can be decomposed as

\[
\rho (r s \tau, r' s' \tau') = \frac{1}{4} \left[ \rho_{00} (r, r') \delta_{ss'} + s_{00} (r, r') \cdot \sigma_{ss'} \right] \delta_{\tau \tau'} + \frac{1}{4} \sum_{t_3=-1}^{+1} \left[ \rho_{1t_3} (r, r') \delta_{ss'} + s_{1t_3} (r, r') \cdot \sigma_{ss'} \right] (\bar{\tau}_{\tau'} t_3) \tag{2.4}
\]

where \( \sigma_{ss'} = (s | \sigma | s') \) and \( \bar{\tau}_{\tau'} = (\tau | \tilde{\tau} | \tau') \) are matrix elements of Pauli matrices in spin and isospin spaces, respectively. Then we have

\[
\rho_{00} (r, r') = \sum_{s \tau} \rho (r s \tau, r' s \tau), \tag{2.5a}
\]

\[
\rho_{1t_3} (r, r') = \sum_{s \tau \tau'} \rho (r s \tau, r' s' \tau') (\bar{\tau}_{\tau'} t_3), \tag{2.5b}
\]

\[
s_{00} (r, r') = \sum_{s s' \tau} \rho (r s \tau, r' s' \tau) \sigma_{s s'}, \tag{2.5c}
\]

\[
s_{1t_3} (r, r') = \sum_{s s' \tau \tau'} \rho (r s \tau, r' s' \tau') \sigma_{s s'} (\bar{\tau}_{\tau'} t_3). \tag{2.5d}
\]

Starting from the non-local density matrix, one can define following local densities and currents.

1. Time-even densities:

\[
\rho_{tt_3}(r) = \rho_{tt_3}(r, r), \tag{2.6a}
\]

\[
\tau_{tt_3}(r) = \nabla \cdot \nabla' \rho_{tt_3} (r, r') \bigg|_{r=r'}, \tag{2.6b}
\]

\[
\mathcal{J}_{tt_3}(r) = \frac{i}{2} \left( \nabla' - \nabla \right) \otimes s_{tt_3} (r, r') \bigg|_{r=r'}. \tag{2.6c}
\]

These are the particle density, kinetic density, and spin-current tensor, respectively. They do not change sign under the time-reversal operation.
2. Time-odd densities:

\[ s_{tt3}(r) = s_{tt3}(r, r), \quad (2.7a) \]
\[ j_{tt3}(r) = \frac{i}{2} \left( \nabla' - \nabla \right) \rho_{tt3}(r, r') \bigg|_{r=r'}, \quad (2.7b) \]
\[ T_{tt3}(r) = \nabla \cdot \nabla' s_{tt3}(r, r') \bigg|_{r=r'}, \quad (2.7c) \]
\[ F_{tt3}(r) = \frac{1}{2} \left( \nabla \otimes \nabla' + \nabla' \otimes \nabla \right) \cdot s_{tt3}(r, r') \bigg|_{r=r'}. \quad (2.7d) \]

These are the spin density, current, spin-kinetic density, and tensor-kinetic density, respectively. They change sign under the time-reversal operation.

For a stationary state of an even-even nuclei, all the time-odd densities vanish due to the time-reversal symmetry. As for the spin-current tensor \( \mathbb{J} \), only its anti-symmetric part is usually considered in the Skyrme EDF, which can be written as the spin-orbit current \( J_{tt3} = \sum_{ijk} \epsilon_{ijk} e_i (e_j \cdot \mathbb{J} \cdot e_k) \), where \( e_i \) is the unit vector in the \( i \) direction and \( \epsilon_{ijk} \) is the Levi-Civita symbol.

Usually the proton-neutron mixing is not considered, which means \( \tau = \tau' \) in Eq. (2.3) and \( t_3 = 0 \) in Eqs. (2.4) \( \sim \) (2.7). In this case we can define density matrices separately for neutrons and protons as

\[ \rho_q(r,s,r',s') = \rho(r s \tau_q, r' s' \tau_q), \quad (2.8) \]

where \( q \in \{ n, p \} \) stands for neutrons or protons and \( \tau_q = \pm \frac{1}{2} \) for \( q = n \) or \( p \). Then the local densities of neutrons and protons are given by

\[ \varrho_{00} = \varrho_n + \varrho_p, \quad \varrho_{10} = \varrho_n - \varrho_p. \quad (2.9) \]
where $\varrho = \rho, \tau, J, s, j, T, F$.

As for the pairing channel, instead of the pairing tensor $\kappa$, it is more convenient to formulate the EDF in terms of the pairing density matrix

$$\tilde{\rho} (r s \tau, r' s' \tau') = -2s' \kappa (r s \tau, r' s' \tau'), \quad \tilde{\rho} (r s \tau, r' s' \tau') = 4s' \tau' \kappa (r s \tau, r' s' \tau') , \quad (2.10)$$

where $\tilde{\rho}$ is usually adopted when there is no proton-neutron mixing, while $\bar{\rho}$ is more convenient to use in the isospin representation.

One can decompose $\tilde{\rho} (r s \tau, r' s' \tau')$ in the same way as Eq. (2.4) and then define various local quantities following Eqs. (2.5) $\sim$ (2.7). But in this dissertation we are only concerned about the time-even pairing density

$$\hat{\rho}_{1 t_3} (r) = \sum_{s t \tau'} \tilde{\rho} (r s \tau, r s' \tau') (\tilde{\tau}_{s' t})_{t_3} , \quad (2.11)$$

and time-odd pairing spin density

$$\tilde{s}_{00} (r) = \sum_{s s' \tau} \tilde{\rho} (r s \tau, r s' \tau) \sigma_{s' s} . \quad (2.12)$$

In the case of no proton-neutron mixing, it is more convenient to use the “tilde” density $\tilde{\rho} (r s \tau, r' s' \tau')$ and define local neutron and proton pairing densities separately:

$$\tilde{\rho}_q (r) = \sum_{s s'} \tilde{\rho}_q (r s \tau_q, r' s' \tau_q) , \quad q \in \{n, p\} . \quad (2.13)$$

While the particle density $\rho_q (r)$ gives the probability of finding a nucleon at position $r$, the pairing density $\tilde{\rho}_q (r)$ describes the enhancement of the probability of finding a pair of
nucleons with opposite spins due to the pairing correlation.

### 2.1.2 Skyrme EDF formalism

Within the Skyrme EDF framework, the total energy of a nucleus can be written as an integral over the whole space:

\[
E = \int dr \mathcal{H}(r) = \int dr \left[ \mathcal{H}_{\text{kin}}(r) + \mathcal{H}_{\text{Sk}}(r) + \mathcal{H}_{\text{pair}}(r) + \mathcal{H}_{\text{Coul}}(r) + \mathcal{H}_{\text{cm}}(r) \right], \tag{2.14}
\]

where \( \mathcal{H}(r) \) is the energy density (ED) that consists of various terms that are discussed in the following.

The kinetic ED is

\[
\mathcal{H}_{\text{kin}}(r) = \sum_{q \in \{n,p\}} \frac{\hbar^2}{2m_q} \sigma_q(r). \tag{2.15}
\]

Some parametrizations assume \( m_n = m_p = m \) and the kinetic ED then becomes \( \frac{\hbar^2}{2m} \sigma_0 \).

The Skyrme interaction ED (in the particle-hole channel) is

\[
\mathcal{H}_{\text{Sk}}(r) = \sum_{t=0}^{1} \sum_{t_3=-t}^{t} \mathcal{H}_{tt_{3}}^{(\text{even})}(r) + \mathcal{H}_{tt_{3}}^{(\text{odd})}(r), \tag{2.16}
\]

where

\[
\mathcal{H}_{tt_{3}}^{(\text{even})}(r) = C_t \rho_{tt_{3}}^2 + C_t^\Delta \rho_{tt_{3}} \nabla^2 \rho_{tt_{3}} + C_t^\tau \rho_{tt_{3}} \tau_{tt_{3}} + C_t^J \nabla^2 J_{tt_{3}} + C_t \nabla J_{tt_{3}} \cdot \mathbf{J}_{tt_{3}} \tag{2.17}
\]
is bilinear in time-even local densities, and

\[
H_{it}^{(\text{odd})} (r) = C_t s_{tt3}^2 + C_t^s s_{tt3} \cdot \nabla^2 s_{tt3} + C_t^j j_{tt3}^2 + C_t^T s_{tt3} \cdot T_{tt3} \\
+ C_t^{\nabla j} s_{tt3} \cdot \nabla \times j_{tt3} + C_t^F s_{tt3} \cdot F_{tt3} + C_t^{\nabla s} \left( \nabla \cdot s_{tt3} \right)^2
\] (2.18)

is bilinear in time-odd local densities. All the coupling constants \( C \) can be density-dependent, but in most Skyrme parameterizations only \( C_t^\rho \) and \( C_t^s \) depend on the isoscalar particle density \( \rho_{00} \) as

\[
C_t [\rho_{00}] = C_t[0] + (C_t [\rho_c] - C_t[0]) \left( \frac{\rho_{00}}{\rho_c} \right)^\gamma,
\] (2.19)

where \( \rho_c \approx 0.16 \text{ fm}^{-3} \) is the nuclear saturation density. The Skyrme interaction ED can be either derived from the Hartree-Fock calculation with an effective zero-range momentum-dependent two-body nuclear force proposed by Skyrme [87, 88], or from the density-matrix expansion without reference to an effective force [89, 90]. The first option results in strong dependencies among coupling constants \( C \), while the latter offers more degrees of freedom.

Terms involving \( C_t^J, C_t^T, C_t^F \) and \( C_t^{\nabla s} \) in Eqs. (2.17) and (2.18) are known as tensor terms, as they are related to the local two-body tensor interaction [87, 88, 91, 15]. There are a number of publications discussing the effects of tensor terms; see, e.g., Refs. [92, 93, 94] for systematic discussions.

The pairing ED (in the particle-particle channel) is usually based on a density-dependent \( \delta \) force. When the isospin symmetry is preserved, it can be written as

\[
\mathcal{H}_{\text{pair}} (r) = \frac{1}{8} \left[ 1 - \frac{\rho_{00}(r)}{\rho_{\text{ref}}} \right] \left( V_0 |\tilde{s}_{00}|^2 + V_1 \sum_{t_3=-1}^{1} |\tilde{\rho}_{1t3}|^2 \right),
\] (2.20)

where \( V_0 \) and \( V_1 \) are isoscalar and isovector pairing strengths, respectively. When the proton-
neutron mixing is absent, we can break the isospin symmetry by assigning different pairing strengths for neutrons and protons:

\[
\mathcal{H}_{\text{pair}}(r) = \frac{1}{4} \sum_{q \in \{n,p\}} V_q \left[ 1 - \frac{\rho_{00}(r)}{\rho_{\text{ref}}} \right] |\tilde{\rho}_q|^2. \tag{2.21}
\]

One can adjust the reference density \(\rho_{\text{ref}}\) for different types of density dependencies [95, 96]. A pure contact interaction (volume pairing) corresponds to \(\rho_{\text{ref}} \to \infty\); a value around the nuclear-matter saturation density \(\rho_{\text{ref}} = \rho_c\) leads to pairing around the nuclear surface (surface pairing); any value in between delivers a mixed-pairing prescription. A widely used value is \(\rho_{\text{ref}} = 0.32 \text{ fm}^{-3} = 2\rho_c\).

The Coulomb term is

\[
\mathcal{H}_{\text{Coul}}(r) = \frac{e^2}{2} \int dr \frac{\rho_p(r)\rho_p(r')}{|r - r'|} - \frac{3e^2}{4} \left( \frac{3}{\pi} \right)^{\frac{1}{3}} \frac{4}{\rho_p^4}(r), \tag{2.22}
\]

where the second term (exchange term) is given by the Slater approximation.

As for the center-of-mass (c.m.) correction \(\mathcal{H}_{\text{cm}}(r)\), there are several recipes and only those employed in this dissertation are discussed. In Skyrme parameterizations SkM* [97] and SLy4 [98], the c.m. correction is introduced by renormalizing the mass of nucleons: \(\frac{1}{m} \to \frac{1}{m} \left( 1 - \frac{1}{A} \right)\), where \(A\) is the number of nucleons in a nucleus. In UNEDF1-HFB [99], a parameterization designed for the study of fission, however, no c.m. correction is added, because the c.m. correction is not additive in the particle number and causes problems in fission calculations [100].

There are a number of model parameters in the Skyrme EDF, which should be adjusted to reproduce selected experimental data. Thus, there exist various Skyrme parameterizations
that are fitted to different data sets with probably different assumptions. The fit data and underlying assumptions are based on the selection of nuclear properties that researchers want to describe well. For example, the energies of fission isomers are included in the fits of UNEDF1 [100] and UNEDF1-HFB [99] so that the nuclear fission can be well described by these two parametrizations.

2.2 Hartree-Fock-Bogoliubov method

2.2.1 HFB equations in the quasiparticle basis

Within the nuclear-DFT framework, the ground state of an even-even nucleus is obtained through the variational principle. We choose the product state as the trial wave function and minimize the total energy under various constraints, which leads to the self-consistent HFB equation [1, 2, 3].

A product HFB state $|\Phi\rangle$ is a vacuum with respect to quasiparticles:

$$\hat{\beta}_k |\Phi\rangle = 0 \text{ for all } k = 1, 2, 3, \cdots ,$$

(2.23)

where the creation and annihilation operators of quasiparticles are defined by the Bogoliubov transformation:

$$\hat{\beta}_k^\dagger = \sum_l \left( U_{lk} \hat{a}_l^\dagger + V_{lk} \hat{a}_l \right), \quad \hat{\beta}_k = \sum_l \left( U_{lk}^* \hat{a}_l^\dagger + V_{lk}^* \hat{a}_l \right).$$

(2.24)
This transformation can be written in a compact manner as

\[
\begin{pmatrix}
\hat{\beta} \\
\hat{\beta}^\dagger
\end{pmatrix} = \mathcal{W}^\dagger
\begin{pmatrix}
\hat{a} \\
\hat{a}^\dagger
\end{pmatrix} = \begin{pmatrix}
U^\dagger & V^\dagger \\
V^T & U^T
\end{pmatrix}
\begin{pmatrix}
\hat{a} \\
\hat{a}^\dagger
\end{pmatrix}.
\]

The matrix \( \mathcal{W} \) must be unitary to ensure that \( \beta \) and \( \beta^\dagger \) obey the canonical anticommutation relations for fermionic operators. Combining Eqs. (2.24) and (2.2), we can express the density matrix and pairing tensor of the quasiparticle vacuum \( |\Phi\rangle \) as

\[
\rho = V^*V^T, \quad \kappa = V^*U^T.
\]  

(2.26)

In general, the product state \( |\Phi\rangle \) does not conserve neutron or proton numbers, so Lagrange multipliers need to be introduced to constrain average particle numbers. The total HFB Routhian is then written as

\[
R[\rho, \kappa, \kappa^*] = E[\rho, \kappa, \kappa^*] - \sum_{q \in \{n, p\}} \epsilon_F^{(q)} \left\langle \hat{N}_q \right\rangle,
\]

where \( \hat{N}_q \) is the particle number operator for neutrons \( (q = n) \) or protons \( (q = p) \), and the Lagrange multipliers \( \epsilon_F^{(q)} \) are also known as chemical potentials or Fermi energies. The minimization of \( R \) with regard to \( \rho \) and \( \kappa \) yields the HFB equation:

\[
H
\begin{pmatrix}
U_k \\
V_k
\end{pmatrix} = \begin{pmatrix}
h - \epsilon_F & \Delta \\
-\Delta^* & -h^* + \epsilon_F
\end{pmatrix}
\begin{pmatrix}
U_k \\
V_k
\end{pmatrix} = E_k
\begin{pmatrix}
U_k \\
V_k
\end{pmatrix},
\]

(2.28)

where \( h \) is called the HF mean field while \( \Delta \) is the pairing field. The eigenvector \( \begin{pmatrix} U_k \\ V_k \end{pmatrix} \) is the \( k \)-th column of \( \mathcal{W} \) and the corresponding eigenvalue \( E_k \) gives the quasiparticle energy.
$U_k$ and $V_k$ are often referred to as the upper and lower components of the quasiparticle wave function, respectively. The matrix elements of $H$ are given by

$$h_{ij} = \frac{\delta E[\rho, \kappa, \kappa^*]}{\delta \rho_{ji}} = h_{ji}^*, \quad \Delta_{ij} = \frac{\delta E[\rho, \kappa, \kappa^*]}{\delta \kappa_{ij}^*} = -\Delta_{ji}, \quad (\epsilon_F)_{ij} = \delta_{ij} \epsilon_F(q_i),$$

(2.29)

where $q_i \in \{n, p\}$ is the isospin of the s.p. state $i$ (we assume no proton-neutron mixing).

Explicit expressions of these matrix elements can be found in a number of publications, e.g., Refs. [1, 15, 86]. Generally speaking, the HFB Hamiltonian depends on the solution $U$ and $V$, so the HFB equation is non-linear and must be solved in a self-consistent iterative approach (e.g., iterative diagonalization or gradient descent). The HFB equation (2.28) is derived from the variation with respect to the density matrix $\rho$ and pairing tensor $\kappa$, but one can also replace $\kappa$ with the pairing density $\tilde{\rho}$ or $\hat{\rho}$ and obtain a similar equation that obviously yields the same physics [15, 70].

When pairing collapses in a closed-shell system, $\kappa = 0$, $\Delta = 0$, and Eq. (2.28) reduces to the HF equation that only involves the diagonalization of the HF mean field $h$. Meanwhile, the product state $|\Phi\rangle$ becomes a Slater determinant that has conserved neutron and proton numbers. Therefore, the HF approach can be treated as a special case of the HFB method. Based on the HF method, the pairing correlation can be included via the BCS approximation, and the HF+BCS approach is less computationally expensive than the full HFB method. However, the HF+BCS method can produce unphysical particle gas surrounding the nucleus and thus a full HFB scheme is preferred [72]. It should be noted that in both HF+BCS and HFB frameworks, the pairing-rearrangement term brought by the density dependence of the pairing functional is also included in the HF mean field $h$.

The HFB equation is numerically solved in a truncated model space, and it is well known
that pairing functionals based on the $\delta$ interaction, such as those defined in Eqs. (2.20, 2.21), diverge as the model space increases, so a cutoff has to be imposed on the pairing-active space [70, 72, 101]. This cutoff can be done in the quasiparticle space: The contribution of one quasiparticle state to all the densities (including the pairing density or pairing tensor) is multiplied by a function of the quasiparticle energy. A commonly used soft cutoff function is [102, 103]

$$w(e) = \frac{1}{1 + \exp\left(\frac{e - E_{\text{cut}}}{\Delta E_{\text{cut}}}\right)},$$

(2.30)

where $E_{\text{cut}}$ is the cutoff energy, $\Delta E_{\text{cut}}$ provides smearing around $E_0$, and $e$ is the reference (equivalent) s.p. energy

$$e_k = (1 - 2P_k) E_k + \epsilon^{(q_k)}_F,$$

(2.31)

where $E_k$ is the quasiparticle energy and $P_k$ represents the norm of the lower component $V_k$.

This reference spectrum is similar to the spectrum of the canonical basis; the definition of the canonical basis is to be presented in the next section. When $\Delta E_{\text{cut}} \to 0$, $w(e)$ becomes a hard cutoff at energy $E_{\text{cut}}$.

### 2.2.2 HFB equations in the canonical basis

Besides the formulation based on quasiparticles, the HFB theory can also be formulated in the s.p. basis of canonical states (or natural orbitals), in which the one-body density matrix $\rho$ is diagonal. The connection between the quasiparticle and canonical states is discussed in Refs. [72, 104]. On the one hand, once all the quasiparticle states are obtained, the canonical basis can be computed by diagonalizing the density matrix $\rho$, and the eigenvalues of $\rho$ represent the occupations of corresponding canonical states. On the other hand, one can directly solve the HFB problem in the canonical basis without any reference to the quasiparticle states.
The canonical-basis HFB formalism adopted in my work was first discussed in Ref. [76] and further developed in [77].

The canonical basis is given by a set of orthonormal s.p. wave functions $\psi_\alpha$ with occupation amplitudes $v_\alpha$:

$$\{\psi_\alpha, v_\alpha, \alpha = 1, \ldots, \Omega\}, \quad v_\alpha \in [0, 1],$$

(2.32)

where $v_\alpha^2$ and $\psi_\alpha$ are the eigenvalues and eigenvectors of the density matrix $\rho$, respectively, and $\Omega$ is the size of the active s.p. space. The non-occupation amplitude is defined as $u_\alpha = \sqrt{1 - v_\alpha^2}$. In the canonical basis, the HFB product state takes the BCS-like form

$$|\Phi\rangle = \prod_{\alpha > 0} \left( u_\alpha + v_\alpha \hat{a}_\alpha \hat{a}^\dagger_\alpha \right) |0\rangle$$

(2.33)

where $|0\rangle$ is the vacuum state, $\hat{a}_\alpha^\dagger$ generates a particle in the state $\psi_\alpha$, and $\alpha$ is the conjugate partner of $\alpha$ that corresponds to the same eigenvalue of $\rho$. In this section we focus on the stationary state of an even-even nucleus, so the partner $\alpha$ is assumed to be the time-reversed state of $\alpha$. Also, from Eq. (2.33) one can immediately see that the HFB product state in general does not have a good particle number.

The density matrix in the coordinate space can be expressed in terms of canonical wave functions:

$$\rho (r s \tau, r' s' \tau') = \sum_\alpha v_\alpha^2 \psi_\alpha (r, s, \tau) \psi^*_\alpha (r', s', \tau'),$$

(2.34)

from which one can calculate all the local densities involved in the particle-hole channel. Meanwhile, the pairing density matrix is

$$\tilde{\rho} (r s \tau, r' s' \tau') = \sum_\alpha u_\alpha v_\alpha (-2s') \psi_\alpha (r', -s', \tau') \psi^*_\alpha (r, s, \tau).$$

(2.35)
Without proton-neutron mixing, the local pairing density can be written as

\[ \tilde{\rho}_q(r) = \sum_{\alpha \in q} u_\alpha v_\alpha \sum_s (-2s) \psi_{\pi}(r, -s) \psi_\alpha(r, s), \tag{2.36} \]

which can be further simplified when the conjugate partner \( \bar{\alpha} \) is the time-reversed state of \( \alpha \):

\[ \tilde{\rho}_q(r) = \sum_{\alpha \in q} \sum_s u_\alpha v_\alpha |\psi_\alpha(r, s)|^2. \tag{2.37} \]

One should note that pairing cutoffs defined in the canonical and quasiparticle bases are not fully equivalent, and one way to achieve equivalence is to employ a pairing functional that does not require a cutoff (e.g., the momentum-dependent pairing functional proposed in Ref. [77]); but the effect brought by the difference should be minor when the cutoff is high enough.

The total HFB Routhian to minimize in the canonical-basis representation is

\[
R[\psi, \psi^*, v] = E\left[ \rho[\psi, \psi^*, v], \tilde{\rho}[\psi, \psi^*, v], \tilde{\rho}^\dagger[\psi, \psi^*, v] \right] \\
- \sum_{q \in \{n,p\}} \epsilon_F^{(q)} \sum_{\alpha \in q} v_\alpha^2 - \sum_{\alpha \beta} \lambda_{\alpha\beta} \left( \langle \psi_\beta | \psi_\alpha \rangle - \delta_{\alpha\beta} \right), \tag{2.38}
\]

where \( \epsilon_F^{(q)} \) is the Fermi energy of neutrons (\( q = n \)) or protons (\( q = p \)), and \( \lambda \) is the matrix of Lagrangian multipliers that guarantee the orthonormality of canonical states. The HFB energy \( E \) here is originally formulated as a functional of the density matrix \( \rho \), pairing density matrix \( \tilde{\rho} \) and its Hermitian conjugate \( \tilde{\rho}^\dagger \) (see Sec. 2.1), which then becomes a functional of canonical wave functions \( \psi \), their complex conjugates \( \psi^* \) and occupation amplitudes \( v \) through Eqs. (2.34, 2.35). Due to \( \langle \psi_\beta | \psi_\alpha \rangle = \langle \psi_\alpha | \psi_\beta \rangle^* \), the quantities we need to constrain for orthonormality constitute a Hermitian matrix, and thus the matrix \( \lambda \) should also be
Hermitian so that the number of independent Lagrange multipliers in it coincides with the number of independent constraints.

The variation of $R$ with respect to $\psi^*_\alpha$ yields the mean-field equation

$$0 = \frac{\delta R}{\delta \psi^*_\alpha} = \frac{\delta E}{\delta \psi^*_\alpha} - \sum_\beta \psi_\beta \lambda_{\beta\alpha} = \hat{H}_\alpha \psi_\alpha - \sum_\beta \psi_\beta \lambda_{\beta\alpha}, \quad (2.39)$$

where

$$\hat{H}_\alpha = v^2_\alpha \hat{h} + u_\alpha v_\alpha \hat{\tilde{h}}, \quad (2.40a)$$

$$\lambda_{\beta\alpha} = \frac{1}{2} \langle \psi_\beta | \hat{H}_\alpha + \hat{H}_\beta | \psi_\alpha \rangle. \quad (2.40b)$$

There are three points worth discussing about the mean-field equation. First, $\frac{\delta E}{\delta \psi^*_\alpha}$ can be reduced to

$$\frac{\delta E}{\delta \psi^*_\alpha} = \frac{\delta E}{\delta \rho} \frac{\delta \rho}{\delta \psi^*_\alpha} + \frac{\delta E}{\delta \tilde{\rho}} \frac{\delta \tilde{\rho}}{\delta \psi^*_\alpha} = v^2_\alpha \hat{h} \psi_\alpha + u_\alpha v_\alpha \hat{\tilde{h}} \psi_\alpha, \quad (2.41)$$

which delivers the explicit expressions of the HF Hamiltonian $\hat{h}$ and the pairing field $\hat{\tilde{h}}$ in the same way as Eq. (2.29). Second, the generalized Hamiltonian $\hat{H}_\alpha$ is state-dependent and hence the full matrix $\lambda$ has to be taken into account to preserve orthonormality. In contrast, the HF calculation only requires Lagrange multipliers for normalization:

$$\hat{h} \psi_\alpha - \varepsilon_\alpha \psi_\alpha = 0, \quad \varepsilon_\alpha = \langle \psi_\alpha | \hat{h} | \psi_\alpha \rangle, \quad (2.42)$$

where all the s.p. states experience the same Hamiltonian $\hat{h}$, and $\varepsilon_\alpha$ is the s.p. HF energy. Third, the Hermiticity of the matrix $\lambda$ is enforced by explicit symmetrization in Eq. (2.40b).
The variation of $R$ with respect to $v_\alpha$ yields the gap equation:

$$0 = 2v_\alpha \left[ h_{\alpha\alpha} - \epsilon_F^{(q_\alpha)} \right] + \left( u_\alpha - \frac{v_\alpha^2}{u_\alpha} \right) \tilde{h}_{\alpha\alpha}, \quad (2.43)$$

where $h_{\alpha\alpha} = \langle \psi_\alpha | \hat{h} | \psi_\alpha \rangle$ is the canonical s.p. energy, $\tilde{h}_{\alpha\alpha} = \langle \psi_\alpha | \hat{\tilde{h}} | \psi_\alpha \rangle$ is the state-dependent pairing gap, and $q_\alpha \in \{n, p\}$ stands for the isospin of the state $\alpha$. The gap equation can be solved in a closed form:

$$\begin{pmatrix} v_\alpha \\ u_\alpha \end{pmatrix} = \frac{1}{\sqrt{2}} \begin{pmatrix} 1 & 1 \\ 2 & 2 \end{pmatrix} \sqrt{h_{\alpha\alpha} - \epsilon_F^{(q_\alpha)}} \left[ h_{\alpha\alpha} - \epsilon_F^{(q_\alpha)} \right]^2 + \tilde{h}_{\alpha\alpha}^2, \quad (2.44)$$

where the Fermi energy $\epsilon_F^{(q)}$ should be adjusted to fulfill the particle-number constraints

$$\sum_{\alpha \in n} v_\alpha^2 = N, \quad \sum_{\alpha \in p} v_\alpha^2 = Z. \quad (2.45)$$

One can note that only the diagonal elements of the HF Hamiltonian $\hat{h}$ and the pairing field $\hat{\tilde{h}}$ in the canonical basis enter the gap equation; therefore, no information about the non-diagonal elements is needed to determine the occupation amplitudes. It should also be noted that the HF+BCS approach solves a gap equation in the same form of Eq. (2.43), but it uses matrix elements computed in the HF basis instead.

The mean-field equation (2.39, 2.40) and gap equation (2.43) together constitute the self-consistent HFB equations in the canonical basis, which can be numerically solved by the gradient descent. The same cutoff scheme as discussed at the end of Sec. 2.2.1 can also be employed in the canonical space: The contribution of one canonical state to all the densities is multiplied by a function of the canonical s.p. energy $\epsilon_\alpha \equiv h_{\alpha\alpha}$, and the cutoff function
can still take the form of Eq. (2.30).

**2.2.3 Constrained calculations**

In previous sections, the HFB Routhian is minimized under particle-number constraints. Other quantities, such as multipole moments, angular momenta, and particle-number fluctuations, can also be constrained in a similar manner [3]. The HFB energy obtained under these constraints becomes a function of the expectation values of these observables, and this function defines the potential-energy curve (one constraint) or surface (multiple constraints) of the system. The constrained HFB calculation can be employed for many problems. For instance, deformation-constrained calculations can help us locate the global minimum that corresponds to the ground state, while an unconstrained calculation can be easily stuck in a local minimum. The potential-energy surface obtained from constrained calculations is also a useful tool to study large-amplitude collective motions.

Nuclear deformations are usually extracted from the multipole moments, which are defined in the spherical coordinate system as

\[ q_{\lambda \mu} = \int d\mathbf{r} \rho(\mathbf{r}) r^{\lambda} Y_{\lambda \mu}(\Omega), \tag{2.46} \]

where \( \rho(\mathbf{r}) \) can be the neutron, proton or total (isoscalar) particle density, which gives the neutron, proton or total multipole moment, respectively. Triaxial components (\( \mu \neq 0 \)) vanish when the axial symmetry is preserved. The total isoscalar dipole moment (\( \lambda = 1 \)) is related to the shift of the center of mass, which must be constrained at zero when the parity symmetry is violated so that the center of mass is fixed at the origin. The quadrupole and
octupole moments can also be expressed in the Cartesian coordinate system:

\[ Q_{20} = \int d\mathbf{r} \rho(\mathbf{r}) \left( 2z^2 - x^2 - y^2 \right), \quad Q_{30} = \int d\mathbf{r} \rho(\mathbf{r}) \left[ z \left( 2z^2 - 3x^2 - 3y^2 \right) \right]. \tag{2.47} \]

The dimensionless deformation parameters are then

\[ \beta_2 = Q_{20} / \left( \sqrt{\frac{16\pi}{5}} \frac{3}{4\pi} AR_0^2 \right), \quad \beta_3 = Q_{30} / \left( \sqrt{\frac{16\pi}{7}} \frac{3}{4\pi} AR_0^3 \right), \tag{2.48} \]

where \( R_0 = 1.2A^{1/3} \) fm is the semi-empirical expression for the nuclear radius. It is more convenient to use dimensionless \( \beta \) values for the comparison of deformations among different nuclei.

There are several methods for constraining multipole moments. One approach is adding a quadratic penalty term to the HFB Routhian

\[ R' = R - c_{\lambda\mu} \left( q_{\lambda\mu} - \bar{q}_{\lambda\mu} \right)^2, \tag{2.49} \]

where \( \bar{q}_{\lambda\mu} \) is the desired value of the multipole moment and the Lagrange multiplier \( c_{\lambda\mu} \) should be large enough to push the minimum to the point of \( q_{\lambda\mu} = \bar{q}_{\lambda\mu} \). Another choice is adding a linear constraint \[105\]

\[ R' = R - c_{\lambda\mu} \left( q_{\lambda\mu} - \bar{q}_{\lambda\mu} \right), \tag{2.50} \]

where the Lagrange multiplier \( c_{\lambda\mu} \) is adjusted based on the quasiparticle random-phase approximation (QRPA). One can also choose the augmented Lagrangian method that combines the quadratic penalty and linear constraint \[106\].
The constraint on the angular-momentum expectation value appears in the cranking calculation for the description of nuclear rotation, and the same cranking formulation can be derived through the introduction of a rotating intrinsic frame [3, 33, 35, 107]. Here we assume that the system rotates around the \( y \) axis, which is not the symmetry axis of the system; then a linear constraint term is added to the Routhian

\[
R' = R - \omega \left( \langle \hat{J}_y \rangle - \bar{J}_y \right),
\] (2.51)

and correspondingly the cranking term \( -\omega \hat{J}_y \) appears in the HF Hamiltonian \( \hat{h} \). In the description of rotation, we usually specify the value of the Lagrange multiplier \( \omega \) instead of the angular momentum \( \bar{J}_y \), because \( \omega \) can be interpreted as the angular velocity of the system.

### 2.2.4 Numerical solvers

There are numerous HF and HFB solvers developed by researchers within the nuclear-DFT framework, and Ref. [75] provides a comprehensive table summarizing some widely employed solvers. In this section I briefly discuss the main features of solvers used in this dissertation. It should be noted that all the solvers discussed below assume no proton-neutron mixing.

The following two codes solve the HFB equation in the quasiparticle basis (Sec. 2.2.1) via direct diagonalization.

- HFBTHO [108, 109, 110, 111] solves the HFB problem with axial and time-reversal symmetries, and one can select whether the parity symmetry is imposed. The HFB Hamiltonian is constructed in the axially symmetric harmonic-oscillator (HO) or transformed HO basis. The HO basis is specified by the number of shells \( N_{\text{HO}} \) and axial
deformation $\beta_2$, and Gaussian quadratures are utilized to calculate integrals in the coordinate space. In HFBTHO, linear constraints (2.50) are added for deformation-constrained calculations. It also provides the kickoff mode, where first 10 iterations are carried out under deformation constraints specified in the input, and then the constraints are released in the following iterations. The kickoff mode helps the program better locate the global minimum without exploring the whole PES.

- HFODD [112, 113, 114, 115, 116, 117, 118, 119, 120] expands quasiparticle wave functions in the three-dimensional (3D) Cartesian deformed HO basis. The oscillator length and numbers of HO quanta in three directions can be varied independently. There is no symmetry restriction, but one can choose to impose the time-reversal symmetry and point-group symmetries like parity or signature. HFODD can perform the cranking calculation for the description of nuclear rotation. As for deformation-constrained calculations, one can choose to add quadratic penalties (2.49) or to use the augmented Lagrange method.

The following three programs solve the HFB equation in the canonical basis (Sec. 2.2.2) using the gradient-descent method.

- HFBFFT [121] solves the HFB equation in the 3D coordinate representation, and the fast Fourier transform (FFT) is employed for numerical differentiation. It assumes the time-reversal symmetry, but no spatial symmetry is imposed. It is based on Sky3D [81, 122, 123], a solver that can perform static HF+BCS and time-dependent HF calculations. The numerical details of HFBFFT are discussed in Chapter 6.

- Sky2D and Sky1D [124] also work in the coordinate space, but Sky2D imposes the axial symmetry while Sky1D the spherical symmetry. Sky2D solves the HFB problem
in the cylindrical coordinate system while Sky1D in the spherical coordinate system.

As for numerical differentiation, Sky2D employs the FFT technique while Sky1D uses the five-point finite difference formula. When the reflection symmetry is imposed in Sky2D, only grid points with \( z > 0 \) are taken into account. It is worth mentioning that Sky2D is based on the HF+BCS solver SkyAx [125].

2.3 Charge-changing finite amplitude method

2.3.1 FAM equations

The HFB theory discussed in Sec. 2.2 is applicable for the ground-state calculation. As for low-lying excited states and giant resonances, the QRPA is often the tool of choice within the nuclear-DFT framework [1, 3, 2]. The problem is that solving the QRPA equation through direct construction and diagonalization of the QRPA matrix is too computationally demanding, especially for deformed systems with no spherical symmetry. The FAM [63, 64], however, provides an efficient scheme for the solution of the QRPA without explicitly constructing a huge matrix.

The FAM is based on the small amplitude limit of the time-dependent HFB (TDHFB) equation. It is assumed the nuclear system evolves with time and stays as a quasiparticle vacuum (HFB product state) all the time. The corresponding quasiparticle operators are thus time-dependent:

\[
\hat{\beta}_k^\dagger(t) = \sum_l \left\{ U_{lk}(t)\hat{a}_l^\dagger + V_{lk}(t)\hat{a}_l \right\}, \quad \hat{\beta}_k(t) = \sum_l \left\{ U_{lk}^*(t)\hat{a}_l + V_{lk}^*(t)\hat{a}_l^\dagger \right\}.
\] (2.52)

The time evolution of the quasiparticle annihilation operator under a time-dependent exter-
nal field $\hat{F}(t)$ is given by the TDHFB equation:

$$
\frac{i}{\partial t} \hat{\beta}_k(t) = \left[ \hat{H}(t) + \hat{F}(t), \hat{\beta}_k(t) \right],
$$

(2.53)

where $\hat{H}(t)$ is the TDHFB Hamiltonian

$$
\hat{H}(t) = \frac{1}{2} \left( \hat{a}^\dagger \hat{a} + \hat{\beta}_k \right) H(t) \left( \begin{array}{c} \hat{a} \\ \hat{\beta}_k \end{array} \right) = \frac{1}{2} \left( \hat{a}^\dagger \hat{a} \right) \begin{pmatrix} h(t) - \epsilon_F & \Delta(t) \\ -\Delta^*(t) & -h^*(t) + \epsilon_F \end{pmatrix} \left( \begin{array}{c} \hat{a} \\ \hat{\beta}_k \end{array} \right),
$$

(2.54)

where normal ordering is assumed, and the matrix $H(t)$ has the same form as the static HFB matrix given in Eq. (2.28). The matrix elements of $H(t)$ are obtained in the same way as Eq. (2.29), but the mean fields $h(t)$ and $\Delta(t)$ are time-dependent in the TDHFB equation because the densities (and underlying solutions $U$ and $V$) are time-dependent.

In the small amplitude limit, the system is perturbed by a weak external field $\hat{F}(t)$ with a fixed frequency $\omega$:

$$
\hat{F}(t) = \eta \left( \hat{F} e^{-i\omega t} + \hat{F}^\dagger e^{i\omega t} \right), \quad \hat{F} = \frac{1}{2} \sum_{kl} \left( F_{kl}^{20} \hat{A}_{kl} + F_{kl}^{02} \hat{B}_{kl} \right) + \sum_{kl} F_{kl}^{11} \hat{B}_{kl},
$$

(2.55)

where $\hat{A}_{kl} = \hat{\beta}_k^\dagger \hat{\beta}_l^\dagger$, $\hat{B}_{kl} \equiv \hat{\beta}_k^\dagger \hat{\beta}_l$, and $\eta$ is small. Because of the canonical anticommutation relations $\{\hat{\beta}_k, \hat{\beta}_l\} = \{\hat{\beta}_k^\dagger, \hat{\beta}_l^\dagger\} = 0$, $F^{20}$ and $F^{02}$ are chosen to be anti-symmetric. The oscillation of $\hat{F}(t)$ induces the oscillations of the density matrix and pairing tensor:

$$
\rho(t) = \rho_0 + \eta \delta \rho(t) = \rho_0 + \eta \left\{ \delta \rho(\omega) e^{-i\omega t} + \delta \rho^\dagger(\omega) e^{i\omega t} \right\},
$$

(2.56a)

$$
\kappa(t) = \kappa_0 + \eta \delta \kappa(t) = \kappa_0 + \eta \left\{ \delta \kappa^+(\omega) e^{-i\omega t} + \delta \kappa^-(\omega) e^{i\omega t} \right\},
$$

(2.56b)
where \( \rho_0 \) and \( \kappa_0 \) are, respectively, the density matrix and pairing tensor of the stationary HFB state that the system oscillates around. The Hamiltonian is hence also oscillating around the stationary HFB Hamiltonian \( H_0 \):

\[
\hat{H}(t) = H_0 + \delta \hat{H}(t) = H_0 + \eta \left\{ \delta \hat{H}(t) e^{-i\omega t} + \delta \hat{H}^\dagger(t) e^{i\omega t} \right\},
\]

\[
\delta \hat{H}(\omega) = \frac{1}{2} \sum_{kl} \left\{ \delta H_{20}^{kl}(\omega) \hat{A}_{kl}^\dagger + \delta H_{02}^{kl}(\omega) \hat{A}_{kl} \right\} + \sum_{kl} \delta H_{11}^{kl}(\omega) \hat{B}_{kl},
\]

where \( \delta H_{20} \) and \( \delta H_{02} \) are anti-symmetric. Moreover, the time-dependent quasiparticle annihilation operator can be decomposed in a similar way:

\[
\hat{\beta}_k(t) = \left\{ \hat{\beta}_k + \delta \hat{\beta}_k(t) \right\} e^{iE_k t}, \quad \delta \hat{\beta}_k(t) = \eta \sum_l \hat{\beta}_l^\dagger \left\{ X_{lk}(\omega) e^{-i\omega t} + Y_{lk}^\ast(\omega) e^{i\omega t} \right\}
\]

where \( E_k \) is the quasiparticle energy of the static HFB Hamiltonian \( H_0 \), and \( \delta \hat{\beta}_k(t) \) is expanded only in terms of quasiparticle creation operators so that the anticommutation relation \( \{ \hat{\beta}_k(t), \hat{\beta}_l^\dagger(t) \} = \delta_{kl} \) is satisfied up to the linear order of \( \eta \). The FAM amplitudes \( X \) and \( Y \) must be anti-symmetric to fulfill the anticommutation relation \( \{ \hat{\beta}_k(t), \hat{\beta}_l(t) \} = 0 \).

With all the definitions given above, the TDHFB equation up to the linear order of \( \eta \) is

\[
i \frac{\partial \delta \hat{\beta}_k(t)}{\partial t} = E_k \delta \hat{\beta}_k(t) + \left[ \hat{H}_0, \delta \hat{\beta}_k(t) \right] + \left[ \delta \hat{H}(t) + \hat{F}(t), \delta \hat{\beta}_k \right],
\]

which yields the FAM equations

\[
(E_k + E_l - \omega) X_{kl}(\omega) + \delta H_{20}^{kl}(\omega) = -F_{20}^{kl}, \quad (E_k + E_l + \omega) Y_{kl}(\omega) + \delta H_{02}^{kl}(\omega) = -F_{02}^{kl}.
\]

It should be noted that \( F_{11} \) and \( \delta H_{11} \) are absent in Eq. (2.60) as they do not contribute in
the linear expansion. Thanks to the anti-symmetric property of $X$ and $Y$, only half of their matrix elements must be solved. Furthermore, the frequency $\omega$ is, in general, a complex number. Now the task is to calculate the induced fields $\delta H^{20}$ and $\delta H^{02}$. One can expand them in terms of amplitudes $X$ and $Y$, and obtain the linear response equation

$$
\begin{pmatrix}
A & B \\
B^* & A^*
\end{pmatrix} - \omega
\begin{pmatrix}
I & 0 \\
0 & -I
\end{pmatrix}
\begin{pmatrix}
X(\omega) \\
Y(\omega)
\end{pmatrix} = -
\begin{pmatrix}
F^{20} \\
F^{02}
\end{pmatrix},
$$

(2.61)

where \( \begin{pmatrix} A & B \\ B^* & A^* \end{pmatrix} \) is the QRPA matrix and $I$ is the identity matrix. Eq. (2.61) becomes the standard QRPA equation when the right-hand side is set to zero:

$$
\begin{pmatrix}
A & B \\
B^* & A^*
\end{pmatrix}
\begin{pmatrix}
\chi_n \\
\gamma_n
\end{pmatrix} = \Omega_n
\begin{pmatrix}
I & 0 \\
0 & -I
\end{pmatrix}
\begin{pmatrix}
\chi_n \\
\gamma_n
\end{pmatrix},
$$

(2.62)

where the eigenvalue $\Omega_n$ is the QRPA energy and the eigenvector \( \begin{pmatrix} \chi_n \\ \gamma_n \end{pmatrix} \) contains QRPA amplitudes (see Ref. [126] for the relation between the FAM amplitudes and QRPA eigenmodes).

The explicit construction of the QRPA matrix is often numerically infeasible, because the dimension of the QRPA matrix, which equals the number of two-quasiparticle excitations, is huge, especially for deformed nuclei. In the FAM, however, the induced fields are directly computed with no explicit evaluation of the QRPA matrix.

Based on Eq. (2.58), we can obtain time-dependent quasiparticle wave functions

$$
U_{lk}(t) = U_{lk}e^{-iE_k t} + n \sum_j \left( V_{lj}^* X_{jk} e^{i\omega t} + V_{lj}^* Y_{jk} e^{-i\omega t} \right) e^{-iE_k t},
$$

(2.63a)

$$
V_{lk}(t) = V_{lk}e^{-iE_k t} + n \sum_j \left( U_{lj}^* X_{jk} e^{i\omega t} + U_{lj}^* Y_{jk} e^{-i\omega t} \right) e^{-iE_k t},
$$

(2.63b)
where \((U, V)\) gives the quasiparticle wave functions of the stationary HFB solution that the system oscillates around. The density matrix \(\rho(t)\) and pairing tensor \(\kappa(t)\) in the s.p. basis are then calculated from \(U_k(t)\) and \(V_k(t)\) via Eq. (2.26). Based on \(\rho(t)\) and \(\kappa(t)\) one can construct the TDHFB Hamiltonian matrix (2.54) in the s.p. basis, which is connected with the induced fields \(\delta H^{20}\) and \(\delta H^{02}\) through the Bogoliubov transformation (2.24). In this procedure we only keep terms up to the first order of \(\eta\); the value of \(\eta\) should be small enough for linearity, but large enough to avoid reaching the numerical precision limit. Explicit expressions for the procedure are presented in Ref. [64]. One can also note that the time-reversal symmetry is broken in the FAM, so the Skyrme-EDF terms involving time-odd densities must be considered for the evaluation of the induced fields.

The general FAM formulation discussed above is applicable for any external transition operator \(\hat{F}\), but for a specific type of \(\hat{F}\) one can use its features to simplify the numerical procedure. As for beta-decay calculations, the external field is generated by the charge-changing operator that transforms a neutron to a proton or vice versa, and the charge-changing FAM is also called the proton-neutron FAM (PNFAM) [65]. In this work we concentrate on the beta-minus decay, which involves one-body transition operators that convert a neutron to a proton. Correspondingly, only proton and neutron quasiparticle states are connected by such operators, and only matrix elements between proton and neutron states are non-zero in the FAM amplitudes \(X\) and \(Y\), induced density \(\delta \rho\), induced pairing tensor \(\delta \kappa\), and induced fields \(\delta H^{02}\) and \(\delta H^{20}\). Therefore, although there is usually no proton-neutron mixing in the stationary HFB state the system oscillates around, such mixing is induced in the PNFAM. Here we only need to solve the proton-neutron part of FAM amplitudes while the neutron-proton part can be obtained via the anti-symmetric property \(X_{\nu\pi} = -X_{\pi\nu}, Y_{\nu\pi} = -Y_{\pi\nu}\).
2.3.2 FAM response function

The relation between the FAM amplitudes and transition matrix elements is revealed by connecting the small-amplitude TDHFB framework with the time-dependent perturbation theory [63]. In the first-order approximation with regard to the external perturbation \( \hat{F}(t) \), the system evolves as

\[
|\Psi(t)\rangle = |\Phi_0\rangle - i \sum_n e^{-i\Omega_n t} \int_{-\infty}^{t} dt' e^{i\Omega_n t'} |\phi_n\rangle \langle \phi_n | \hat{F}(t') |\Phi_0\rangle
\]

\[
= |\Phi_0\rangle + \sum_n |\phi_n\rangle \left( \frac{\eta \langle \phi_n | \hat{F} | \phi_0 \rangle}{\omega - \Omega_n + i\epsilon} e^{-i\omega t} - \frac{\eta^* \langle \phi_n | \hat{F}^\dagger | \phi_0 \rangle}{\omega + \Omega_n - i\epsilon} e^{i\omega t} \right),
\]

(2.64)

where \( |\Phi_0\rangle \) is the ground state while \( |\phi_n\rangle \) represents the \( n \)-th excited state whose excitation energy is \( \Omega_n \). Here the frequency \( \omega \) in \( \hat{F}(t) \) is replaced by \( \omega \pm i\epsilon \) so that the external field vanishes when \( t \to -\infty \). The expectation value of \( \hat{F}^\dagger \) is then

\[
\langle \Psi(t) | \hat{F}^\dagger | \Psi(t) \rangle = \langle \Phi_0 | \hat{F}^\dagger | \Phi_0 \rangle + \eta S(\hat{F}; \omega) e^{-i\omega t} + ...
\]

(2.65a)

\[
S(\hat{F}; \omega) = \sum_n \left( \frac{|\langle \phi_n | \hat{F} | \phi_0 \rangle|^2}{\omega - \Omega_n + i\epsilon} - \frac{|\langle \phi_n | \hat{F}^\dagger | \phi_0 \rangle|^2}{\omega + \Omega_n - i\epsilon} \right)
\]

(2.65b)

where \( S(F; \omega) \) is the response function. Taking the limit \( \epsilon \to 0 \), we have the transition strength distribution

\[
\frac{dB(\hat{F}; \omega)}{d\omega} \equiv \sum_n |\langle \phi_n | \hat{F} | \phi_0 \rangle|^2 \delta (\omega - \Omega_n) = \frac{1}{\pi} \text{Im} S(\hat{F}; \omega).
\]

(2.66)

For a complex frequency \( \omega = \omega_r + i\Gamma \) (\( \Gamma > 0 \)), we have

\[
- \frac{1}{\pi} \text{Im} S(\hat{F}; \omega_r + i\Gamma) = \frac{\Gamma}{\pi} \sum_n \left\{ \frac{|\langle \phi_n | \hat{F} | \phi_0 \rangle|^2}{(\omega_r - \Omega_n)^2 + \Gamma^2} - \frac{|\langle \phi_n | \hat{F}^\dagger | \phi_0 \rangle|^2}{(\omega_r + \Omega_n)^2 + \Gamma^2} \right\},
\]

(2.67)
which is the strength distribution smeared with a Lorentzian function of width $\Gamma$.

One can also evaluate the expectation value $\langle \Psi(t)|\hat{F}^\dagger|\Psi(t)\rangle$ with the TDHFB state in the small amplitude limit, and find the relation

$$S(\hat{F};\omega) = \frac{1}{2} \sum_{kl} \left\{ F_{kl}^{20*} X_{kl}(\omega) + F_{kl}^{02*} Y_{kl}(\omega) \right\}$$

$$= \sum_n \left( \frac{\langle \Phi_n|\hat{F}|\Phi_0\rangle^2}{\omega - \Omega_n} - \frac{\langle \Phi_n|\hat{F}^\dagger|\Phi_0\rangle^2}{\omega + \Omega_n} \right),$$

which connects the transition matrix elements with the FAM solution. Based on Eqs. (2.61, 2.62), Ref. [126] shows that the excitation energy $\Omega_n$ equals the QRPA energy and the state $|\Phi_n\rangle$ is the QRPA state (including the QRPA ground state $|\Phi_0\rangle$). When the QRPA stability condition is met, all the energies $\Omega_n$ are real-valued and we have $S(\omega^*) = S^*(\omega)$.

It is also possible to construct a response function that includes the interference between two distinct transition operators [65]. Starting from $\langle \Psi(t)|\hat{G}^\dagger|\Psi(t)\rangle$, we obtain

$$\chi(\hat{F},\hat{G};\omega) = \frac{1}{2} \sum_{kl} \left\{ G_{kl}^{20*} X_{kl}(\hat{F};\omega) + G_{kl}^{02*} Y_{kl}(\hat{F};\omega) \right\}$$

$$= \sum_n \left( \frac{\langle \Phi_n|\hat{F}^\dagger|\Phi_0\rangle \langle \Phi_0|\hat{G}^\dagger|\Phi_n\rangle}{\omega - \Omega_n} - \frac{\langle \Phi_0|\hat{F}|\Phi_n\rangle \langle \Phi_n|\hat{G}^\dagger|\Phi_0\rangle}{\omega + \Omega_n} \right),$$

where $G^{20}$ and $G^{02}$ are defined in the same manner as $F^{20}$ and $F^{02}$ in Eq. (2.55).

### 2.3.3 Beta-decay rate

As shown in Ref. [65], the beta-decay rate can be calculated via the contour integration of PNFAM response functions, which eliminates the necessity to extract individual QRPA modes from the response functions. Here we take the Fermi transition as an example to illustrate the procedure. Equations for the rates of allowed (Fermi and Gamow-Teller) and
first-forbidden transitions can all be found in Ref. [65].

The Fermi transition rate is proportional to the sum of individual transition strengths \( B_i(\hat{F}) \) from the ground state of the parent nucleus \(|\Phi_0\rangle\) to all the energetically allowed states \(|\Phi_i\rangle\) of the daughter nucleus, weighted by a phase-space factor:

\[
\lambda_{\hat{F}} = \frac{\ln 2}{\kappa} \sum_n f(E_n) B_n(\hat{F}),
\]

(2.70)

where \( \kappa = (6147.0 \pm 2.4) \) s, \( \hat{F} = \tau_- \) is the Fermi transition operator (isospin lowering operator) for the \( \beta^- \) decay, \( E_n \) is the energy released in the transition, and \( f(E_n) \) is the phase-space factor derived from the final-state lepton kinematics. The explicit expression of \( f \) can be found in Refs. [65, 127], and the released energy \( E_n \) is computed via [58]

\[
E_n = \omega_{\text{max}} - \Omega_n = \epsilon^{(n)}_F - \epsilon^{(p)}_F + \Delta M_{n-H} - \Omega_n,
\]

(2.71)

where \( \Omega_n \) is the QRPA energy, and \( \Delta M_{n-H} = 0.78227 \) MeV is the mass difference between the neutron and hydrogen atom. Here \( \epsilon^{(n)}_F \) and \( \epsilon^{(p)}_F \) represent neutron and proton Fermi energies of the HFB ground state of the parent nucleus, respectively. Based on Eq. (2.68), the transition strength can be expressed as the residue of the response function, and one can thus perform a contour integration in the complex plane of \( \omega \) to evaluate the rate:

\[
\lambda_{\hat{F}} = \frac{\ln 2}{\kappa} \sum_n g(\Omega_n) B_n(\hat{F}) = \frac{\ln 2}{\kappa} \sum_n g(\Omega_n) \text{Res} \left[ S(\hat{F}), \Omega_n \right]
\]

\[
= \frac{\ln 2}{\kappa} \sum_n \text{Res} \left[ \tilde{g}S(\hat{F}), \Omega_n \right] = \frac{\ln 2}{\kappa} \frac{1}{2\pi i} \oint_C d\omega \tilde{g}(\omega) S(\hat{F};\omega),
\]

(2.72)

where \( g(\omega) = f(\omega_{\text{max}} - \omega) \), \( \tilde{g}(\omega) \) is the analytic continuation of \( g(\omega) \), and the contour \( C \) encloses all the poles entering the summation (2.70). Our choice is a circular contour that
crosses the real axis at origin and $\omega_{\text{max}}$:

$$\omega(\theta) = \frac{\omega_{\text{max}}}{2} \left( 1 + e^{i\theta} \right), \; \theta \in [-\pi, \pi]. \tag{2.73}$$

Thanks to the relation $S(\omega^*) = S^*(\omega)$, we only need to compute the integrand on the upper or lower semicircle.

The remaining problem is how the analytic continuation of the phase-space factor $f$ should be done numerically. We employ a polynomial or rational function to interpolate $f$ on the real axis; the interpolating function is analytic in the complex plane and can thus be used in the integrand. Because of the Runge’s phenomenon [128], a better choice is the Thiele’s interpolation formula [67, 129]. A rational function in the form of a continued fraction is utilized to interpolate $f$ on a grid between 0 and $\omega_{\text{max}}$:

$$f_{\text{interp}}(x) = f(x_1) + \frac{x - x_1}{\rho(x_1, x_2) + \frac{x - x_2}{\rho_2(x_1, x_2, x_3) - f(x_1) + \frac{x - x_3}{\rho_3(x_1, x_2, x_3, x_4) - \rho(x_1, x_2) + \cdots}}} \tag{2.74}$$

where $x_1, x_2, \cdots$ constitute the grid on which $f$ is evaluated and $\rho$ denotes the reciprocal difference. With this interpolation the integrand is smooth, and we can employ the Gauss-Legendre quadrature rule to calculate the integral.

The total beta-decay rate $\lambda$ is obtained by summing the rates of all the possible decay transitions. Currently we include allowed and first-forbidden transitions in our calculations. Once we have the total rate, the half life can be evaluated via $T_{1/2} = \frac{\ln 2}{\lambda}$. 

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2.3.4 Gamow-Teller resonance

Apart from the decay rate, the Gamow-Teller resonance (GTR) is another phenomenon worth investigating through the PNFAM formalism. Among all the final states that can be reached via GT transitions from the ground state of the parent nucleus, the GTR takes a large portion of the total GT transition strength, and it can be identified as a strong enhancement (peak) in the total GT transition strength distribution

$$\frac{-1}{\pi} \sum_{K=0, \pm 1} S(GT, K; \omega)$$

(2.75)

where $K$ is the angular momentum projection of the GT operator. For a spherical parent nucleus, the response function $S$ is independent of $K$ and we only need consider one $K$ value.

To locate the GTR, one can evaluate $S(GT, K; \omega)$ along a horizontal line close to the real axis in the $\omega$ plane, i.e., $\omega = \omega_r + i\Gamma$ where $\Gamma$ is a small positive constant; then the maximum of the smeared total GT strength (2.75) corresponds to the GTR. A direct scan along the line is computationally expensive, because a dense grid of $\omega_r$ with a tiny smearing width $\Gamma$ is required for high resolution but the PNFAM does not converge well when $\Gamma$ approaches zero. Besides, the range of $\omega_r$ must be large enough to ensure that the GTR is included in the scan. For efficient determination of the GTR, we turn to a sparser $\omega_r$ grid and a larger $\Gamma$ when the GTR is well separated from other states, and interpolate the response function between grid points to locate the maximum. The width $\Gamma$ here should be smaller than the grid spacing to avoid missing peaks. A good choice is to interpolate $1/S$ with the Thiele’s formula (2.74), because it produces a rational function with the degree of the numerator being either the same (odd number of grid points) or one more than (even number of grid points) the degree of the denominator, and $1/S$ has the latter property according to Eq.
Complex-conjugate points $\omega^* = \omega_r - i\Gamma$ are also added to the interpolation to ensure $S(\omega^*) = S^*(\omega)$.

The GTR energy presented in experimental works are usually the excitation energy with respect to the ground state of the daughter nucleus, while the $\omega_r$ value corresponding to the GTR peak is the QRPA energy relative to the ground state of the parent nucleus. As Ref. [130] points out, the QRPA energy needs to be converted to

$$E_x = E_{\text{QRPA}} - m(Z + 1, A) + m(Z, A) - \Delta M_{n-H} + \epsilon_F^{(p)} - \epsilon_F^{(n)},$$

(2.76)

for direct comparison with the experimental GTR energy. Here $m(Z, A)$ and $m(Z + 1, A)$ are the atomic masses of parent and daughter nuclei, respectively.

### 2.3.5 Numerical PNFAM solver

The development and test of the numerical PNFAM solver has been discussed in Ref. [65], while its applications can be found in Refs. [62, 66, 67]. In PNFAM, the calculation starts from a HFBTHO solution with the axial symmetry imposed. A Python package called PyNFAM [67] has been developed to manage HFBTHO and PNFAM calculations in a parallel manner with MPI, and to calculate observables like GTRs and decay rates based on PNFAM outputs. The parallelization in PyNFAM is straightforward: First, calculations of different nuclei can always be done in parallel; second, for one nucleus, HFBTHO runs with different deformation kickoffs are performed simultaneously, and PNFAM computations at distinct frequencies $\omega$ can also run at the same time.

In the PNFAM program, the performance has been significantly boosted ($\gtrsim 10 \times$ speedup) by replacing explicit loops with BLAS [131] matrix-matrix multiplication routines. The
Hamiltonian matrix element in the PNFAM is constructed via the Gaussian quadrature:

\[
h_{\alpha\beta} = \sum_{s,s'} \int d^3r \psi_\alpha^*(r, s) h(r, s, s') \psi_\beta(r, s') \approx \sum_{s,s',i} w_i \psi_\alpha^*(r_i, s) h(r_i, s, s') \psi_\beta(r_i, s'), \tag{2.77}
\]

where \( \psi(r, s) \) is the basis wave function at point \( r \) with spin \( s \), \( h(r, s, s') \) the mean-field matrix element in the coordinate space, and \( w_i \) the quadrature weight. We can construct matrices \( A^{(s)} \) and \( B^{(s)} \) such that \( A^{(s)}_{\alpha i} = \sqrt{w_i} \psi_\alpha^*(r_i, s) \), \( B^{(s)}_{j\beta} = \sum_{s'} h(r_i, s, s') \sqrt{w_i} \psi_\beta(r_i, s') \). Then the matrix element \( h_{\alpha\beta} \) can be expressed in the form of a matrix-matrix multiplication:

\[
h_{\alpha\beta} = \sum_s \sum_i A^{(s)}_{\alpha i} B^{(s)}_{i\beta} = \sum_s A^{(s)} B^{(s)},
\]

which can be efficiently computed by optimized BLAS libraries. A similar strategy can be used to accelerate the evaluation of local densities.

For instance, the particle density is given by

\[
\rho(r_i) = \sum_s \sum_{\alpha\beta} \psi_\alpha(r_i, s) \rho_{\alpha\beta} \psi_\beta^*(r_i, s), \tag{2.78}
\]

where \( \rho_{\alpha\beta} \) is the density matrix in the HO basis. We immediately see that \( \sum_\beta \rho_{\alpha\beta} \psi_\beta^*(r_i, s) \) can be computed by the matrix-matrix multiplication routine.
Chapter 3

Nucleon localization function in rotating nuclei

This chapter investigates the nucleon localization function (NLF) in rotating systems. Results of this work has been published in Ref. [132]. This chapter is organized as follows. Sec. 3.1 first discusses the definition and interpretation of the NLF, and then the details of cranked calculations are given in Sec. 3.2. Results are presented in Sec. 3.3, with a short summary given in Sec. 3.4.

3.1 Nucleon localization function

3.1.1 Definition

Let $\mu = x, y, z$ be the spin-quantization axis. Starting from $\rho_q (r s_\mu, r' s'_\mu)$, the non-local density of neutrons ($q = n$) or protons ($q = p$) defined in Eq. (2.8), one can construct the particle density, kinetic density and current of a specific spin $s_\mu = \pm \frac{1}{2}$ and isospin $q$ as

\begin{align}
\rho_{qs_\mu} (r) &= \rho_q (r s_\mu, r s_\mu) = \frac{1}{2} \rho_q (r) + \frac{1}{2} \sigma_\mu s_q (r) \cdot e_\mu, \\
\tau_{qs_\mu} (r) &= \nabla \cdot \nabla' \rho_q (r s_\mu, r' s'_\mu) \bigg|_{r = r'} = \frac{1}{2} \tau_q (r) + \frac{1}{2} \sigma_\mu T_q (r) \cdot e_\mu, \\
j_{qs_\mu} (r) &= \frac{i}{2} (\nabla' - \nabla) \rho_q (r s_\mu, r' s'_\mu) \bigg|_{r = r'} = \frac{1}{2} j_q (r) + \frac{1}{2} \sigma_\mu \sigma_q (r) \cdot e_\mu,
\end{align}
where $\sigma_\mu = 2s_\mu = \pm 1$, $e_\mu$ is the unit vector in the $\mu$ direction, and $\rho_q$, $s_q$, $\tau_q$, $T_q$, $j_q$ and $J_q$ are local densities defined in Eq. (2.9). In a rotationally-invariant and spin-unpolarized system, densities defined above are independent of the choice of the quantization axis $\mu$, but in a deformed and rotating nucleus one has to explicitly specify $\mu$.

For a HF product state $|\Psi\rangle$, the definition of the NLF starts from the probability of finding two nucleons of a given isospin $q$ and spin $s_\mu$ at spatial locations $r$ and $r'$:

$$P_{qs\mu}(r, r') = \langle \Psi | a_r^{s\mu q} a_r^{s\mu q} a_{r'}^{s\mu q} a_{r'}^{s\mu q} | \Psi \rangle = \rho_q(r s_\mu, r s_\mu) \rho_q(r' s_\mu, r' s_\mu) - |\rho_q(r s_\mu, r' s_\mu)|^2. \quad (3.2)$$

Because of the Pauli exclusion principle, $P_{qs\mu}(r, r) = 0$. Given that a nucleon with spin $s_\mu$ and isospin $q$ is located at position $r$, the conditional probability of finding another nucleon with the same spin and isospin at position $r'$ is

$$R_{qs\mu}(r, r') = \frac{P_{qs\mu}(r, r')}{\rho_q(r s_\mu, r s_\mu)}. \quad (3.3)$$

To find the conditional probability of like-spin and like-isospin nucleons in the vicinity of each other, we assume that the second nucleon is located within a small spherical shell with radius $\delta$ around $r$; the conditional probability can then be written as:

$$R_{qs\mu}(r, r + \delta) = e^{\delta \nabla'} R_{qs\mu}(r, r') \bigg|_{r=r'}. \quad (3.4)$$

Let $\langle \cdots \rangle$ denote an average over the spherical shell. The first non-zero term in the Taylor
expansion of \( \langle R_{qs\mu}(\mathbf{r}, \mathbf{r} + \delta) \rangle \) with respect to the radius \( \delta \) is [133]

\[
\frac{1}{6} \delta^2 \nabla^2 R_{qs\mu}(\mathbf{r}, \mathbf{r}') \bigg|_{\mathbf{r} = \mathbf{r}'} = \frac{1}{3} D_{qs\mu}(\mathbf{r}) \delta^2, \tag{3.5}
\]

where

\[
D_{qs\mu} = \tau_{qs\mu} - \frac{1}{4} \frac{\left| \nabla \rho_{qs\mu} \right|^2}{\rho_{qs\mu}} - \frac{\left| j_{qs\mu} \right|^2}{\rho_{qs\mu}}, \tag{3.6}
\]

is a localization measure that captures the short-range behavior of \( R_{qs\mu}(\mathbf{r}, \mathbf{r}') \). The derivation for Eq. (3.6) is based on the density-matrix expansion technique [89, 134]; details can be found in Ref. [135]. Following Ref. [36], we define a dimensionless and normalized NLF:

\[
C_{qs\mu}(\mathbf{r}) = \left[ 1 + \left( \frac{D_{qs\mu}(\mathbf{r})}{\tau_{qs\mu}^{\text{TF}}(\mathbf{r})} \right)^2 \right]^{-1}, \tag{3.7}
\]

where \( \tau_{qs\mu}^{\text{TF}}(\mathbf{r}) = \frac{3}{5} \left( 6\pi^2 \right)^{2/3} \left( \frac{5}{3} \right)^{5/3} \rho_{qs\mu}(\mathbf{r}) \) is the Thomas-Fermi kinetic density. The value of \( C \) can vary in the range of \([0, 1]\); the smaller the probability of finding two like-spin particles near each other, the more localized they are, and the larger the NLF is.

According to Eq. (3.1), densities entering the NLF are composed of both time-even and time-odd terms. In a time-reversal invariant system that is spin-saturated or governed by spin-independent interactions, we have \( D_{q,s\mu} = \frac{1}{2} \tau_q - \frac{1}{8} \left| \nabla \rho_q \right|^2 / \rho_q \). We also note that the \( \mathbb{J}_q(\mathbf{r}) \) term does not vanish even when the time-reversal symmetry is conserved [136]. Hence, the current \( j_{qs\mu}(\mathbf{r}) \) does not vanish in the ground-state configuration of an even-even nucleus unless the system is spin-saturated. Although the contribution of the current \( j_{qs} \) to the NLF was ignored in previous works [40, 44, 43], it actually almost vanishes inside the nucleus (see Sec. 3.3.2), so we can safely neglect it when using the NLF as a visualization tool.
3.1.2 Alternative interpretation

Besides interpreting the NLF as a measure of the conditional probability of like-spin pairs, there is an alternative interpretation which allows the generalization of NLF’s definition. As discussed in Refs. [37, 137], the localization function can also be interpreted in terms of the Pauli exclusion principle. Let us assume that one isolated fermion of given spin \( s_\mu \) and isospin \( q \) is located in some region. The wave function of this particle can be written as

\[ \psi_{qs\mu}(r) = \sqrt{\rho_{qs\mu}} e^{i\chi(r)} \],

where \( \chi(r) \) is a position-dependent phase factor whose gradient is related to the current density as \( j_{qs\mu} = \rho_{qs\mu} \nabla \chi \). The corresponding s.p. kinetic density is the sum of last two terms in \( D_{qs\mu} \) (3.6):

\[ \tau_{qs\mu}^\text{s.p.} = \left| \nabla \psi_{qs\mu} \right|^2 = \frac{1}{4} \left( \frac{\nabla \rho_{qs\mu}}{\rho_{qs\mu}} \right)^2 + \frac{\left| j_{qs\mu} \right|^2}{\rho_{qs\mu}}, \tag{3.8} \]

where the first term is the von Weizsacker kinetic energy density [138]. Therefore, the localization measure \( D_{qs\mu} = \tau_{qs\mu} - \tau_{qs\mu}^\text{s.p.} \) can be interpreted as the excess of kinetic density due to the Pauli exclusion principle. This interpretation of the NLF is more flexible since it does not involve the notion of the conditional probability; with the new interpretation one can straightforwardly generalize the NLF to the case of point-group symmetries.

For the rotating systems to be discussed in following sections, parity \((\hat{P})\), \(y\)-signature \((\hat{R}_y = e^{-i\pi \hat{J}_y})\), and \(y\)-simplex \((\hat{R}_y = \hat{P} \hat{R}_y)\) symmetries are conserved. To display the effects caused by different s.p. orbits, it is convenient to study the NLF of a given \( y \) signature \( r_y \) or \( y \)-simplex \( \tau_y \). This can be done by expressing local densities in terms of their symmetry-conserving components. For instance, when \( y \) simplex is conserved, we have

\[ \rho_q(r) = \rho_{q\bar{y}=+1}(r) + \rho_{q\bar{y}=-1}(r), \tag{3.9} \]
where $\sigma_y \equiv r_y/i = \pm 1$. In practice, the component $\rho_{q\sigma_y}$ is computed via summing up the contributions from s.p. levels with $y$ simplex $r_y = i\sigma_y$ [112, 139, 140]. The kinetic density $\tau_q(r)$ and current $j_q(r)$ can be similarly decomposed. With $\rho_{q\sigma_y}$, $\tau_{q\sigma_y}$ and $j_{q\sigma_y}$, the NLF for quantum number $q\sigma_y$ can be defined as

$$D_{q\sigma_y} = \tau_{q\sigma_y} - \frac{1}{4} \left| \nabla \rho_{q\sigma_y} \right|^2 - \left| \frac{j_{q\sigma_y}}{\rho_{q\sigma_y}} \right|^2, \quad \mathcal{C}_{q\sigma_y}(r) = \left[ 1 + \left( \frac{D_{q\sigma_y}(r)}{\tau_{TF}^{q\sigma_y}(r)} \right)^2 \right]^{-1},$$

(3.10)

where $\tau_{TF}^{q\sigma_y}(r) = \frac{3}{5} \left( 6\pi^2 \right)^{2/3} \rho_{q\sigma_y}^{5/3}(r)$. One should note that densities $\rho_{q\sigma_y}$, $\tau_{q\sigma_y}$ and $j_{q\sigma_y}$ can also be decomposed into time-even and time-odd components in the same manner as Eq. (3.1).

### 3.2 Cranked calculations

The two examples we employ for the study of the NLF in rotating nuclear systems are superdeformed (SD) $^{152}$Dy and the cranked harmonic-oscillator (CHO) model. The cranked Hartree-Fock (CHF) calculation for the rotating $^{152}$Dy is presented in Sec. 3.2.1; then the CHO model is discussed in Sec. 3.2.2.

#### 3.2.1 Cranked Hartree-Fock calculation

The CHF method is utilized to study the rotational band of SD $^{152}$Dy. Extreme single-particle behavior is seen in this system [141, 142] and the pairing correlation is absent due to large SD gaps at $Z = 66$ and $N = 86$ as well as rapid rotation [143, 144, 145]. The collective rotation of $^{152}$Dy has been investigated in many works, e.g., Refs. [146, 147, 148, 94]. We follow the procedure given in Ref. [146] and perform CHF calculations with HFOOD.
Single-particle wave functions are expanded in a deformed Cartesian HO basis with frequencies $h\omega_z = 6.246$ MeV and $h\omega_\perp = 11.200$ MeV along the directions parallel and perpendicular to the symmetry axis, respectively, and the total number of basis states is 1013 with HO quanta not exceeding 15 in each direction. The Skyrme parametrization SkM* [97] with its generic time-odd terms [146, 149] is adopted for the CHF calculations.

The main idea of the cranked calculation has already been discussed around Eq. (2.51) in Sec. 2.2.3. Here we follow the notation adopted in Sec. 2.2.3 and assume that the system rotates around the $y$ axis. In our CHF calculations, parity, $y$-signature, and $y$-simplex symmetries are preserved while time-reversal and axial symmetries are broken; see Refs. [112, 139, 140] for more discussions. One should note that operator $\hat{R}_y$ ($\hat{R}_y$) is time-odd and thus time-reversed s.p. states belong to opposite signature (simplex) eigenvalues. With conserved parity and $y$ signature, the CHF configuration can be labeled in terms of parity-signature blocks $[N_+,+i, N_+-,i, N_-,+i, N_-,+i]$, where $N_{\pi y}$ denotes the number of occupied s.p. orbits with parity $\pi$ ind $y$-signature $r_y$. The yrast configuration of SD $^{152}$Dy is $[22, 22, 21, 21]_n \otimes [16, 16, 17, 17]_p$ [146]. The relative variation of the quadrupole moment $Q_{20}$ within this configuration is less than 1% in the frequency range $h\omega = 0.2 \sim 0.5$ MeV [141], so we constrain $Q_{20}$ at 42 b to eliminate its possible influence on the NLF.

Single-particle Routhians for the SD yrast band of $^{152}$Dy are shown in Fig. 3.1, where SD shell closures are clearly exhibited. In the figure, rotation-aligned s.p. orbits are marked by thicker lines; they are dramatically impacted by rotation and tend to align their angular momenta along the rotational axis ($y$ axis) as the rotational frequency $\omega$ rises. Some of them are intruder orbits denoted by their main HO components with large principal quantum numbers $N$; they are the lowest $N = 7$ neutron and $N = 6$ proton levels [150, 151]. Other rotation-aligned orbits are labeled by the asymptotic quantum numbers (Nilsson quantum
Figure 3.1: Single-particle neutron (a) and proton (b) Routhians as functions of $\omega$, obtained from the CHF+SkM* calculations for the SD yrast band of $^{152}$Dy. The $(\pi, r_y)$ combinations are denoted by solid lines ($+, +i$), dotted lines ($+, -i$), dot-dashed lines ($-, +i$), and dashed lines ($-, -i$). Single-particle Routhians of the lowest neutron $N = 7$, proton $N' = 6$, and proton $[541]1/2$ levels are marked by thicker lines.

numbers) $[N n_z \Lambda]\Omega$ of their dominant HO components. On the other hand, some s.p. states around the Fermi level are weakly affected by rotation and thus known as deformation-aligned [32, 152, 153]; they tend to align their angular momenta along the $z$ axis, the symmetry axis at $\omega = 0$.  

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Figure 3.2: Single-particle Routhians of the SD CHO model belonging to supershells $N_{\text{shell}} = 6$ and 7. The CHO quantum numbers $[n_1, n_2, n_3]$ are given in brackets. Positive-parity and negative-parity orbits are marked by solid and dashed lines, respectively. The rotational frequency $\omega$ is expressed in units of $\omega_0 = 3^{1/3} \sqrt{\omega_z \omega_{\perp}^2}$ while the Routhians $E$ in units of $\hbar \omega_z$. Each level is doubly degenerate due to the two possible spin orientations. The crossing between the lowest $N = 7$ Routhian $[0,0,7]$ and the $[3,0,0]$ Routhian at $\omega/\omega_0 \approx 0.2$ is denoted by the arrow.

### 3.2.2 Cranked harmonic-oscillator (CHO) calculation

In the previous study of the NLF, the deformed harmonic-oscillator model was employed to provide illustrative guidance [41]. For a rotating nucleus, the corresponding model for illustration is the cranked harmonic oscillator (CHO) that rotates around the $y$ axis. In the CHO model we neglect the alignment of spin along the rotational axis and assume that the
s.p. Hamiltonian is spin-independent:

\[ \hat{H}_{\text{CHO}}^{\text{s.p.}} = \frac{\hat{p}^2}{2m} + \frac{1}{2} m \omega_\perp^2 (\hat{x}^2 + \hat{y}^2) + \frac{1}{2} m \omega_z^2 \hat{z}^2 - \omega \hat{L}_y, \]  

(3.11)

where \( \omega_z \) and \( \omega_\perp = 2 \omega_z \) are HO frequencies, \( \omega \) is the angular velocity of rotation, and \( \hat{L}_y = \hat{z} \hat{p}_x - \hat{x} \hat{p}_z \) is the orbital angular-momentum operator in the \( y \) direction. All the s.p. levels are doubly degenerate and the NLFs of different spins (\( y \) signatures or \( y \) simplexes) are identical. Analytical expressions for the s.p. Routhians and wave functions of the CHO model can be found in a number of publications, e.g., Refs. [33, 154, 155]. The CHO s.p. Routhian is

\[ E = \hbar \Omega_+ (n_1 + \frac{1}{2}) + \hbar \omega_y (n_2 + \frac{1}{2}) + \hbar \Omega_- (n_3 + \frac{1}{2}) \]  

(3.12)

where

\[ \Omega_\pm = \sqrt{\frac{\omega_\perp^2 + \omega_z^2}{2} + \omega^2 \pm \sqrt{(\omega_\perp^2 - \omega_z^2)^2 + 8 \omega^2 (\omega_\perp^2 + \omega_z^2)}}, \]  

(3.13)

and \( n_1, n_2 \) and \( n_3 \) become HO quantum numbers in \( x, y \) and \( z \) directions when \( \omega \) is zero. It should be noted that the consistency relation between the mean-field ellipsoidal deformation and average density distribution [32, 155] is not adopted here.

As an analytical counterpart of SD doubly-magic \(^{152}\text{Dy}\), the CHO model should also be closed-shell. Figure 3.2 presents the s.p. Routhians of the CHO system with 60 fermions filling SD supershells up to \( N_{\text{shell}} \equiv 2(n_1 + n_2) + n_3 = 6 \) [32, 153, 156, 157]. As discussed in Ref. [33] and shown in Fig. 3.2, orbits with no CHO quanta along the rotation axis \( (n_2 = 0) \) and the largest possible value of the difference \( (n_3 - n_1) \) carry most of the angular momentum, and thus are most highly aligned; those are the \([0, 0, 7]\) and \([0, 0, 6]\) Routhians shown in the figure.

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3.3 Results and discussions

3.3.1 Time-odd local densities

In a rotating system, the collective rotational behavior is well characterized by the current density $\mathbf{j}$ [148, 158, 159, 160, 161, 162, 163, 164, 165]. Figure 3.3 displays how the current emerges in the CHO model as the rotational frequency $\omega$ increases to $0.2\omega_0$. As $\omega$ increases, a flow pattern close to a rigid-body rotation gradually develops inside the system. At $\omega = 0.2\omega_0$ there is dramatic growth in $|\mathbf{j}|^2$, indicating significant angular-momentum alignment; the system becomes more elongated as well. This effect results from the band crossing marked by the arrow in Fig. 3.2, where prolate s.p. level $[0, 0, 7]$ with large angular momentum becomes occupied while oblate $[3, 0, 0]$ level becomes empty.

Figure 3.4 presents the current distribution of $^{152}\text{Dy}$ obtained from CHF calculations, with the rotational frequency $\hbar\omega$ ranging from 0 up to 0.8 MeV (angular momentum $J_y$ from 0 to $90\hbar$). The leftmost column in the figure displays the result of the benchmark FAM-QRPA calculation [166], which corresponds to the limit of $\omega \to 0$. One can find flow patterns close to the rigid-body rotation in both FAM-QRPA and CHF results. Since the
irrotational flow is mainly attributed to pairing correlations [161, 166], it is expected that no significant irrotational current is produced in the CHF calculations without static pairing.

Besides the current $\vec{j}$, three other time-odd densities are also present in the expression of the NLF: the spin density $\vec{s}$, spin-kinetic density $\vec{T}$, and spin-current tensor $\vec{J}$. Figure 3.5 shows the distributions of $\vec{s}$ and $\vec{T}$ at a number of rotational frequencies, while Fig. 3.6 displays $\vec{J} \cdot \vec{e}_y$, the projection that enters the NLF when the $y$ axis is selected as the spin-quantization axis. Both $\vec{s}$ and $\vec{T}$ are polarized in the $y$ direction, parallel to the aligned angular momentum. As frequency $\omega$ increases, the magnitudes of $\vec{s}$ and $\vec{T}$ gradually increase while their directions hardly vary. The distributions of $\vec{J} \cdot \vec{e}_y$ shown in Fig. 3.6 are mainly present at the nuclear surface, similar to the current $\vec{j}$. On the other hand, the spin-current tensor does not vanish at $\omega = 0$ and depends weakly on the rotational frequency.
Figure 3.5: Spin density $s$ (top) and spin-kinetic density $T$ (bottom) in the $x$-$y$ ($z = 0$) plane for neutrons in the SD yrast band of $^{152}$Dy, as functions of the rotational frequency $\omega$ (in MeV/$\hbar$). The magnitudes, $|s|$ (in fm$^{-3}$) and $|T|$ (in fm$^{-5}$), are shown by color and line thickness. The FAM-QRPA results are presented in the leftmost column with a different color range.

### 3.3.2 Simplified nucleon localization function

An important consequence of the surface characters of $j$ and $J \cdot e_y$ is that they only contribute significantly to the NLF at the surface. This observation should be valid in most cases even if the irrotational flow exists (see examples in Refs. [161, 166]). The same feature is also observed in the distribution of squared density gradient $|\nabla \rho_{q\mu}|^2$ since the particle density is quite flat in the nuclear interior. Therefore, we can define a simplified localization function

Figure 3.6: Spin-current tensor density $\mathcal{J} \cdot e_y$ in the $x$-$z$ ($y = 0$) plane for neutrons in the SD yrast band of $^{152}$Dy, as a function of the rotational frequency $\omega$ (in MeV/$\hbar$). Its magnitude (in fm$^{-4}$) is shown by color and line thickness.
as

$$C_{qs\mu}^\tau (r) = \left[ 1 + \left( \frac{\tau_{qs\mu}}{\tau_{TF}} \right)^2 \right]^{-1},$$

(3.14)

which does not include the current $j_{qs\mu}$ and density gradient $\nabla \rho_{qs\mu}$. Figure 3.7 presents the NLFs (3.7), simplified NLFs (3.14), and their differences in the CHO model. In the figure, the simplified NLF shows the same pattern as the usual NLF, except the lack of strong enhancement resulting from the current and density gradient at the surface. Similar behavior is also observed in the yrast band of $^{152}$Dy: Figure 3.8 shows the comparison between the distributions of $C$ and $C^\tau$ for neutrons with $\hat{\sigma}_y = -1$ ($y$-simplex $v_y = -i$) at rotational frequency $\hbar \omega = 0.9 \text{ MeV}$, and the difference $C - C^\tau$ is only significant at the surface. This difference is less pronounced at lower rotational frequencies. One can see that the simplified NLF eliminates the large value of the NLF at the nuclear surface, and it is
Figure 3.8: $C$ (left), $C^\tau$ (middle), and their difference (right) in the $x$-$z$ ($y = 0$) plane for neutrons with $\hat{\sigma}_y = -1$ ($y$-simplex $\tau_y = -1$) in the SD yrast band of $^{152}$Dy at rotational frequency $\hbar \omega = 0.9$ MeV.

thus unnecessary to normalize the NLF as $C_{qs} \rightarrow C_{qs} \rho_{qs} \big/ \max \rho_{qs}$ [44].

Considering the interpretation of $D_{qs\mu}$ as the excess of the kinetic density due to the Pauli principle, we are not surprised to see that $|\nabla \rho_{qs\mu}|^2$ and $|\hat{j}_{qs\mu}|^2$ are non-negligible only at the surface where only a limited number of s.p. orbits are prominent and “localized.” Therefore, we can replace the old NLF with the simplified NLF in most cases, except perhaps some extremely dynamic processes where the current and density gradient can become appreciable inside the nucleus. In addition, this interpretation also helps the study of the Pauli energy in heavy-ion fusion reactions [51].

3.3.3 Dependence on the choice of spin-quantization axis

As mentioned in Sec. 3.1, the NLF $C_{qs\mu}$ (3.7) and its simplified version $C^\tau_{qs\mu}$ (3.14) depend on the choice of the spin quantization direction $\mu$. This dependence is visualized in Figs. 3.9 ($y = 0$ plane) and 3.10 ($x = 0$ plane) for the rotating $^{152}$Dy at frequency $\hbar \omega = 0.5$ MeV. It is shown that the NLF depends slightly on the choice of the quantization axis $\mu$ as well as the plane selection. One can also notice the relation $C^\tau_{qs\mu} \approx C_{qs\mu}$ inside the nuclear volume, no matter which quantization axis $\mu$ and which cross section are selected.
3.3.4 Angular-momentum alignment in the CHO model

Figure 3.7 has shown the NLFs of the CHO model in the $y = 0$ plane at different rotational frequencies; patterns in the $x = 0$ plane are similar. A pattern similar to that shown in Ref. [43] for the deformed HO model can be clearly observed at $\omega = 0$, but as $\omega$ increases the pattern gradually becomes blurred. At $\omega = 0.2\omega_0$ where the band crossing occurs, the NLF changes dramatically: The number of maxima along the $z$ axis increases due to the occupation of the $[0, 0, 7]$ state; the number of maxima in the $x$ direction, on the other hand, decreases since the $[3, 0, 0]$ orbit becomes empty. Thus, the NLF is a good indicator for the shell structure and can easily visualize the effect of band crossing.
Figure 3.10: Similar to Fig. 3.9 but shown in the \( y-z (x=0) \) plane.

To better display the evolution of the simplified NLF \( C^\tau \) induced by rotation, in Fig. 3.11 we show the difference

\[
\Delta C^\tau (r; \omega) \equiv C^\tau (r; \omega) - C^\tau (r; \omega = 0),
\]

(3.15)
together with density variations \( \Delta \tau \equiv \tau (r; \omega) - \tau (r; \omega = 0) \) and \( \tau^{\text{TF}} \equiv \tau^{\text{TF}} (r; \omega) - \tau^{\text{TF}} (r; \omega = 0) \). Figure 3.11 shows a clear correspondence between the peaks of \( \Delta C^\tau \) and valleys (peaks) of \( \Delta \tau \) (\( \Delta \tau^{\text{TF}} \)), which is consistent with Eq. (3.14). This phenomenon indicates that \( \Delta \tau \) and \( \Delta \tau^{\text{TF}} \) oscillates in antiphase, causing a “constructive interference” when we compute their ratio.

One can notice that this out-of-phase oscillation already exists in the patterns of \( \tau \) and \( \tau^{\text{TF}} \) at \( \omega = 0 \). Figure 3.12(a) shows \( \tau \), \( \tau^{\text{TF}} \) and \( C^\tau \) of the non-rotating HO model along the \( z \) axis \((x=y=0)\), as well as the density profile of the \([0,0,6]\) orbit. We see that the valleys
Figure 3.11: $C^\tau$ (top), $\tau$ (in fm$^{-5}$, middle) and $\tau_{\text{TF}}$ (in fm$^{-5}$, bottom) in the $x$-$z$ ($y = 0$) plane, obtained from the CHO model. The leftmost column shows the reference plots at $\omega = 0$ while the other columns show the rotational dependence relative to the $\omega = 0$ reference as a function of the rotational frequency $\omega$ (in units of $\omega_0$).

(peaks) of $\tau$ ($\tau_{\text{TF}}$ or $C^\tau$) roughly coincide with the maximum points of the s.p. density $|\psi_{006}|^2$, while lower s.p. levels provide a smooth background for the total $\tau$ ($\tau_{\text{TF}}$). This observation is more evident in the one-dimensional (1D) HO model, as presented in Fig. 3.12(b); in this 1D HO model s.p. orbits with quantum number $N \leq 6$ are occupied. The antiphase correspondence is expected because the kinetic density $\tau$ is based on the gradient of s.p. wave functions while the $\tau_{\text{TF}}$ depends on the particle density $\rho$ and involves no derivatives. The oscillating patterns of $\tau$ and $\tau_{\text{TF}}$ basically reflect the characteristic nodal structure of high-$N$ s.p. orbits, allowing us to have a sense of the shell structure by counting the number of maxima; the NLF then successfully magnifies this nodal pattern thanks to the “constructive interference” between $\tau$ and $\tau_{\text{TF}}$. In addition, the nodal structure is closely associated with clustering, which has been discussed in many works, e.g., Refs. [165, 167, 168, 169]; and hence the NLF can also visualize clusters inside the nucleus.
Figure 3.12: \( C^\tau \) (thick solid line), \( \tau \) (solid line), and \( \tau^{\text{TF}} \) (dashed line) in the non-rotating HO model along the \( z \) axis \((x = y = 0)\). (a) 3D SD HO potential with 60 particles. The density profile of the \([0,0,6]\) orbit is marked by a dotted line. (b) 1D case. HO orbits with quantum number \( N \leq 6 \) are occupied. The density profile of the \( N = 6 \) orbit is marked by a dotted line; here \( \tau^{\text{TF}} = \pi^2 \rho^3 / 3 \). Some quantities are scaled for better visualization.

From the s.p. perspective, the density differences shown in Fig. 3.11 are related to the particle-hole (p-h) excitations across the Fermi level induced by the cranking operator \( \omega \hat{L}_y \), especially the low-energy transitions with \( \Delta n_1 = \pm 1 \), \( \Delta n_2 = 0 \), \( \Delta n_3 = \mp 1 \). Crossings and mixtures of s.p. levels far below the Fermi energy, on the other hand, can barely impact densities. Figure 3.13 displays the kinetic-density variations generated by the p-h excitations from the occupied supershell \( N_{\text{shell}} = 6 \) to the unoccupied \( N_{\text{shell}} = 7 \) (see Fig. 3.2 for supershell notations). By summing all the p-h contributions one can obtain the last panel of Fig. 3.13, where we see a pattern similar to that of Fig. 3.11 at \( \omega = 0.15 \omega_0 \). The oscillating pattern along the \( z \) axis primarily comes from the excitation of \([0,0,6] \rightarrow [1,0,5]\). Other transitions produce more structures in the horizontal direction; the most extreme
Figure 3.13: Variations in the kinetic energy density $\tau$ due to p-h excitations (at $\omega = 0$) from supershell $N_{\text{shell}} = 6$ to $N_{\text{shell}} = 7$ shown in Fig. 3.2. These excitations are induced by the cranking term in the CHO model. The rightmost panel displays the average of all the p-h contributions.

One is $[2, 0, 2] \rightarrow [3, 0, 1]$. Both $[0, 0, 6]$ and $[1, 0, 5]$ orbits are prolate and rotation-aligned, corresponding to $[660]1/2$ ($61_2$) and $[651]3/2$ ($63_4$) Nilsson levels, respectively; $[2, 0, 2]$ and $[3, 0, 1]$, on the other hand, are deformation-aligned and resemble $[420]1/2$ ($[422]3/2$) and $[411]3/2$ ($[413]5/2$). In short, the strong imprints left by p-h transitions involving rotation-aligned levels can be identified through the characteristic NLF variation along the major axis, while those involving deformation-aligned orbits contribute to patterns along the minor axis.

### 3.3.5 Angular momentum alignment in the CHF calculation

In this section we analyze the NLF patterns obtained from CHF calculations for the yrast band of $^{152}\text{Dy}$. Figure 3.14 shows the simplified NLFs $C^\tau_{q\bar{q}y}$ for different $y$ simplexes and isospins in the $y = 0$ plane; patterns in the $x = 0$ plane are similar. The first column of the figure gives the NLFs of the non-rotating case, where one can identify the characteristic pattern of a deformed nucleus, similar to those presented in Ref. [44]. As frequency $\omega$ rises, new patterns gradually develop inside the nuclear volume, and the NLFs of opposite simplexes deviate from each other because of time-reversal symmetry breaking.

Similar to Fig. 3.11, NLF differences $\Delta C^\tau_{q\bar{q}y}$ (3.15) in the $y = 0$ plane are presented in
Figure 3.14: The NLF $C_{q\tilde{y}}^\tau$ in the $x$-$z$ ($y = 0$) plane as a function of $\omega$ (in MeV/$h$), obtained from CHF calculation for the SD yrast band of $^{152}$Dy. The symbols $\uparrow$ and $\downarrow$ represent $\tilde{\sigma}_y = +1$ and $-1$ ($y$-simplex $r_y = +i$ and $-i$), respectively.

Fig. 3.15 for the yrast band of $^{152}$Dy; density variations $\Delta \tau_{q\tilde{y}}$ and $\Delta \tau_{TF}^\tau$ are also shown in Figs. 3.16 and 3.17, respectively. One can see a clear connection between the patterns of $\Delta \tau$ ($\Delta \tau_{TF}$) and $\Delta C^\tau$, similar to those in the CHO model, so the mechanism of the “constructive interference” discussed in Sec. 3.3.4 is still valid in the realistic case. Moreover, in Figs. 3.15, 3.16 and 3.17, $\Delta \tau$, $\Delta \tau_{TF}$, and $\Delta C$ of opposite simplexes vary in a roughly opposite manner when frequency $\omega$ increases. This can be explained via decomposing $\Delta \tau_{q\tilde{y}}$ and $\Delta \rho_{q\tilde{y}}$ into
time-even and time-odd components. For example, the difference $\Delta \tau_{q\bar{\sigma}_y}$ can be written as

$$\Delta \tau_{q\bar{\sigma}_y}(r; \omega) = \frac{1}{2} \Delta \tau_q(r; \omega) + \frac{1}{2} \bar{\sigma}_y T'_q(r; \omega),$$  \hspace{1cm} (3.16)$$

where the time-even term $\Delta \tau_q(r; \omega) = \tau_q(r; \omega) - \tau_q(r; \omega = 0)$ produces the same background in $\Delta \tau_{q\bar{\sigma}_y} = \pm 1$, while the time-odd term $T'_q(r; \omega) = \tau_{q\bar{\sigma}_y} = 1(r; \omega) - \tau_{q\bar{\sigma}_y} = -1(r; \omega)$ results in the difference between the NLFs of different simplexes.
Figure 3.16: Similar to Fig. 3.15, but for $\Delta \tau_{q\vec{\sigma}y}$ (in fm$^{-5}$). The reference value of $\tau_{q\vec{\sigma}y}$ at $\omega = 0$ is shown in the leftmost column.

For the connection between the NLFs and s.p. orbits in the rotating $^{152}$Dy, we follow Sec. 3.2.2 and concentrate on the study of $\Delta \tau$. It is difficult to figure out individual p-h excitations in the realistic case, so we track individual s.p. levels near the Fermi energy from $\hbar \omega = 0$ up to 0.1 MeV (see Fig. 3.1) and investigate the evolution of their kinetic densities. When the rotational frequency becomes larger than 0.1 MeV, it is difficult to identify contributions from different s.p. orbits due to strong level mixing. Figure 3.18 presents the kinetic-density variations $\Delta \tau$ for different parity-signature combinations when $\omega$ grows from 0 to 0.1 MeV. These patterns can be approximately reproduced with contributions from a few occupied s.p. levels close to the Fermi energy, as shown in Fig. 3.19. Among neutron orbits with negative parity, rotation-aligned levels $7_{1,2}$ with high $N$ provide important contributions to
Figure 3.17: Similar to Fig. 3.16, but for $\Delta \tau_{TF}^{q\sigma_y}$ (in fm$^{-5}$).

the vertical oscillation of $\Delta \tau_n$. Among neutron orbits with positive parity, the highest four occupied levels, $[651]1/2$, $[642]5/2$, $[413]5/2$, and $[411]1/2$, are closely lying and deformation-aligned; they are responsible for the patterns of $\Delta \tau_n$, especially the horizontal structures.

For protons, rotation-aligned orbits $6_{1,2,3,4}$ and $[541]1/2$ are most crucial and the quantum numbers of their dominant HO components explains their contributions to the difference $\Delta \tau_p$.

One can now reach the same conclusion as Sec. 3.3.4: The nodal structures of $\Delta \tau$ ($\Delta \tau_{TF}$ and $\Delta \mathcal{C}^\tau$) along the $z$ axis are attributed to rotation-aligned s.p. orbits below the Fermi energy with large $N$ and $n_3$, while their horizontal features are associated with deformation-aligned levels.
Figure 3.18: $\Delta \tau$ (in fm$^{-5}$) of neutrons (top) and protons (bottom) in the $x$-$z$ ($y = 0$) plane for different parity-signature combinations $(\pi, r_y)$ in the SD yrast band of $^{152}$Dy at rotational frequency $\hbar \omega = 0.1$ MeV.

3.4 Summary

In this chapter, the nucleon localization function has been studied in anisotropic, spin-unsaturated and spin-polarized rotating systems. The concept of the NLF is first generalized to the case of point-group symmetries with the help of the interpretation that the NLF measures the kinetic-density excess owing to the Pauli principle. Then we propose a simplified NLF that is easier to compute and interpret. It is shown that the “constructive interference” between the kinetic density $\tau$ and Thomas-Fermi kinetic density $\tau_{TF}$ makes the NLF a powerful tool for the visualization of the nodal structure of high-lying s.p. orbits, which explains why the NLF pattern is closely connected with the shell structure and clustering. In the rotating $^{152}$Dy, time-odd effects are clearly displayed in the NLFs of opposite simplexes. Also, we see the relation between s.p. levels and the NLF pattern’s evolution: The NLF structure along the major axis is related to occupied rotation-aligned states close
Figure 3.19: Contributions to the variation $\Delta \tau$ (in fm$^{-5}$) in the $x$-$z$ ($y = 0$) plane for different parity-signature blocks from individual s.p. Routhians in $^{152}$Dy at $\hbar \omega = 0.1$ MeV: the four positive-parity neutron levels $[651]1/2$, $[642]5/2$, $[413]5/2$ and $[411]1/2$ with signature $r_y = +i$ (a) and $-i$ (b) that lie below the $N = 86$ shell gap in Fig. 3.1 (see Fig. 1 of Ref. [151] for the asymptotic quantum numbers $[Nn_z\Lambda]\Omega$ of s.p. levels in the SD $^{152}$Dy); the $N = 7$ neutron intruder states $7_1$ (c) and $7_2$ (d); the $N = 6$ proton intruder states $6_2 + 6_4$ (e) and $6_1 + 6_3$ (f); and the $[541]1/2_2$ (g) and $[541]1/2_1$ (h) proton states.

to the Fermi energy, while the pattern in the perpendicular direction results from high-lying deformation-aligned states.

Results discussed in this chapter demonstrates that the NLF is helpful for the study of nuclear rotation, and we expect its use for other collective motions and time-dependent processes. We are also considering the extension of the NLF to the HFB case and beyond-mean-field frameworks for new insights into many-body correlations. The kinetic density has already been computed within the no-core shell model [170], which paves the way for the application of the NLF in the $ab initio$ framework.
Chapter 4

Origin of reflection-asymmetric shapes

This chapter focuses on the microscopic origin of ground-state reflection-asymmetric deformations of even-even nuclei and investigates this problem from the perspectives of the multipole expansion and s.p. spectrum. Results discussed in the following has been published in Ref. [121], and main conclusions are presented for Ra and Yb isotopes. The multipole expansion of the total energy is discussed in Sec. 4.1, followed by the behaviors of various multipole components shown in Sec. 4.2. In Sec. 4.3 the origin of pear-like deformations are studied through the spectra of canonical s.p. states.

4.1 Multipole expansion of the EDF

The multipole expansion of the total energy was first adopted in Ref. [171] and further studied in Ref. [172] to reveal the origin of quadrupole deformations. We generalize the concept of the expansion for the EDF, and employ it for reflection-asymmetric shapes. Since the octupole moment is the lowest nonzero moment related to the reflection-symmetry violation, the pear-like shape is often referred to as “octupole” deformation.

In the Skyrme EDF presented in Sec. 2.1, each term inside the integrand can be written in the form of \( \Gamma(r) \rho(r) \), where \( \rho \) is some density. When the time-reversal symmetry is conserved, \( \rho \) can be \( \rho, \bar{\rho}, \tau, \nabla^2 \rho, \nabla \cdot J \) or \( \mathbb{J}_{ij} \) \( (i, j = x, y, z) \). For the term \( C_i^\tau \rho_{tt3}(r) \tau_{tt3}(r) \), one can choose \( \Gamma = C_i^\tau \rho_{tt3} \) and \( \rho = \tau \), and it does not matter whether the coupling \( C^\tau \)
is density-dependent. With the axial symmetry imposed, the field \( \Gamma \) and density \( \rho \) can be expanded with the help of spherical harmonics \( Y_{LM} \) as [13]

\[
\xi(r) = \sum_{L} \xi_{[L]}(r) Y_{L, M=0}(\Omega), \quad \xi = \Gamma \text{ or } \rho,
\]

where \( \xi_{[L]}(r) = \int d\Omega \xi(r) Y_{L, M=0}^*(\Omega) \). The corresponding energy becomes

\[
E^{(\Gamma \rho)} = \int d\mathbf{r} \Gamma(\mathbf{r}) \rho(\mathbf{r}) = \sum_{L} \int d\mathbf{r} \Gamma_{[L]} \rho_{[L]} = \sum_{L} E^{(\Gamma \rho)}_{[L]},
\]

where \( \int d\mathbf{r} \Gamma_{[L]} \rho_{[L']} (L \neq L') \) vanishes thanks to the orthogonality of spherical harmonics.

In the isospin representation, the density that determines \( \Gamma \) is in the same isospin channel of \( \rho \), and we can calculate the multipole components of isoscalar \((t = 0)\) and isovector \((t = 1)\) energies to compare their contributions to the appearance of reflection-asymmetric shapes. The kinetic ED enters the isoscalar energy as the Skyrme parametrization we use in this chapter adopts the same proton and neutron masses. Like-particle pairing EDF (2.21) with \( V_p \neq V_n \), and the Coulomb interaction (2.22), however, are neither isoscalar nor isovector because they breaks the isospin symmetry. Without proton-neutron mixing, one can also write the EDF in terms of neutron and proton densities with the help of Eq. (2.9), and the total energy is then decomposed into neutron-neutron (n-n), proton-proton (p-p) and proton-neutron (p-n) components. The Coulomb energy contributes to the p-p energy, while neutron and proton pairing energies are included in n-n and p-p energies, respectively. One can also investigate the multipole components of n-n, p-p and p-n energies to examine their roles in the onset of reflection-asymmetric deformations.
Figure 4.1: The octupole deformation energy $\Delta E$ of $^{224}$Ra (a), its multipole expansion $\Delta E_{[L]}$ (b), and the components of the octupole part $\Delta E_{[3]}$ in isospin and proton-neutron representations (c), as functions of the octupole deformation $\beta_3$.

4.2 Energy expansion and reflection asymmetry

A recent global nuclear-DFT survey [31] showed that a number of even-even Ra isotopes have parity-violating ground states. We use $^{224}$Ra as an example and study its deformation with the HFBTHO program [110] and SLy4 Skyrme parametrization [98]. First the equilibrium quadrupole deformation $\beta_2$ is determined by parity-conserving HFB calculations; then with $\beta_2$ fixed at the equilibrium value, we vary the constraint on the octupole deformation $\beta_3$ and obtain the potential-energy curve shown in Fig. 4.1(a). The octupole deformation energy in the figure is defined as $\Delta E(\beta_3) = E(\beta_3) - E(\beta_3 = 0)$, and we see that the minimum-energy point is octupole-deformed.

Figure 4.1(b) presents the multipole components of the deformation energy

$$\Delta E_{[L]}(\beta_3) = E_{[L]}(\beta_3) - E_{[L]}(\beta_3 = 0), \quad (4.3)$$

as functions of $\beta_3$. We notice that the magnitudes of these components are much larger than the magnitude of the total deformation energy, so strong cancellation between them leads to the small deformation energy, and high-order components can be crucial for the determination of the minimum-energy point. Figure 4.1(c) shows the decomposition of $\Delta E_{[3]}$.
Figure 4.2: The octupole deformation energy $\Delta E$ (a), its multipole components $\Delta E_{[L]}$ (b), and the cumulative sum $\Delta E_{[0-L]}$ (c), along the isotopic chain of Ra. They are calculated at a fixed octupole deformation $\beta_3 = 0.05$. Isotopes with negative total deformabilities (octupole-deformed ground states) are marked by shading.

Figure 4.3: Similar to Fig. 4.2 but along the isotopic chain of Yb.

in both isospin and proton-neutron representations. The dominant contribution comes from the isoscalar part $\Delta E_{[3]}^{t=0}$ or p-n part $\Delta E_{[3]}^{pn}$, which is consistent with the simple estimate with a schematic particle-vibration coupling Hamiltonian [121]. Similar phenomenon has already been observed in the study of quadrupole deformations [171].

Now we turn to the behavior of energy decomposition along the isotopic chain of Ra. We plot in Fig. 4.2(a) octupole deformabilities, i.e., the octupole deformation energies at a small octupole deformation $\beta_3 = 0.05$, for even-even Ra isotopes. Figure 4.2(b) then shows the multipole components of these deformabilities. In the figure, significant variations of monopole and octupole parts are observed at around $N = 130$, the transition point of reflection-symmetry breaking. As the neutron number $N$ increases to 142, the region of pear-shaped nuclei ends and curves in Fig. 4.2(b) become flat. For comparison, Fig. 4.3(a)
shows the octupole deformabilities along the isotopic chain of Yb. One can see that there are no pear-shaped Yb ground states, and the corresponding multipole components given in Fig. 4.3(b) vary slowly as the neutron number $N$ increases.

To illustrate the cancellation between various multipolarities, Fig. 4.2(c) displays the cumulative sum

$$\Delta E_{[0-L]}(\beta_3 = 0.05) = \sum_{L'=0}^{L} \Delta E_{[L']}(\beta_3 = 0.05),$$

along the Ra isotopic chain. It is seen that a summation up to $L = 7$ is necessary to converge due to the strong cancellation between monopole and octupole components, especially for isotopes with negative total deformabilities, and that dotriacontapole ($L = 5$) components significantly contribute to the appearance of pear-like shapes. This observation is in agreement with previous works that show the important roles of high-order deformations in pear-shaped nuclei [173, 174, 175, 176, 177, 178, 179, 180].

### 4.3 Relation to single-particle spectra

From the s.p. perspective, the octupole deformation results from the coupling between closely-lying opposite-parity orbits (parity doublets) with angular-momentum difference $\Delta \ell = \Delta j = 3$. One can find such pairs of orbits across the Fermi energy when neutron or proton numbers are around $N_{\text{oct}} = 34, 56, 88$ or 134, slightly larger than the magic numbers, where the unique-parity intruder shell $(\ell, j)$ becomes close to the normal-parity shell $(\ell - 3, j - 3)$. Refs. [177, 179] noted that the dotriacontapole (and even higher-order) coupling is also related to the $\Delta \ell = \Delta j = 3$, consistent with the essential role of high-order multipolarities shown in the previous section. One should note that the s.p. argument regarding the appearance of reflection-asymmetric shapes is qualitative and cannot be used to exactly determine which
isotope is octupole-deformed. The quadrupole deformation makes the simple criterion on \(\Delta \ell\) and \(\Delta j\) difficult to apply as \(\ell\) and \(j\) are no longer good quantum numbers, and pairing correlations disfavor the emergence of deformations.

Figure 4.4 displays the canonical s.p. energies of \(^{224}\text{Ra}\) as functions of the quadrupole deformation \(\beta_2\). HFB calculations for this figure are carried out with the reflection symmetry imposed (\(\beta_3 = 0\)). The Fermi energies and equilibrium quadrupole deformations of a series of Ra isotopes are also marked. Around the Fermi energy, we find two pairs of shells, \(\pi 1i_{13/2} \leftrightarrow 2f_{7/2}\) and \(\nu 1j_{15/2} \leftrightarrow 2f_{2g_{9/2}}\), whose couplings drive the octupole shape. As the neutron number grows from 130 to 136, the quadrupole deformation emerges and breaks the spherical symmetry; then the octupole deformation appears and causes more level repulsion due to the interactions between opposite-parity pairs of states with the same angular-momentum projection \(\Omega\). After we pass the neutron number 136, the pear-like shape gradually becomes less favored as more s.p. levels originating from octupole-driving pairs of shells become occupied.

For comparison, the s.p. diagram of \(^{176}\text{Yb}\) is given in Fig. 4.5. Compared with \(^{224}\text{Ra}\), \(^{176}\text{Yb}\) has a larger equilibrium quadrupole deformation but no close-lying opposite-parity pairs of levels with the same \(\Omega\) value around the Fermi energy, and its ground state is thus reflection symmetric.

4.4 Summary

The microscopic origin of the reflection-asymmetric ground-state deformations is investigated in this chapter, with the help of the multipole expansion of the EDF and the spectrum of canonical s.p. states. The monopole and octupole energy components significantly contribute
to the appearance of pear-like deformations, but higher-multipolarity interactions are also crucial due to the strong cancellation between $\Delta E_{[0]}$ and $\Delta E_{[3]}$. We also see that the isoscalar part $\Delta E_{[3]}^{I=0}$ or the proton-neutron part $\Delta E_{[3]}^{pn}$ plays a dominant role in $\Delta E_{[3]}$. From the s.p. perspective, the emergence of reflection-asymmetric shapes is mainly attributed to the coupling between parity doublets with $\Delta \ell = \Delta j = 3$ around the Fermi energy; both octupole and dotriacontapole couplings between such pair of states are strong.
Figure 4.4: Canonical single-particle energies for neutrons (top) and protons (bottom) in $^{224}\text{Ra}$ as functions of the quadrupole deformation $\beta_2$. The reflection symmetry is imposed ($\beta_3 = 0$). Solid (dashed) lines represent levels with positive (negative) parity. The equilibrium quadrupole deformation of $^{224}\text{Ra}$ is given by a vertical dotted line, and the Fermi energies of $^{224}\text{Ra}$ are marked by dash-dotted lines. Quadrupole deformations and Fermi energies of even-even Ra isotopes with neutron number $N = 130 \sim 144$ are denoted by circles. All the Fermi energies are shifted with respect to the positions of spherical neutron $2g_9/2$ and proton $1h_9/2$ shells. The line color indicates the angular-momentum projection $\Omega$. 

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Figure 4.5: Similar to Fig. 4.4 but for $^{176}$Yb.
Chapter 5

Model calibration for beta-decay studies

This chapter discusses the EDF calibration for beta-decay calculations. The physics model we employ in the calibration has been presented in Sec. 2.3, and the $\chi^2$ optimization is carried out to determine optimal model parameters. Sections 5.1 and 5.2 discuss model parameters and experimental observables considered in the fit. Then the basic framework of the $\chi^2$ optimization is presented in Sec. 5.3, with numerical details discussed in Sec. 5.4. Results are shown in Sec. 5.5 and a summary is given in Sec. 5.6.

5.1 Model parameters

For beta-decay studies within the Skyrme-HFB-PNFAM framework, it is necessary to fit parameters that play important roles in the PNFAM but are not constrained in static HFB calculations. In the Skyrme interaction ED (2.16), only isovector ($t = 1$) terms with $t_3 = \pm 1$ contribute to the proton-neutron induced fields in the PNFAM. Because of time-reversal symmetry breaking, $\mathcal{H}^{(\text{odd})}$ (2.18) that is bilinear in time-odd densities must be taken into account, but corresponding coupling constants are not constrained by the properties of even-even nuclei and thus need to be calibrated.

The Skyrme parametrization we utilize for time-even Skyrme couplings and like-particle pairing strengths is UNEDF1-HFB [99], whose parameters have been well calibrated within the $\chi^2$-optimization framework. Following Ref. [66], we preset some time-odd couplings and
exclude them from the fit for simplicity. First, we assume that there is no density dependence in
the time-odd couplings we fit. Second, the local gauge invariance [146] can relate $C^j_t$ and
$C_t\nabla J$ to time-even couplings as $C^j_t = -C^r_t$, $C_t\nabla j = C_t\nabla J$. Third, we set $C_{1}^{\Delta s} = 0$ to avoid
finite-size instabilities [181]. Fourth, as for time-odd tensor terms, we only take $C_{1}^{T}$ into
account while $C_{1}^{F}$ and $C_{1}^{\nabla s}$ are set to zero so that the number of free parameters is limited.
In the end, the time-odd couplings that we need to calibrate are $C_{1}^{s}$ and $C_{1}^{T}$.

As for the particle-particle channel, we utilize the pairing ED in the isospin representation
(2.20), where neutron-neutron, proton-proton and proton-neutron pairing correlations are all
included. Both isoscalar and isovector terms in Eq. (2.20) contribute in the PNFAM, but
their strengths are determined in different ways. We use the average of neutron-neutron and
proton-proton pairing strengths as the isovector pairing strength $V_1$, i.e., $V_1 = (V_p + V_n) / 2$,
where $V_p$ and $V_n$ are defined in Eq. (2.21) and their values have been fitted together with
time-even Skyrme couplings to reproduce selected properties of even-even nuclei. Since
$V_p$ and $V_n$ will be the same as $V_1$ when the isospin symmetry is strictly preserved, our
choice of $V_1$ partially fulfills the consistency requirement between static HFB and PNFAM
calculations. The isoscalar pairing strength $V_0$, on the other hand, should be calibrated in
the same manner as $C_{1}^{s}$ and $C_{1}^{T}$. Although the isoscalar pairing is related to the Wigner
effect in nuclei with $N = Z$ [182], it is not yet reliably constrained in static HFB calculations
for these nuclei.

Besides EDF parameters, the effective axial-vector coupling constant $g_A$ is also included
as a model parameter in the fit. It impacts the strengths of Gamow-Teller (GT) and forbidden
transitions. For a free neutron its value is $g_A^{\text{free}} \approx 1.27$ [183], but in nuclei it is quenched
due to nuclear-medium effects and deficiencies in nuclear models; see Refs. [184, 185] for
reviews on the quenching of the axial-vector coupling. It should be noted that the quenching
in principle depends on the nucleus and transition type, and heavy systems usually require strong quenching. In this work, however, we adopt a universal $g_A$ parameter for simplicity. In previous global beta-decay studies within the Skyrme-HFB-PNFAM framework [66, 67], an empirical value $g_A = 1.0$ is used and the results are acceptable when compared with experimental data. Hence, this empirical value provides a starting point for our fits.

It is a good practice to exploit dimensionless parameters with similar variations in the $\chi^2$-optimization routine. The effective axial-vector coupling $g_A$ is already dimensionless, with its value constrained within the range of $[0, 2]$. A natural choice for the dimensionless isoscalar pairing strength is $v_0 = V_0/|V_1| \leq 0$. As for the time-odd Skyrme couplings, we transform them into dimensionless Landau-Migdal parameters that are defined by the Landau interaction, a residual-interaction form suitable for the studies of nuclear-matter properties and low-lying excitations [186, 187]. The corresponding transformation is [94, 188, 189]:

\begin{align}
  g'_0 &= N_0 \left( 2C_1^s + 2C_1^T k_F^2 + \frac{2}{3}C_1^F k_F^2 \right), \\
  g'_1 &= -2N_0C_1^T k_F^2 - \frac{2}{3}N_0C_1^F k_F^2, \\
  h'_0 &= \frac{1}{3}N_0 k_F^2 C_1^F ,
\end{align}

where $k_F = (3\pi^2 \rho_c/2)^{2/3}$ is the Fermi momentum of nuclear matter at the saturation density $\rho_c$. The normalization factor is $N_0 = 2m^* k_F/ (\pi^2 \hbar^2)$, where

\begin{equation}
  m^* = \left( \frac{2}{\hbar^2} \frac{\partial E}{\partial \tau_{00}} \right)_{\rho_{00}=\rho_c}^{-1} = \left( \frac{1}{m} + \frac{2}{\hbar^2} C_0^\tau \rho_c \right)^{-1}
\end{equation}

is the isoscalar effective mass of symmetric nuclear matter [190, 191]. One can notice that
Table 5.1: Four optimization schemes with different free and fixed parameter sets.

<table>
<thead>
<tr>
<th>Scheme</th>
<th>Free parameters</th>
<th>Fixed parameters</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>$g'<em>{0}$, $v</em>{0}$</td>
<td>$g_{A} = 1$, $g'_{1} = 0$</td>
</tr>
<tr>
<td>B</td>
<td>$g'<em>{0}$, $v</em>{0}$, $g_{A}$</td>
<td>$g'_{1} = 0$</td>
</tr>
<tr>
<td>C</td>
<td>$g'<em>{0}$, $g'</em>{1}$, $v_{0}$</td>
<td>$g_{A} = 1$</td>
</tr>
<tr>
<td>D</td>
<td>$g'<em>{0}$, $g'</em>{1}$, $v_{0}$, $g_{A}$</td>
<td>None</td>
</tr>
</tbody>
</table>

Table 5.2: GTR energies and their experimental errors (in MeV) taken from Refs. [198, 196, 199, 200] for the four nuclei selected in this work.

<table>
<thead>
<tr>
<th>No.</th>
<th>Nucleus</th>
<th>$E_{GTR}$</th>
<th>Exp error</th>
<th>No.</th>
<th>Nucleus</th>
<th>$E_{GTR}$</th>
<th>Exp error</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>$^{208}$Pb</td>
<td>15.6</td>
<td>0.2</td>
<td>3</td>
<td>$^{90}$Zr</td>
<td>8.7</td>
<td>-</td>
</tr>
<tr>
<td>2</td>
<td>$^{132}$Sn</td>
<td>16.3</td>
<td>0.6</td>
<td>4</td>
<td>$^{112}$Sn</td>
<td>8.94</td>
<td>0.25</td>
</tr>
</tbody>
</table>

$g'_{1}$ and $h'_{0}$ are purely determined by the couplings of tensor terms in the Skyrme EDF. In the fit we have $h'_{0} = 0$ as we set $C_{1}^{F} = 0$, and the stability condition of nuclear matter in the spin-isospin channel yields $g'_{1} > -3$ when $h'_{0} = 0$ [192], providing a constrain on $g'_{1}$. On the other hand, no constrain is placed on $g'_{0}$, but experiments on nuclear spin-isospin responses give a value of $g'_{0} \sim 1.6$ [193, 194, 195, 196], which is a good starting point for our fits.

In summary, model parameters considered in the calibration are $g'_{0}$, $g'_{1}$, $v_{0}$ and $g_{A}$. Table 5.1 summarizes the four optimization schemes we use, where $g'_{0}$ and $v_{0}$ are always free parameters but $g'_{1}$ and $g_{A}$ can be either fixed or free. The comparison between these schemes will answer if $g'_{1}$ and $g_{A}$ can be well constrained by data and if they should be included in the fits.

5.2 Fit observables

As shown in Tables 5.2 and 5.3, two types of observables are employed in the calibration: Gamow-Teller-resonance (GTR) energies $E_{GTR}$ and $\beta^{-}$-decay half lives $T_{1/2}$. The half lives

\footnote{It should be noted that experimental papers usually adopt the $\pi + \rho + g'$ model with a different normalization factor $1/N'_{0} = 392$ MeV $\cdot$ fm [197], so a transformation of normalization is necessary here.}
Table 5.3: Beta-minus-decay half lives and their experimental errors (in second) taken from Ref. [201] for the 25 nuclei selected in this work. Half lives are listed in ascending order.

<table>
<thead>
<tr>
<th>No.</th>
<th>Nucleus</th>
<th>$T_{1/2}$</th>
<th>Exp error</th>
<th>No.</th>
<th>Nucleus</th>
<th>$T_{1/2}$</th>
<th>Exp error</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>$^{98}$Kr</td>
<td>0.043</td>
<td>0.004</td>
<td>18</td>
<td>$^{169}$Gd</td>
<td>4.8</td>
<td>1.0</td>
</tr>
<tr>
<td>6</td>
<td>$^{58}$Ti</td>
<td>0.058</td>
<td>0.009</td>
<td>19</td>
<td>$^{156}$Nd</td>
<td>5.26</td>
<td>0.2</td>
</tr>
<tr>
<td>7</td>
<td>$^{102}$Sr</td>
<td>0.069</td>
<td>0.006</td>
<td>20</td>
<td>$^{204}$Pt</td>
<td>10.3</td>
<td>1.4</td>
</tr>
<tr>
<td>8</td>
<td>$^{82}$Zn</td>
<td>0.166</td>
<td>0.011</td>
<td>21</td>
<td>$^{74}$Zn</td>
<td>95.6</td>
<td>1.2</td>
</tr>
<tr>
<td>9</td>
<td>$^{48}$Ar</td>
<td>0.475</td>
<td>0.04</td>
<td>22</td>
<td>$^{52}$Ti</td>
<td>102</td>
<td>6</td>
</tr>
<tr>
<td>10</td>
<td>$^{60}$Cr</td>
<td>0.49</td>
<td>0.01</td>
<td>23</td>
<td>$^{180}$Yb</td>
<td>144</td>
<td>30</td>
</tr>
<tr>
<td>11</td>
<td>$^{126}$Cd</td>
<td>0.515</td>
<td>0.017</td>
<td>24</td>
<td>$^{114}$Pd</td>
<td>145.2</td>
<td>3.6</td>
</tr>
<tr>
<td>12</td>
<td>$^{114}$Ru</td>
<td>0.54</td>
<td>0.03</td>
<td>25</td>
<td>$^{242}$U</td>
<td>1008</td>
<td>30</td>
</tr>
<tr>
<td>13</td>
<td>$^{134}$Sn</td>
<td>1.05</td>
<td>0.011</td>
<td>26</td>
<td>$^{134}$Te</td>
<td>2508</td>
<td>48</td>
</tr>
<tr>
<td>14</td>
<td>$^{152}$Ce</td>
<td>1.4</td>
<td>0.2</td>
<td>27</td>
<td>$^{92}$Sr</td>
<td>9399.6</td>
<td>61.2</td>
</tr>
<tr>
<td>15</td>
<td>$^{78}$Zn</td>
<td>1.47</td>
<td>0.15</td>
<td>28</td>
<td>$^{156}$Sm</td>
<td>33840</td>
<td>720</td>
</tr>
<tr>
<td>16</td>
<td>$^{72}$Ni</td>
<td>1.57</td>
<td>0.05</td>
<td>29</td>
<td>$^{208}$Pt</td>
<td>45360</td>
<td>1080</td>
</tr>
<tr>
<td>17</td>
<td>$^{92}$Kr</td>
<td>1.84</td>
<td>0.008</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

are given in ascending order, and their logarithms $\lg T_{1/2}$ (base 10) are adopted in the fit as the half lives can vary by several orders of magnitudes.

Our data selection is similar to that of Ref. [66], but some data points have been excluded for various reasons. As for the resonance data, only GTRs of doubly-magic ($^{208}$Pb and $^{132}$Sn) and semi-magic ($^{90}$Zr and $^{112}$Sn) systems are considered, while transitional soft systems $^{76}$Ge, $^{130}$Te and $^{150}$Nd are not well described by the mean-field model and thus excluded. The GTR of $^{48}$Ca and spin-dipole resonances (SDRs) of $^{90}$Zr and $^{208}$Pb are also excluded because their experimental spectra do not exhibit clear resonance peaks (see Refs. [202, 203, 204] for the GTR of $^{48}$Ca and Refs. [205, 195] for the SDRs of $^{90}$Zr and $^{208}$Pb). As for beta-decay data, systems with possible octupole ground states ($^{148}$Ba, $^{226}$Rn) [31] are ruled out as the reflection symmetry is imposed in our calculations.
5.3 Least-squares fit

Reference [68] provides a comprehensive guide and a compilation of examples for the application of the least-squares fit in nuclear physics. Here we briefly summarize conclusions that are useful for this work. The optimal model parameters are determined through minimizing the weighted sum of squared residuals (errors):

$$\chi^2(x) = \frac{1}{n_d - n_x} \sum_{k=1}^{n_d} \varepsilon_k^2(x) = \frac{1}{n_d - n_x} \sum_{k=1}^{n_d} \left[ \frac{s_k(x) - d_k}{w_k} \right]^2,$$

where $x \in \mathbb{R}^{n_x}$ is the (column) vector constituted by model parameters, $n_x = |x|$ is the number of model parameters, $n_d$ is the total number of observables, subscript $k$ denotes the observable index, $s_k(x)$ is the model prediction, $d_k$ is the experimental value, and $w_k$ is known as the weight or adopted error. When there is more than one observable type, the weight is necessary to make the residual $\varepsilon_k = [s_k(x) - d_k]/w_k$ dimensionless. For an inexact model, there is arbitrariness on the values of weights and one can adjust them to vary the importance of different observables [206]. However, the statistical assumption we adopt for uncertainty estimation can provide a guide for the weight determination.

It is assumed that all weighted residuals $\varepsilon_k$ are independent and follow the same normal distribution with expectation 0 and variance $\sigma^2$, and the $\chi^2$ value at the optimal point $\hat{x}$ is approximately the variance $\sigma^2$. To satisfy this assumption, we should choose the weight $w_k$ close to the error of model $s_k$, including theoretical, numerical and experimental errors, and then the $\chi^2(\hat{x})$ should be approximately 1. Although each point can have a distinct weight, the points of the same type usually share the same weight as their errors are expected to be close. One can note that what actually matters is the relative weights between observable
types, because we can introduce a global scale factor $s$ (Birge factor [207]) such that

$$
\chi^2(\hat{x}) \rightarrow \tilde{\chi}^2(\hat{x}) = \chi^2(\hat{x})/s = 1, \ w_k \rightarrow \tilde{w}_k = w_k\sqrt{s}.
$$

(5.4)

Then, for consistency between weights and residual distributions, the scaled weight for a given observable type $\tilde{w}_{\text{typ}}$ should be close to

$$
r_{\text{typ}} = \sqrt{\frac{n_d}{n_{\text{typ}}(n_d - n_x)} \sum_{k \in \text{typ}} [s_k(\hat{x}) - d_k]^2},
$$

(5.5)

where $n_{\text{typ}}$ is the number of points of the given type.

With the linear expansions of weighted residuals $\varepsilon_k(x)$ around the optimal point $\hat{x}$, we can transform the nonlinear optimization problem to a linear one and employ the linear-regression framework. Rigorous mathematical discussions can be found in Ref. [208]; in the following we only list important conclusions without proof. Let $x^*$ be the true parameter vector. Then the difference $(\hat{x} - x^*)$ approximately follows a multivariate normal distribution: $\hat{x} - x^* \sim N(0, \text{Cov}(\hat{x}))$. The covariance matrix is

$$
\text{Cov}(\hat{x}) \approx \chi^2(\hat{x}) \left[ J^T(\hat{x})J(\hat{x}) \right]^{-1},
$$

(5.6)

where the $n_d \times n_x$ Jacobian matrix $J(x)$ is defined by $J_{kl} = \frac{\partial \varepsilon_k}{\partial x_l}$. There exist several approximate evaluations for the covariance matrix, but Eq. (5.6) is simpler, numerically cheaper and stabler than other choices [209]. The correlation matrix is then

$$
R_{kl} = \frac{\text{Cov}(\hat{x})_{kl}}{\sigma_k \sigma_l},
$$

(5.7)
where $\sigma_k = \sqrt{\text{Cov}(\hat{x})_{kk}}$ is the standard deviation of $\hat{x}_k$. The $(1 - \alpha)$ confidence region for $\mathbf{x}^*$ is

$$\left\{ \mathbf{x}^* \in \mathbb{R}^{n_x} : (\mathbf{x}^* - \hat{\mathbf{x}})^T \text{Cov}^{-1}(\hat{\mathbf{x}})(\mathbf{x}^* - \hat{\mathbf{x}}) \leq n_x F_{n_x, n_d-n_x, 1-\alpha} \right\}, \quad (5.8)$$

where $F_{n_x, n_d-n_x, 1-\alpha}$ is the $(1 - \alpha)$ quantile of the $F$ distribution with $n_x$ and $(n_d-n_x)$ degrees of freedom. Suppose $\mathbf{x} = (\mathbf{x}_1^T, \mathbf{x}_2^T)^T$, where $\mathbf{x}_1 \in \mathbb{R}^{n_1}$ and $\mathbf{x}_2 \in \mathbb{R}^{n_x-n_1}$; the $(1 - \alpha)$ confidence region for the parameter subset $\mathbf{x}_1^*$ is

$$\left\{ \mathbf{x}_1^* \in \mathbb{R}^{n_1} : (\mathbf{x}_1^* - \hat{\mathbf{x}}_1)^T \text{Cov}^{-1}(\hat{\mathbf{x}}_1)(\mathbf{x}_1^* - \hat{\mathbf{x}}_1) \leq n_1 F_{n_1, n_d-n_x, 1-\alpha} \right\}, \quad (5.9)$$

where $\text{Cov}(\hat{\mathbf{x}}_1)$ is the upper left $n_1 \times n_1$ submatrix of $\text{Cov}(\hat{\mathbf{x}})$. When $n_1 = 1$ we obtain the $(1 - \alpha)$ confidence interval for the $k$-th parameter: \( \left\{ x_k^* \in \mathbb{R} : |x_k^* - \hat{x}_k| \leq \sigma_k t_{n_d-n_x, 1-\alpha/2} \right\} \), where $t_{n_d-n_x, 1-\alpha/2}$ is the $(1 - \alpha/2)$ quantile of the $t$ distribution with $(n_d-n_x)$ degrees of freedom. As discussed in Ref. [191], the sensitivity matrix is

$$S = \left[J^T(\hat{x}) J(\hat{x})\right]^{-1} J^T(\hat{x}), \quad (5.10)$$

whose matrix elements are $S_{kl} \approx \frac{\partial \hat{x}_k}{\partial \epsilon_l}$, representing the variations of optimal parameters when experimental data are slightly changed. To eliminate the impact of different parameter scales, we introduce normalized model parameters $y_k = x_k / \sigma_k$; correspondingly, the normalized Jacobian matrix is $\tilde{J}_{kl}(\mathbf{y}) = \frac{\partial x_k}{\partial y_l} = \sigma_l J_{kl}(\mathbf{x})$, and the normalized sensitivity matrix is

$$\tilde{S}_{kl} = S_{kl} / \sigma_k = \left\{ \left[J^T(\tilde{y}) J(\tilde{y})\right]^{-1} J^T(\tilde{y}) \right\}_{kl} \approx \frac{\partial \tilde{y}_k}{\partial \epsilon_l}. \quad (5.11)$$

The principal component analysis (PCA) [210, 211, 212] is a useful tool to explore the
possibility of reducing the dimension of the parameter space. It has been employed to find the number of effective parameters in nuclear mass models [69]. The PCA provides an orthogonal transformation among model parameters such that the first few new parameters can explain most variation in data. In the PCA, we calculate the singular value decomposition (SVD) of the normalized Jacobian matrix $\hat{J}$ at the optimal point $\hat{y}$ to obtain principal components:

$$
\sum_l J(\hat{y})_kl\hat{V}_{lm} = s_m\hat{U}_{km}, \quad s_1 \geq s_2 \geq \cdots \geq s_{nx} \geq 0,
$$

(5.12)

where $\hat{V} \in \mathbb{R}^{nx \times nx}$ and $\hat{U} \in \mathbb{R}^{nd \times nd}$ are orthogonal matrices. The transformation $\hat{V}$ defines a new set of parameters as $z_m = \sum_l y_l \hat{V}_{lm}$, and the singular value $s_m$ indicates the relevance of $z_m$ in the fit. One can prove that the approximate covariance matrix of the new parameter set is $\text{Cov}(\hat{z})_{kl} = [\hat{V}^T \text{Cov}(\hat{y})\hat{V}]_{kl} \propto s_k^{-2}\delta_{kl}$. Therefore, $\hat{z}_k$ and $\hat{z}_l$ ($k \neq l$) are uncorrelated, and the larger the singular value $s_k$ is, the better the parameter $z_k$ is constrained (the less soft the $\chi^2$ surface is in the corresponding direction). To determine the number of effective model parameters, we define a cumulative quantity:

$$
S_m = \frac{\sum_{k=1}^{m} s_k^2}{\sum_{k=1}^{nx} s_k^2},
$$

(5.13)

which is the percentage of the variation in data that first $m$ components (columns) of matrices $\hat{U}$ and $\hat{V}$ account for. The number of effective parameters is the minimum $m$ that satisfies $S_m > S_{th}$. A typical threshold value is $S_{th} = 0.99$, and the corresponding parameter set $\{z_1, z_2, \cdots, z_m\}$ explains 99% variation.
5.4 Numerical details

The program HFBTHO is utilized to calculate the HFB ground states of parent nuclei whose GTRs or beta-decay half lives are fit observables. To be consistent with the fit of UNEDF1-HFB, the number of HO shells is \( N_{\text{HO}} = 20 \). The spherical symmetry is assumed in the ground states of \(^{208}\text{Pb}\), \(^{90}\text{Zr}\) and \(^{132}\text{Sn}\), while other HFB calculations start from a series of kickoff deformations \( \beta_2 = -0.4 + 0.1k \) (\( k = 0, 1, \cdots, 8 \)) with the HO-basis deformation taking the same \( \beta_2 \) value, and then the minimum-energy point corresponds to the ground state.

Based on HFB results, the PyNFAM program calls the PNFAM routine to compute \( \beta^- \) transition strength distributions, and then evaluates the GTR energy or beta-decay half life; see Sec. 2.3 for details. In GTR calculations, the resonance is searched on a horizontal line in the complex \( \omega \) plane with a constant imaginary part (smearing width) \( \Gamma = 0.5 \) MeV, while the grid spacing of the real part is 0.4 MeV. In half-life calculations, 60 sample points are used for the Gauss-Legendre integration on the full circular contour (2.73); PNFAM runs on half of the points are actually needed thanks to the relation \( S(\omega^*) = S^*(\omega) \). In addition, a 15-point Chebyshev grid is adopted for the rational interpolation of phase-space factors.

The least-squares fit routine we choose is POUNDERS (practical optimization using no derivatives for sums of squares) [213] in PETSc/TAO [214, 215, 216]. It is first employed for the fit of Skyrme parametrization UNEDF0 [191], and shown to be robust and efficient among derivative-free optimizers [217, 218]. In the POUNDERS algorithm, weighted residuals \( \varepsilon_i \) are modeled by a quadratic function within a trust region to guide the \( \chi^2 \) minimization. To find the statistical quantities discussed in Sec. 5.3, the Jacobian matrix \( J \) is approximately
Table 5.4: Optimal parameter values obtained from fits with GTR energies only. Standard deviations are given in brackets. The first letter of the fit indicates the scheme (see Table 5.1). Schemes B and D are absent as $g_A$ does not affect the GTR energy. Root-mean-square errors (RMSEs) are also presented for comparison.

<table>
<thead>
<tr>
<th>Fit</th>
<th>$g'_0$</th>
<th>$g'_1$</th>
<th>$v_0$</th>
<th>RMSE (MeV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A-GTR</td>
<td>1.60553 (0.025)</td>
<td>0</td>
<td>−1.73430 (0.561)</td>
<td>0.118</td>
</tr>
<tr>
<td>C-GTR</td>
<td>1.47434 (0.779)</td>
<td>0.38129 (2.353)</td>
<td>−1.70663 (0.699)</td>
<td>0.116</td>
</tr>
</tbody>
</table>

Table 5.5: Similar to Table 5.4 but with beta-decay half-life data only. Optimization schemes C and D with $g'_1$ as a free parameter are absent because of severe numerical instability.

<table>
<thead>
<tr>
<th>Fit</th>
<th>$g'_0$</th>
<th>$v_0$</th>
<th>$g_A$</th>
<th>RMSE</th>
</tr>
</thead>
<tbody>
<tr>
<td>A-β</td>
<td>2.42992 (0.503)</td>
<td>−1.06817 (0.169)</td>
<td>1</td>
<td>0.924</td>
</tr>
<tr>
<td>B-β</td>
<td>−0.07262 (0.212)</td>
<td>−1.06339 (0.274)</td>
<td>0.12243 (0.059)</td>
<td>0.718</td>
</tr>
</tbody>
</table>

evaluated via the central difference formula:

$$J_{kl}(\hat{x}) = \frac{\partial \varepsilon_k}{\partial x_l} \bigg|_{\hat{x}} \approx \frac{\varepsilon_k(\hat{x} + d e_l) - \varepsilon_k(\hat{x} - d e_l)}{2d}, \quad (5.14)$$

where $e_l \in \mathbb{R}^{nx}$ denotes a standard unit vector with a 1 in the $l$-th component and zeros elsewhere, and we choose $d = 10^{-3}$ in our calculations. A Python interface has been developed to connect the PyNFAM program and POUNDERS routine. This interface is also capable of performing calculations for a given group of parameter vectors, and the results can then be used as training or test data to build a physics-model emulator for the Bayesian calibration.
5.5 Results and discussions

5.5.1 Optimizations involving one observable type

In order to examine the role of different data types and to estimate their weights for a fit with the full data set, we first carry out fits with only one type of data and results are shown in Tables 5.4 (GTR data) and 5.5 (beta-decay half-life data). Optimization schemes B and D are not considered in Table 5.4 since $g_A$ does not affect the GTR location, while schemes C and D are not listed in Table 5.5 because in these schemes model parameters are pushed to unphysical regions where severe numerical instability is observed.

In Fit A-GTR, we see that $g'_0$ is well constrained by the GTR data but $v_0$ is not, since only the GTR locations of semi-magic $^{90}$Zr and $^{112}$Sn weakly depend on $v_0$. On the other hand, in Fit A-$\beta$ the half-life data yield a large uncertainty of $g'_0$ and drive it far from the experimentally accepted value, while the standard deviation of $v_0$ is small. Therefore, when both data types are considered in A, $g'_0$ will be primarily determined by the GTR data while $v_0$ be well constrained by the half-life data. As shown in Table 5.4, when $g'_1$ is included as a free parameter (Fit C-GTR), the root-mean-square error (RMSE) is barely reduced and the uncertainty of $g'_1$ is quite large; in addition, $g'_0$ and $g'_1$ are strongly correlated with a correlation coefficient of $-0.999$. Therefore, $g'_1$ should not be considered in the fit with the GTR data only. Fit C-$\beta$ takes $g_A$ into account and yields unphysical values of $g'_0$ and $g_A$, far away from those determined in previous studies (see Sec. 5.1); but the RMSE significantly decreases in this fit and it is difficult to say whether $g_A$ should be freely varied.
Figure 5.1: Values of $\tilde{w}$ and $r$ for fits with a series of relative weights $\gamma = w_{\text{GTR}}/w_\beta$. The optimization scheme A in Table 5.1 is used and all the data in Tables 5.2 and 5.3 are included in the fits. Actual calculations are performed with $w_\beta = 1$ and $w_{\text{GTR}} = \gamma$.

### 5.5.2 Weight determination

As discussed in Sec. 5.3, only relative weights between different observable types need to be determined, and the main idea is to match the values of $\tilde{w}$ and $r$ defined in Eqs. (5.4) and (5.5). Let $w_{\text{GTR}}$ and $w_\beta$ be the weights of GTR energies $E_{\text{GTR}}$ and logarithms of beta-decay half lives $\lg T_{1/2}$, respectively. The relative weight is then the ratio $\gamma = w_{\text{GTR}}/w_\beta = \tilde{w}_{\text{GTR}}/\tilde{w}_\beta$. Using optimization scheme A, we perform a series of fits with a fixed value of $w_\beta = 1$ and various values of $w_{\text{GTR}}$, and Fig. 5.1 shows the values of $\tilde{w}$ and $r$ of both data types. One can notice that the requirement $\tilde{w} \approx r$ is fulfilled at around $\gamma = 0.15$ MeV, where $\tilde{w}_{\text{GTR}} \approx 0.15$ MeV and $w_\beta \approx 1.03$. On the other hand, the quantity $\tilde{w}$ is supposed to include theoretical, numerical and experimental errors, and thus $\tilde{w}_{\text{GTR}}$ cannot be that small given the experimental errors in Table 5.2 (the experimental errors of $\lg T_{1/2}$ is smaller than 1 and thus negligible). Based on these two considerations, we can make a trade-off and choose $\gamma = 0.3$ MeV, which yields $\tilde{w}_{\text{GTR}} \approx 0.29$ MeV and $\tilde{w}_\beta \approx 0.96$. Figure 5.2 displays
the RMSEs of the two types of data for the same series of fits shown in Fig. 5.1, and we see that the fit of $\gamma = 0.3$ MeV produces a good balance between the RMSEs of the two types. Other optimization schemes produce similar results and are thus not discussed here.

5.5.3 Optimizations involving two observable types

Table 5.6 presents optimal parameters obtained from the four optimization schemes with weights $w_{GTR} = 0.3$ MeV and $w_{\beta} = 1$ (relative weight $\gamma = 0.3$ MeV). As expected in Sec.

The RMSEs of two types of data (GTR energies $E_{GTR}$ and logarithms of beta-decay half lives $\log T_{1/2}$), obtained from the fits shown in Fig. 5.1. Relative weights $\gamma$ (in MeV) are represented by the colors of square markers.

Table 5.6: Optimal parameter values obtained from fits with weights $w_{GTR} = 0.3$ MeV and $w_{\beta} = 1$ (relative weight $\gamma = 0.3$ MeV). Standard deviations are given in brackets, and the $\chi^2$ values are also presented for comparison.

<table>
<thead>
<tr>
<th>Fit</th>
<th>$g'_0$</th>
<th>$g'_1$</th>
<th>$v_0$</th>
<th>$g_A$</th>
<th>$\chi^2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>1.59560 (0.039)</td>
<td>0</td>
<td>$-0.99993$ (0.178)</td>
<td>1</td>
<td>25.057</td>
</tr>
<tr>
<td>B</td>
<td>1.59184 (0.034)</td>
<td>0</td>
<td>$-1.19745$ (0.179)</td>
<td>0.50345 (0.143)</td>
<td>19.385</td>
</tr>
<tr>
<td>C</td>
<td>1.73245 (0.820)</td>
<td>$-0.37034$ (2.143)</td>
<td>$-0.99920$ (0.183)</td>
<td>1</td>
<td>25.019</td>
</tr>
<tr>
<td>D</td>
<td>2.72206 (0.422)</td>
<td>$-2.54125$ (0.781)</td>
<td>$-1.23511$ (0.179)</td>
<td>0.41168 (0.132)</td>
<td>17.788</td>
</tr>
</tbody>
</table>

Figure 5.2: Relation between the RMSEs of two types of data (GTR energies $E_{GTR}$ and logarithms of beta-decay half lives $\log T_{1/2}$), obtained from the fits shown in Fig. 5.1. Relative weights $\gamma$ (in MeV) are represented by the colors of square markers.
5.5.1, both $g'_0$ and $v_0$ are well constrained in Fit A. In Fit B, where $g_A$ becomes a free parameter, the value of $\chi^2$ is dramatically reduced as decay half lives are better fitted. In Fit C, however, the introduction of free $g'_1$ does not improve the fit much, and both $g'_0$ and $g'_1$ become badly constrained. When both $g'_1$ and $g_A$ are freely varied in Fit D, the optimal values of $g'_0$ and $g'_1$ are pushed far from other fits with large uncertainties. Based on the results in Table 5.6, we conclude that $g'_1$ cannot be constrained by our data and should not be included as a free parameter in the fit.

Figure 5.3 displays individual weighted residuals $\varepsilon_k$ given by the four fits in Table 5.6, which are consistent with the $\chi^2$ values shown in the table. One can notice a general trend that the errors of decay half lives are significantly larger for long-lived nuclei (those with large data-point numbers). These nuclei usually have a small beta-decay $Q$ number, and Refs. [66, 67] show that their leptonic phase-space factors incur greater theoretical errors. In addition, a small $Q$ value also means that the radius of the integration contour (2.73) is small.
Table 5.7: Correlation matrix (5.7) obtained from Fit A in Table 5.6.

<table>
<thead>
<tr>
<th></th>
<th>$g_0'$</th>
<th>$v_0$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$g_0'$</td>
<td>1.000</td>
<td></td>
</tr>
<tr>
<td>$v_0$</td>
<td></td>
<td>−0.135 1.000</td>
</tr>
</tbody>
</table>

Table 5.8: Similar to Table 5.7 but for Fit B given in Table 5.6.

<table>
<thead>
<tr>
<th></th>
<th>$g_0'\</th>
<th>v_0 \</th>
<th>g_A</th>
</tr>
</thead>
<tbody>
<tr>
<td>$g_0'$</td>
<td>1.000</td>
<td></td>
<td></td>
</tr>
<tr>
<td>$v_0$</td>
<td></td>
<td>−0.097 1.000</td>
<td></td>
</tr>
<tr>
<td>$g_A$</td>
<td></td>
<td></td>
<td>−0.004 0.615 1.000</td>
</tr>
</tbody>
</table>

and a very limited number of residues are enclosed, so one peak that is poorly determined in the calculation can lead to a large error in the half life. Hence, one may consider assigning different weights for short- and long-lived nuclei in future fits.

5.5.4 Number of effective model parameters

The correlation matrix (5.7) can give us hints on the number of effective model parameters as two well correlated variables should be combined into one effective parameter. Tables 5.7, 5.8, 5.9 and 5.10 present the correlation matrices of the four fits given in Table 5.6. One can note that $g_0'$ and $g_1'$ are highly correlated in Fits C and D, similar to the case of Fit C-GTR in Table 5.4. Also, $v_0$ and $g_A$ are moderately correlated in Fits B and D as both of them depend heavily on the half-life data.

As discussed in Sec. 5.3, more concrete determination of effective parameters is provided

Table 5.9: Correlation matrix (5.7) obtained from Fit C in Table 5.6. Correlations larger than 0.8 in absolute value are marked in italics.

<table>
<thead>
<tr>
<th></th>
<th>$g_0'$</th>
<th>$g_1'$</th>
<th>$v_0$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$g_0'$</td>
<td>1.000</td>
<td></td>
<td></td>
</tr>
<tr>
<td>$g_1'$</td>
<td></td>
<td>−0.999 1.000</td>
<td></td>
</tr>
<tr>
<td>$v_0$</td>
<td></td>
<td>0.057 −0.064 1.000</td>
<td></td>
</tr>
</tbody>
</table>
Table 5.10: Similar to Table 5.9 but for Fit D given in Table 5.6.

<table>
<thead>
<tr>
<th>$D$</th>
<th>$g'_0$</th>
<th>$g'_1$</th>
<th>$v_0$</th>
<th>$g_A$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$g'_0$</td>
<td>1.000</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$g'_1$</td>
<td>−0.996</td>
<td>1.000</td>
<td></td>
<td></td>
</tr>
<tr>
<td>$v_0$</td>
<td>−0.259</td>
<td>0.256</td>
<td>1.000</td>
<td></td>
</tr>
<tr>
<td>$g_A$</td>
<td>−0.500</td>
<td>0.507</td>
<td>0.622</td>
<td>1.000</td>
</tr>
</tbody>
</table>

Figure 5.4: Squared singular values (in descending order) obtained from Eq. (5.12), the SVD of the normalized Jacobian matrix $\hat{J}$, for the four fits presented in Table 5.6.

by the PCA. Figure 5.4 displays squared singular values of normalized Jacobian matrices $\hat{J}$ obtained from Eq. (5.12) for the four fits listed in Table 5.6. In Fits A and B, the number of effective parameters is the same as that of free parameters. In Fits C and D, the 99% threshold is reached when the first 2 and 3 singular values are included in the cumulative quantity $S_m$ (5.13), respectively. The parameter corresponding to the smallest singular value should thus be removed from the set of effective parameters in Fits C and D.

Figure 5.4 shows the squared principal components $\hat{V}^2_{kl}$ corresponding to the first ($l = 1$) and second ($l = 2$) largest singular values in the four fits; they represent how the two most effective parameters determined by the PCA are constructed from the normalized parameter
Figure 5.5: Squared amplitudes of the first (left) and second (right) principal components (column vectors of $\hat{V}$) that correspond to the first and second largest singular values, for the four fits presented in Table 5.6. They are obtained from Eq. (5.12), the SVD of the normalized Jacobian matrix $\hat{J}$.

set $\{y_k\}$. In Fit A, $g'_0$ and $v_0$ are equally important and democratically contribute to $\hat{V}_{k1}^2$ and $\hat{V}_{k2}^2$. In Fit B, the first principal component is constituted by $v_0$ and $g_A$ that are mainly related to the half-life data, while the second component is purely $g'_0$ that has almost no correlation with other parameters. In Fits C and D, the first principal component is composed of $g'_0$ and $g'_1$ with almost the same amplitudes because of their strong correlation.

5.5.5 Sensitivity analysis

The matrix elements of the normalized sensitivities (5.11) are shown in Fig. 5.6 in absolute value for the four fits given in Table 5.6. The sensitivity quantifies the variations of parameter
Figure 5.6: Normalized sensitivity matrices (5.11) in absolute value for the fits presented in Table 5.6, with weights \( w_{\text{GTR}} = 0.3 \text{ MeV} \) and \( w_\beta = 1 \) (relative weight \( \gamma = 0.3 \text{ MeV} \)). The labeling of data points is consistent with that in Tables 5.2 and 5.3. The vertical dashed line separates the points of GTR energies and the logarithms of beta-decay half lives.

values when one observable is changed. In Fits A and B the value of \( g'_0 \) is highly sensitive to the GTR data, while in Fits C and D the introduction of \( g'_1 \) as a free parameter spreads the sensitivity distribution of \( g'_0 \). Furthermore, in Fits C and D the sensitivities of \( g'_0 \) and \( g'_1 \) on the same point are close to each other in absolute value but have opposite signs, indicating the strong anticorrelation between these two parameters. In all the fits, \( v_0 \) is equally sensitive to all the data, except the half lives of \(^{52}\text{Ti}\) and \(^{92}\text{Sr}\). These two nuclei are long-lived and it is unclear whether there are any underlying physics problems attributed to their strong sensitivities. As expected, the sensitivity of \( g_A \) to the half-life data is greater than that to
the GTR data, and the point of $^{52}$Ti still provides the largest sensitivity.

5.6 Summary

In this chapter we calibrate time-odd Skyrme couplings (Landau parameters), the isoscalar pairing strength and the effective axial-vector coupling in the Skyrme-HFB-PNFAM model for beta-decay calculations. The calibration is carried out within the $\chi^2$-optimization framework, i.e., in the frequentist picture. Selected GTR energies and $\beta^-$-decay half lives are included as experimental data in the fits, and we see that different parameters depend differently on the two types of data. The weights of the two types are carefully determined based on the statistical assumption and experimental errors. The comparison between the results of various schemes shows that $g_{1T}'$, the parameter related to the tensor term, cannot be well constrained by data. The correlation matrices and principal component analysis help us reduce the dimension of the effective parameter space.

Based on current results, there are a couple of possible extensions for future works. First, the density dependence of $C_1^s$ should be taken into account in the fit to investigate its role in GTR and beta-decay calculations. Second, as noted in Ref. [219] the two-body weak current plays a crucial role in the quenching of $g_A$, and Ref. [220] has included it in the PNFAM framework. In this new model, low-energy constants involved in the chiral effective field theory becomes model parameters to calibrate while $g_A$ should take the value of $g_A^{\text{free}}$. Moreover, we consider adding new data or including other types of data (e.g., the GTR strength and beta-decay rate of a given order) to better constrain model parameters. One should notice that in the total decay rate, contributions from various transitions can compensate each other; thus, the total half life may not well constrain some model parameters.
and individual transitions can provide more information.

In addition to the $\chi^2$ optimization, we are also working on the Bayesian model calibration with the help of the Kennedy-O’Hagen (KOH) framework [221]. The Bayesian framework avoids the assumption employed in the $\chi^2$ optimization and directly produces the posterior distributions of model parameters through Monte Carlo sampling. An emulator is necessary for such sampling as the physics model we are using is not fast enough to finish sampling in a reasonable time. Our choice is the Gaussian-process (GP) emulator [222], a widely used statistical model for emulation. The Python toolkit discussed in this chapter can also be conveniently utilized to generate data for emulator building.
Chapter 6

New HFB solver HFBFFT in three-dimensional coordinate space

This chapter describes the numerical implementations and benchmarks of HFBFFT, a new HFB solver for even-even nuclei in the 3D coordinate space. The theoretical framework of the new solver has been discussed in Sec. 2.2.2, and this chapter starts from its numerical implementation (Sec. 6.1). Then the benchmarks of the new solver against well-established HFB solvers are discussed in Sec. 6.2, with a brief summary presented in Sec. 6.3. The material discussed in this chapter has recently been published in Ref. [223].

6.1 Numerical framework

6.1.1 Numerical realization on a grid

HFBFFT is adapted from the HF+BCS solver Sky3D, whose numerical framework has been presented in Refs. [81, 122, 123]. This section briefly summarizes the numerical representation of densities, fields and wave functions in the two solvers, and discusses how we correct the problem of Hermiticity violation.

All densities, fields and wave functions in HFBFFT are defined on a 3D Cartesian grid. Grid points along one ($x$, $y$ or $z$) direction are equidistant, but one can in principle choose
different numbers of points and different grid spacings in different directions. In our calculations we use identical geometries along the three directions. For simplicity, the numerical framework will be explained for the 1D case in the following; one can straightforwardly generalize it to the 3D case.

The grid points in the $x$ direction are $x_\nu = \left(-\frac{N_x+1}{2} + \nu\right) \delta x$, $\nu = 1, \ldots, N_x$, where $N_x$ is the (even) number of grid points and $\delta x$ is the grid spacing. On a grid, the action of a local operator $U(x)$ on a wave function $\psi(x)$ is a simple multiplication, i.e., $U(x_\nu)\psi(x_\nu)$. The action of the momentum operator requires numerical derivatives defined in the Fourier space. The Fourier technique has been shown to be precise and advantageous for large grids [80]. In the Fourier space, the discrete grid points $k_n$ are defined as:

\[
k_n = \begin{cases} 
(n - 1)\delta k, & n = 1, \ldots, \frac{N_x}{2} \\
(n - N_x - 1)\delta k, & n = \frac{N_x}{2} + 1, \ldots, N_x
\end{cases}, \quad \delta k = \frac{2\pi}{N_x\delta x}. \tag{6.1}
\]

A coordinate-space wave function $\psi(x_\nu)$ is connected to its Fourier-space counterpart $\tilde{\psi}(k_n)$ via the discrete Fourier transform and its inverse

\[
\tilde{\psi}_n = \sum_{\nu=1}^{N_x} \exp(-ik_n x_\nu) \psi_\nu, \tag{6.2a}
\]

\[
\psi_\nu = \frac{1}{N_x} \sum_{n=1}^{N_x} \exp(i k_n x_\nu) \tilde{\psi}_n, \tag{6.2b}
\]

where $\psi_\nu \equiv \psi(x_\nu)$, $\tilde{\psi}_n \equiv \tilde{\psi}(k_n)$. The discrete Fourier transform and its inverse can be efficiently evaluated through the FFT routine provided by the FFTW3 library [224]. One should note that the use of the Fourier space indicates that the wave function $\psi$ is periodic, i.e., $\psi(x + N_x \delta x) = \psi(x)$. The long-range Coulomb field is not compatible with periodicity,
so we deal with it in the same way as Sky3D, i.e., solve Poisson’s equation for an isolated charged distribution [81, 225, 226]. The appropriate integration method that works with the FFT technique and the periodic boundary condition is the trapezoidal rule

\[
\int_{-\frac{N_x}{2}\delta x}^{\frac{N_x}{2}\delta x} dx f(x) \approx \sum_{\nu=1}^{N_x} f(x\nu)\delta x,
\]

(6.3)

where all the points have equal weights.

Algorithm 1 presents the numerical evaluation of \( \frac{d^m\psi}{dx^m} \) in the FFT scheme, while algorithm 2 shows the method for the calculation of position-varying differentiation \( \frac{d}{dx} \left[ B(x) \frac{d\psi}{dx} \right] \). The position-varying derivative in the HF mean field \( \hat{h} \) comes from the functional term \( C_i^T \rho t t_3 \tau t_3 \).

Mathematical details about the two algorithms can be found in Ref. [227]. Here we only discuss issues related to the point \( k_{N_x/2+1} \) and the product rule.

One should note that \( k_{N_x/2+1} \) in principle can be \( \pm \frac{N_x}{2}\delta k \). This arbitrariness does not impact the transforms (6.2) but leads to different \( \psi^{(m)} \) when \( m \) is odd. We can equally split \( \tilde{\psi}(k_{N_x/2+1}) \) between positive and negative momenta, and they will cancel each other in the final result of an odd-order derivative, which is equivalent to setting \( \tilde{\psi}(k_{N_x/2+1}) = 0 \) in Algorithm 1. On the one hand, this ensures that the derivative of a real-valued function is still real; on the other hand, it means that the second derivative is not equivalent to two consecutive first derivatives in this framework.

As noted in Refs. [123, 227], the FFT-based differentiation is not compatible with the

---

**Algorithm 1** Compute the 1D differentiation \( \psi^{(m)} \equiv \frac{d^m\psi}{dx^m}, \ m = 1, 2, 3, \ldots \).

1: Compute Fourier transform \( \tilde{\psi}_n = \text{FFT}[\psi_\nu] \) via Eq. (6.2a).
2: If \( m \) is odd, set \( \tilde{\psi}_{N_x/2+1} = 0 \). Then compute \( \tilde{\psi}^{(m)}_n = (i k_n)^m \tilde{\psi}_n \).
3: Compute inverse Fourier transform \( \psi^{(m)}_\nu = \text{FFT}^{-1}[\tilde{\psi}^{(m)}_n] \) via Eq. (6.2b).
Algorithm 2 Compute the 1D position-varying differentiation $\chi \equiv \frac{d}{dx} \left[ B(x) \frac{d\psi}{dx} \right]$.

1: Compute Fourier transform $\tilde{\psi}_n = \text{FFT}[\psi_n]$ via Eq. (6.2a).
2: Save $\tilde{\psi}_{N_x/2+1} \rightarrow \tilde{\Psi}$ where $\tilde{\Psi}$ is a temporary variable.
3: Set $\tilde{\psi}_{N_x/2+1} = 0$ and compute $\tilde{\psi}^{(1)}_n = i k_n \tilde{\psi}_n$ (see Algorithm 1).
4: Compute inverse Fourier transform $\psi^{(1)}_n = \text{FFT}^{-1}[\tilde{\psi}^{(1)}_n]$ via Eq. (6.2b).
5: Calculate $\phi_\nu = B_\nu \psi^{(1)}_\nu$ with $B_\nu = B(x_\nu)$, then compute $\tilde{\phi}_n = \text{FFT}[\phi_\nu]$.
6: Compute $\tilde{\phi}^{(1)}_n = i k_n \tilde{\phi}_n$ and set $\tilde{\phi}^{(1)}_{N_x/2+1} = -\frac{1}{N_x} \left( \sum_{\nu=1}^{N_x} B_\nu \right) \left( \frac{N_x}{2} \delta k \right)^2 \tilde{\Psi}$.
7: Compute inverse Fourier transform $\chi_\nu = \text{FFT}^{-1}[\tilde{\phi}^{(1)}_n]$.

product rule. However, in Sky3D the position-varying derivative is still computed via the product rule

\[
\frac{d}{dx} \left[ B(x) \frac{d\psi}{dx} \right] = \frac{dB}{dx} \frac{d\psi}{dx} + B \frac{d^2\psi}{dx^2}, \tag{6.4}
\]

which leads to Hermiticity violation. To restore Hermiticity, one should compute $\frac{d}{dx} \left[ B \frac{d\psi}{dx} \right]$ with two consecutive first derivatives, but special treatment must be performed for the point $k_{N_x/2+1}$ to ensure that we return to Algorithm 1 with $m = 2$ when $B = 1$. As suggested in Ref. [227], we keep the term $\tilde{\psi}_{N_x/2+1}$ in the two first derivatives and then average the results of $k_{N_x/2+1} = \pm \frac{N_x}{2} \delta k$ for the symmetry in the Fourier space. One can prove that this treatment gives Algorithm 2 (Algorithm 3 in [227]), the algorithm implemented in HFBFFT.

6.1.2 Iteration scheme in coordinate space

The self-consistent HFB equations (2.39, 2.40, 2.43) in the canonical basis are solved via the damped gradient iteration [76, 78, 79, 80, 81]. The main iteration scheme is presented in Algorithm 3. The iteration starts from a number of HF+BCS steps (default is 30 HF+BCS steps), and the HF+BCS calculation is initialized by a 3D HO wave function. The initial HO potential can be spherical, axially deformed or triaxially deformed, but triaxiality is not
considered in examples shown this work.

Both Sky3D and HFBFFT are parallelized with OpenMP and MPI. The parallelization design for Sky3D can be found in Ref. [122], and it is adapted in HFBFFT to perform scalable HFB calculations. In steps 1 and $4 \sim 6$, each thread / process is responsible for a subset of s.p. wave functions. Matrix operations in steps $7 \sim 9$ and 11 are performed with BLAS [131] + LAPACK [228] (sequential or OpenMP) or ScaLAPACK [229] (MPI or OpenMP/MPI hybrid) libraries. Steps 2, 3 and 10 are not parallelized, i.e., all the threads / processes do the same work at the same time.

6.1.3 Sub-iteration scheme in the configuration space

To accelerate the convergence, we develop the sub-iteration scheme (step 11 in Algorithm 3) in which damped gradient iterations are performed in the configuration space. In the sub-iteration step, the program tries to find a unitary transformation among canonical s.p. states and a new set of occupation amplitudes to further minimize the HFB energy. Since the number of s.p. states included in the calculation is much less than the number of grid points (several hundreds vs. at least $10^4$), one iteration in the configuration space is much less computationally expensive than that on a full 3D grid, and many iterations can be performed in the sub-iteration scheme to achieve fast convergence.

In the configuration space, all the s.p. wave functions are expanded in some basis \( \{ \varphi_n \} \) as

\[
\psi_{\alpha} = \sum_{n=1}^{\Omega} \varphi_n c_{n\alpha},
\]  

(6.12)

and we choose an expansion basis such that \( c_{n\alpha}^{(0)} = \delta_{n\alpha} \) when entering the sub-iteration.
Algorithm 3 Damped gradient iteration scheme of HFBFFT. For simplicity we do not distinguish proton and neutron states.

1: Given canonical s.p. states and their occupation fractions \( \{ \psi_\alpha, v_\alpha, \alpha = 1, ..., \Omega \} \), compute local densities \( \rho, \tau, J \) and \( \bar{\rho} \) on the grid.

2: Linear mix new densities with old ones for better convergence:

\[
\varrho^{(\text{new})} = (1 - \gamma)\varrho^{(\text{old})} + \gamma\varrho_\psi, \quad \varrho = \rho, \tau \text{ or } \bar{\rho},
\]

where subscript \( \psi \) denotes the density directly computed from wave functions and \( \gamma \) is the mixture ratio whose default value is 0.2.

3: Calculate HF mean field \( \hat{h} \) and pairing field \( \hat{\tilde{h}} \) based on densities.

4: Apply \( \hat{h} \) and \( \hat{\tilde{h}} \) on all the wave functions \( \psi_\alpha \):

\[
\hat{h}\psi_\alpha \rightarrow \Psi_\alpha, \quad \hat{\tilde{h}}\psi_\alpha \rightarrow \tilde{\Psi}_\alpha,
\]

5: Compute canonical s.p. energies and pairing gaps:

\[
h_{\alpha\alpha} = \langle \psi_\alpha | \Psi_\alpha \rangle, \quad \tilde{h}_{\alpha\alpha} = \langle \psi_\alpha | \tilde{\Psi}_\alpha \rangle.
\]

6: Evaluate the action of the generalized Hamiltonian \( \hat{\mathcal{H}} \) on \( \psi_\alpha \) and overwrite \( \Psi_\alpha \):

\[
\hat{\mathcal{H}}_\alpha \psi_\alpha = v_\alpha^2 \Psi_\alpha + u_\alpha v_\alpha \tilde{\Psi}_\alpha \rightarrow \Psi_\alpha.
\]

7: Compute

\[
\Psi_\alpha - \sum_\beta \psi_\beta \lambda_{\beta\alpha} \rightarrow \Psi_\alpha,
\]

where the matrix of Lagrange multipliers is taken into account.

8: Apply the damping operator \( \hat{\mathcal{D}} \) and orthonormalization \( \hat{\mathcal{O}} \)

\[
\psi_\alpha^{(\text{new})} = \hat{\mathcal{O}} \left\{ \psi_\alpha - \hat{\mathcal{D}}\Psi_\alpha \right\}, \quad \hat{\mathcal{D}} = \frac{x_0}{v_\alpha^2 (\hat{T} + E_0) + \frac{1}{2} u_\alpha v_\alpha \hat{h}_0},
\]

where \( x_0 \) and \( E_0 \) are adjustable numerical parameters, and \( \hat{h}_0 = \max \left[ \hat{h}_n(\mathbf{r}), \hat{h}_p(\mathbf{r}) \right] \). Empirical values \( x_0 = 0.45, E_0 = 100 \text{ MeV} \) are employed.

9: Reevaluate the action of the generalized Hamiltonian and compute the matrix of Lagrange multipliers

\[
\lambda_{\beta\alpha} = \frac{\langle \psi_\beta | \hat{\mathcal{H}}_\alpha | \psi_\alpha \rangle + \langle \psi_\alpha | \hat{\mathcal{H}}_\beta | \psi_\beta \rangle^*}{2}.
\]

10: With new \( h_{\alpha\alpha} \) and \( \tilde{h}_{\alpha\alpha} \), compute new occupations amplitudes \( v_\alpha \) via Eq. (2.44).

11: Perform iterations in the configuration space (see Sec. 6.1.3).

12: If convergence is achieved (see Sec. 6.1.4), exit the iteration; otherwise, return to step 1.
The corresponding damped gradient iteration is

\[
\langle \text{new} \rangle = \hat{O} \left\{ c_{n\alpha} - \frac{\delta}{h_{nn} - h_{11} + E_0} \left[ \sum_m \mathcal{H}_{\alpha,nm} c_{m\alpha} - \sum_{\beta} c_{n\beta} \lambda_{\beta\alpha} \right] \right\},
\]

\[
= \hat{O} \left\{ c_{n\alpha} - \frac{\delta}{h_{nn} - h_{11} + E_0} \sum_{\beta} c_{n\beta} \lambda_{\beta\alpha} \right\},
\]

(6.13)

where \( \mathcal{H}_{\alpha,nm} = \langle \varphi_n | \hat{H}_\alpha | \varphi_m \rangle \), \( \lambda_{\beta\alpha} = \frac{1}{2} \sum_{mn} c_{n\beta}^* (\mathcal{H}_{\alpha,nm} + \mathcal{H}_{\beta,nm}) c_{m\alpha} \) and

\[
\lambda_{\beta\alpha} = \frac{1}{2} \langle \psi_\beta | \hat{H}_\alpha - \hat{H}_\beta | \psi_\alpha \rangle = \frac{1}{2} \sum_{mn} c_{n\beta}^* \langle \varphi_n | \hat{H}_\alpha - \hat{H}_\beta | \varphi_m \rangle c_{m\alpha}.
\]

(6.14)

Default values of \( E_0 \) and \( \delta \) are 10 MeV and 2, respectively. The gap equation (2.43) is solved between two damped gradient steps (6.13), but the HF Hamiltonian \( \hat{h} \) and pairing potential \( \hat{\tilde{h}} \) are not updated in the sub-iteration scheme. The best combination of iterations in coordinate and configuration spaces should be determined by numerical experiments; the default choice is to perform 100 iterations in the configuration space between two coordinate-space steps.

6.1.4 Convergence criterion

In a gradient-descent framework, a natural choice for the convergence check is the norm of the gradient vector. The \( \alpha \)-th component of the gradient vector in the coordinate space is

\[
\mathcal{H}_\alpha \psi_\alpha - \sum_\beta \psi_\beta \lambda_{\beta\alpha},
\]

whose projection in the configuration space is given by

\[
\langle \psi_\beta | \mathcal{H}_\alpha \psi_\alpha - \sum_\beta \psi_\beta \lambda_{\beta\alpha} \rangle = \lambda_{\beta\alpha}. \]

The matrix elements of \( \lambda^- \) can be combined into one convergence measure:

\[
\Delta S \equiv \frac{1}{2} \sum_{q \in \{n,p\}} \left[ \frac{1}{\Omega_q^2} \sum_{\alpha,\beta \in q} \left| \lambda_{\beta\alpha}^- \right|^2 \right].
\]

(6.15)

The iteration stops once \( \Delta S \) is smaller than a specific threshold.
This convergence measure, however, does not always work for the pure HF case with no pairing present. Starting from the Hermiticity relation $\langle \psi_\alpha | \hat{h} | \psi_\beta \rangle = \langle \psi_\beta | \hat{h} | \psi_\alpha \rangle^*$, one can show that $\lambda_{\alpha\beta}^{-}$ vanishes when both $\psi_\alpha$ and $\psi_\beta$ are occupied ($v_\alpha = v_\beta = 1$) or unoccupied ($v_\alpha = v_\beta = 0$). Hence, in the pure HF calculation the quantity $\Delta S$ measures overlaps of occupied and unoccupied orbits, which reaches zero at the HF solution. Without pairing included, it is allowed that the size of the active s.p. space equals the number of particles ($\Omega_n = N$, $\Omega_p = Z$) and that all the s.p. levels are occupied. Under this circumstance, $\Delta S$ always stays zero and is thus not an appropriate measure of convergence; it instead becomes a measure of Hermiticity violation in our numerical implementations (see Secs. 6.1.1 and 6.2.1). Therefore, we should adopt the convergence measure utilized in Sky3D for the pure HF calculation:

$$\sqrt{\sum_\alpha \left( \langle \psi_\alpha | \hat{h}^2 | \psi_\alpha \rangle - \langle \psi_\alpha | \hat{h} | \psi_\alpha \rangle^2 \right)}.$$ (6.16)

### 6.1.5 Strategies for pairing

Since the HF solution with no pairing is a valid but probably unstable HFB solution, the iteration can be easily locked in a no-pairing solution for a long time; this phenomenon is called pairing breakdown. There exist many strategies to avoid this problem, and here we use a method similar to simulated annealing [230]. The iteration begins with an enhanced pairing strength which is gradually reduced to the physical value as the iteration goes on:

$$V_{\text{pair}}^{(\text{effective})} = V_{\text{pair}}^{(\text{physical})} \left[ \eta_{\text{enh}} \max(N_{\text{enh}} - n, 0) \right. \left. N_{\text{enh}} \right] + 1,$$ (6.17)

where $n$ is the iteration number. Default values of $\eta_{\text{enh}}$ and $N_{\text{enh}}$ are 2 and 400, respectively.

It is worth mentioning that the pairing annealing strategy also mitigates the point collapse
issue [77]. Here we only discuss the origin of the point collapse in a qualitative way. Assume that there exists a s.p. state $\psi_\alpha$ with $h_{\alpha\alpha} \gg \epsilon_F$ and thus $v_\alpha \approx 0$, $u_\alpha \approx 1$. Then we have $u_\alpha v_\alpha \gg v_\alpha^2$ and $\hat{H}_{\alpha} \approx u_\alpha v_\alpha \hat{h}$, which means that the generalized Hamiltonian becomes a local operator with no derivative when $\hat{h}$ is derived from a density-dependent $\delta$ interaction.

The corresponding solution is $\psi_\alpha(r) \propto \delta(r - r_{\text{min}})$, where $r_{\text{min}}$ is the minimum point of the pairing potential. Then the s.p. state $\alpha$ acquires a huge kinetic energy and thus a huge canonical s.p. energy $h_{\alpha\alpha}$, which drives the occupation fraction even closer to zero ($v_\alpha \to 0$) and forms a positive feedback. This phenomenon is called the point collapse since the corresponding wave function collapses into a spatial point around $r_{\text{min}}$. The enhanced pairing strength at the early stage of iterations ensures that the occupation amplitude $v_\alpha$ cannot be easily deadlocked at a small value and thus mitigates the point collapse. Otherwise, it will be difficult to recover correct pairing if some s.p. states are kicked out of the pairing window at an early stage prior to the convergence.

As for the pairing cutoff, the soft cutoff scheme discussed in Sec. 2.2.1 is adopted. In HFBFFT the cutoff weight $w$ given by Eq. (2.30) is a function of $e = h_{\alpha\alpha} - \epsilon_F$ and we choose $\Delta E_{\text{cut}} = 0.1 E_{\text{cut}}$ by default.

### 6.2 Benchmarks

Following the procedure presented in Ref. [231], we have benchmarked our new solver HFBFFT against several well established solvers, including HFBTHO, Sky1D (spherical systems only) and Sky2D. The Skyrme parameterization SLy4 [98] in the particle-hole channel and the mixed density-dependent $\delta$ interaction ($\rho_{\text{ref}} = 0.32$ fm$^{-3}$ in Eq. (2.21)) in the particle-particle channel are employed for benchmarks. The nuclei we use for benchmarks are: (i)
Table 6.1: Parameters adopted in different solvers for benchmark calculations: HFBTHO – the number of HO shells $N_{HO}$ and the axial deformation of the HO basis $\beta_2$; Sky1D, Sky2D and HFBFFT – the number of points $N_i$ and grid spacing $\delta i$ in the direction of $i$; Common parameter – the pairing cutoff energy $E_{cut}$.

<table>
<thead>
<tr>
<th>Solver</th>
<th>Parameters</th>
</tr>
</thead>
<tbody>
<tr>
<td>HFBTHO</td>
<td>$N_{HO} = 25, \beta_2 = 0$ (spherical systems) or 0.2, $E_{cut} = 60$ MeV</td>
</tr>
<tr>
<td>Sky1D</td>
<td>$N_r = 141, \delta r = 0.15$ fm, $E_{cut} = 15$ MeV</td>
</tr>
<tr>
<td>Sky2D</td>
<td>$N_r = N_z = 31, \delta r = \delta z = 0.7$ fm, $E_{cut} = 15$ MeV</td>
</tr>
<tr>
<td>HFBFFT</td>
<td>$N_x = N_y = N_z = 48, \delta x = \delta y = \delta z = 0.8$ fm, $E_{cut} = 15$ MeV</td>
</tr>
</tbody>
</table>

spherical closed-shell nuclei – $^{208}$Pb, $^{132}$Sn, (ii) spherical superfluid nucleus – $^{120}$Sn, (iii) axially deformed superfluid nuclei – $^{102}$Zr, $^{110}$Zr, and (iv) superfluid nucleus with a superdeformed fission isomer – $^{240}$Pu. All the nuclear systems used for benchmarks have axial and reflection symmetries, but they are rather different: $^{110}$Zr is a weakly bound system while other nuclei in (i), (ii) and (iii) are well bound; $^{120}$Sn and $^{102}$Zr have no proton pairing while static pairing correlations exist for both protons and neutrons in $^{110}$Zr and $^{240}$Pu.

Table 6.1 summarizes some parameters utilized in HFBTHO, Sky1D, Sky2D and HFBFFT. The reflection symmetry is imposed in both HFBTHO and Sky2D. The numbers of active neutron and proton s.p. states in three coordinate-space solvers (Sky1D, Sky2D and HFBFFT) for nuclei in items (i), (ii) and (iii) are $\Omega_n = 176$ and $\Omega_p = 126$, respectively; they are large enough as the total energy does not vary significantly ($< 10$ keV) when we increase $(\Omega_n, \Omega_p)$ to (200, 150). For $^{240}$Pu, however, the active s.p. space should be expanded to accommodate its large number of nucleons: We choose $(\Omega_n, \Omega_p) = (300, 200)$ for the ground state and $(\Omega_n, \Omega_p) = (400, 300)$ for the fission isomer. The box size adopted in HFBFFT is also large enough so that proton and neutron densities are small enough ($< 10^{-7}$ fm$^{-3}$) at the boundary.
Table 6.2: Total energies $E_{\text{tot}}$ (in MeV) and $\Delta S$ (in MeV) obtained from HFBFFT without and with Hermiticity restoration. Digits that do not coincide are marked in bold.

<table>
<thead>
<tr>
<th></th>
<th>Hermiticity broken</th>
<th>Hermiticity restored</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$E_{\text{tot}}$</td>
<td>$\Delta S$</td>
</tr>
<tr>
<td>$^{132}\text{Sn}$</td>
<td>$-1103.5429$</td>
<td>$3.44 \times 10^{-7}$</td>
</tr>
<tr>
<td>$^{208}\text{Pb}$</td>
<td>$-1635.6817$</td>
<td>$3.16 \times 10^{-7}$</td>
</tr>
<tr>
<td>$^{120}\text{Sn}$</td>
<td>$-1018.3310$</td>
<td>$3.44 \times 10^{-7}$</td>
</tr>
<tr>
<td>$^{110}\text{Zr}$</td>
<td>$-893.8578$</td>
<td>$4.59 \times 10^{-7}$</td>
</tr>
<tr>
<td>$^{102}\text{Zr}$</td>
<td>$-859.4696$</td>
<td>$4.94 \times 10^{-7}$</td>
</tr>
</tbody>
</table>

6.2.1 Effect of Hermiticity restoration

As mentioned in Sec. 6.1, Algorithm 2 for the position-varying derivative avoids the Hermiticity breaking caused by the incompatibility between the product rule and FFT-based differentiation. The effect of this Hermiticity restoration is shown in Table 6.2, where one can compare the results of some closed- and open-shell nuclei before and after the restoration.

For closed-shell $^{132}\text{Sn}$ and $^{208}\text{Pb}$ the HFB problem is reduced to the pure HF case, and thus we choose $(\Omega_n, \Omega_p) = (50, 82)$ for $^{132}\text{Sn}$ and $(\Omega_n, \Omega_p) = (82, 126)$ for $^{208}\text{Pb}$. The Hermiticity violation is then demonstrated by their non-vanishing $\Delta S$ in the third column of Table 6.2, and the magnitudes of their $\Delta S$ indicate the order of errors brought by the Hermiticity breaking. After the restoration, the values of $\Delta S$ obtained from HF calculations reach almost zero as Hermiticity is well preserved in the new implementation. On the other hand, the impact of the Hermiticity breaking on the total energy is usually a few keV and thus insignificant. Moreover, the convergence measure $\Delta S$ of a superfluid system does not vary significantly before and after the restoration, and is also larger than the error resulting from the Hermiticity violation. Hence, the Hermiticity breaking is not a critical issue for static HFB calculations. But it should not be neglected in a time-dependent framework where such errors can accumulate through time steps and become appreciable.
Table 6.3: Total energies $E_{\text{tot}}$ of $^{132}\text{Sn}$ and $^{208}\text{Pb}$ obtained from HFBTHO, HFBFFT, Sky1D and Sky2D. Various energy components are also listed: Subscript of densities denotes contributions from corresponding ED terms in Eq. (2.17), “Coul” the Coulomb energy, and “kin” the kinetic energy. Digits that do not coincide with HFBFFT are marked in bold.

<table>
<thead>
<tr>
<th></th>
<th>HFBTHO</th>
<th>HFBFFT</th>
<th>Sky1D</th>
<th>Sky2D</th>
</tr>
</thead>
<tbody>
<tr>
<td>$^{132}\text{Sn}$</td>
<td>$E_{\text{tot}}$</td>
<td>$-1103.49$</td>
<td>$-1103.54$</td>
<td>$-1103.57$</td>
</tr>
<tr>
<td></td>
<td>$E_{\text{kin,n}}$</td>
<td>$1637.71$</td>
<td>$1637.97$</td>
<td>$1638.01$</td>
</tr>
<tr>
<td></td>
<td>$E_{\text{kin,p}}$</td>
<td>$808.44$</td>
<td>$808.57$</td>
<td>$808.59$</td>
</tr>
<tr>
<td></td>
<td>$E_{\rho\rho}$</td>
<td>$-4876.26$</td>
<td>$-4877.02$</td>
<td>$-4877.04$</td>
</tr>
<tr>
<td></td>
<td>$E_{\rho\tau}$</td>
<td>$821.49$</td>
<td>$821.70$</td>
<td>$821.73$</td>
</tr>
<tr>
<td></td>
<td>$E_{\rho\nabla^2\rho}$</td>
<td>$248.11$</td>
<td>$248.23$</td>
<td>$248.25$</td>
</tr>
<tr>
<td></td>
<td>$E_{\rho\nabla\cdot\rho}$</td>
<td>$-84.40$</td>
<td>$-84.43$</td>
<td>$-84.44$</td>
</tr>
<tr>
<td></td>
<td>$E_{\text{Coul}}$</td>
<td>$341.42$</td>
<td>$341.44$</td>
<td>$341.44$</td>
</tr>
</tbody>
</table>

| $^{208}\text{Pb}$ | $E_{\text{tot}}$   | $-1635.46$ | $-1635.68$ | $-1635.70$ | $-1635.70$ |
|       | $E_{\text{kin,n}}$ | $2528.42$ | $2529.13$ | $2529.16$ | $2529.12$ |
|       | $E_{\text{kin,p}}$ | $1336.71$ | $1337.06$ | $1337.07$ | $1337.08$ |
|       | $E_{\rho\rho}$     | $-7845.66$ | $-7847.54$ | $-7847.63$ | $-7847.57$ |
|       | $E_{\rho\tau}$     | $1329.79$ | $1330.20$ | $1330.22$ | $1330.20$ |
|       | $E_{\rho\nabla^2\rho}$ | $315.12$ | $315.29$ | $315.29$ | $315.30$ |
|       | $E_{\rho\nabla\cdot\rho}$ | $-96.42$ | $-96.45$ | $-96.45$ | $-96.45$ |
|       | $E_{\text{Coul}}$  | $796.56$ | $796.63$ | $796.63$ | $796.62$ |

6.2.2 Benchmarks without pairing

We start our benchmarks from systems without pairing correlations, i.e., closed-shell nuclei. Table 6.3 shows the energies of doubly magic $^{132}\text{Sn}$ and $^{208}\text{Pb}$ obtained from various solvers; contributions from different functional terms are also given for comparison. These two nuclei are spherical so we can use 1D, 2D and 3D solvers for them. One can see that the results of three coordinate-space solvers agree with each other quite well, and the small differences are mainly attributed to their different grid geometries. The agreement between the results of HFBFFT and HFBTHO is acceptable but not perfect; the main reason is the slow convergence of the total energy (especially the kinetic energy) with respect to the number of HO shells [232, 233]. This is clearly shown by the large discrepancies of $E_{\text{kin}}$ and
between HFBTHO and HFBFFT. According to Refs. [232, 234], the relation between the total energy \( E_{\text{tot}} \) obtained from a HO-basis-based solver and the number of HO shells \( N \) can be approximated by
\[
E_{\text{tot}}(L) = E_\infty + a_0 e^{-2k_\infty L},
\]
where \( L \equiv \sqrt{2(N + 3/2 + 2)}b \), \( b \) is the oscillator length of the HO basis, and \( a_0, k_\infty \) and \( E_\infty \) should be obtained by fitting. Then \( E_\infty \) is the energy corresponding to an infinitely large model space. The fit for \(^{208}\text{Pb}\) yields \( E_\infty = -1635.786 \) MeV, which is closer to the energy given by HFBFFT.

### 6.2.3 Benchmarks with pairing

Due to different numerical representations adopted in different solvers, their discretized quasiparticle continua are also different. In addition, inconsistent cutoff schemes are adopted in these solvers, as we prefer a low cutoff energy and a large smearing factor in grid-based solvers due to the dense continuum in the coordinate-space representation. Thus, pairing strengths \( V_q \) \((q \in \{p, n\})\) are not portable and must be renormalized for benchmarks. Our strategy is to adjust pairing strengths in grid-based solvers to match the spectral pairing gap \( \Delta_q \) [70, 72, 235] given by HFBTHO. The pairing gap is defined in the canonical basis as

\[
\Delta_q = \frac{\sum_{\alpha \in q} w_\alpha v_\alpha^2 |\tilde{h}_{\alpha\alpha}|}{\sum_{\alpha \in q} w_\alpha v_\alpha^2}, \tag{6.18}
\]

where \( w_\alpha \) is the pairing cutoff weight (2.30). In HFBTHO, the neutron pairing strength \( V_n \) is tuned to reproduce the average experimental neutron pairing gap of \(^{120}\text{Sn}\), which is \( \Delta_n = 1.25 \) MeV, while the proton pairing strength \( V_p \) takes the same value as \( V_n \). Another quantity based on which we can tune pairing strengths is the sum of kinetic and pairing energies, \( \tilde{E}_{\text{kin}} = E_{\text{kin}} + E_{\text{pair}} \), as it is less sensitive to the pairing cutoff energy than its components [101, 236]. But the pairing gap is preferred because it is closely connected with
experimental observables and usually yields better total-energy agreement among different solvers.

Tables 6.4, 6.5 and 6.6 show the results obtained with different solvers for spherical, deformed and superdeformed systems. As shown in these tables, the values of various observables (total energy, total root-mean-square radius and quadrupole moments) in different columns agree fairly well after the renormalization, although energy components differ significantly. It is worth noting that the calculation for a deformed nucleus should start from the wave functions of a HO (in grid-based solvers) or Woods-Saxon (in HFBTHO) potential with an appropriate deformation. Besides, the fission isomer of $^{240}$Pu in principle should be determined by locating the local minimum on the potential-energy curve obtained from quadrupole-moment constrained calculations (see Fig. 7 in Ref. [231] for an example). This can be easily carried out in HFBTHO and Sky2D; in HFBFFT, however, the constrained calculation has not been implemented yet and the fission isomer is found by initializing iterations with various HO deformations.

6.3 Summary

We have developed a new HFB solver HFBFFT in the 3D coordinate space, using the canonical-basis formalism and damped gradient method. The development of the new program is based on Sky3D, a well-optimized, highly-parallelized HF+BCS program; the parallelization framework of HFBFFT is similar to that of Sky3D. A number of implementations has been done to ensure correct results and quick convergence, including the sub-iteration method, soft pairing cutoff, pairing annealing, and the new algorithm to restore Hermiticity in FFT-based numerical derivatives. We analyze a variety of nuclei with different deforma-
Table 6.4: Results of spherical $^{120}$Sn obtained from HFBTHO, HFBFFT, Sky1D and Sky2D, including total energies $E_{\text{tot}}$, some energy components, average pairing gaps $\Delta$ and total root-mean-square (rms) radii $r_{\text{rms}}$ (in fm). All energies are in MeV. Pairing strengths $V_n$ and $V_p$ adopted in these calculations are equal and listed in the last row. Digits that do not coincide with HFBFFT are marked in bold and numbers used for the pairing renormalization are in italics.

<table>
<thead>
<tr>
<th></th>
<th>HFBTHO</th>
<th>HFBFFT</th>
<th>Sky1D</th>
<th>Sky2D</th>
</tr>
</thead>
<tbody>
<tr>
<td>$^{120}$Sn</td>
<td>$E_{\text{tot}}$</td>
<td>(-1018.77)</td>
<td>(-1018.78)</td>
<td>(-1018.92)</td>
</tr>
<tr>
<td></td>
<td>$E_{\text{kin},n}$</td>
<td>(1340.51)</td>
<td>(1339.17)</td>
<td>(1339.14)</td>
</tr>
<tr>
<td></td>
<td>$E_{\text{kin},p}$</td>
<td>(830.75)</td>
<td>(831.25)</td>
<td>(831.31)</td>
</tr>
<tr>
<td></td>
<td>$E_{\text{pair},n}$</td>
<td>(-12.48)</td>
<td>(-9.29)</td>
<td>(-9.14)</td>
</tr>
<tr>
<td></td>
<td>$E_{\text{pair},p}$</td>
<td>(0.00)</td>
<td>(0.00)</td>
<td>(0.00)</td>
</tr>
<tr>
<td></td>
<td>$\tilde{E}_{\text{kin},n}$</td>
<td>(1328.03)</td>
<td>(1329.88)</td>
<td>(1330.01)</td>
</tr>
<tr>
<td></td>
<td>$\Delta_n$</td>
<td>(1.25)</td>
<td>(1.25)</td>
<td>(1.25)</td>
</tr>
<tr>
<td></td>
<td>$\Delta_p$</td>
<td>(0.00)</td>
<td>(0.00)</td>
<td>(0.00)</td>
</tr>
<tr>
<td></td>
<td>$r_{\text{rms}}$</td>
<td>(4.67)</td>
<td>(4.67)</td>
<td>(4.67)</td>
</tr>
<tr>
<td></td>
<td>$V_n$, $V_p$</td>
<td>(-284.57)</td>
<td>(-361.80)</td>
<td>(-367.30)</td>
</tr>
</tbody>
</table>

Table 6.5: Results of axially deformed $^{102}$Zr and $^{110}$Zr obtained from HFBTHO, HFBFFT and Sky2D. Besides quantities shown in Table 6.4, quadrupole moments $Q_{20}$ (in fm$^2$) are also included here. Pairing strengths $V_n$ and $V_p$ adopted in these calculations are listed in the last two rows. Digits that do not coincide with HFBFFT are marked in bold and numbers used for the pairing renormalization are in italics.

<table>
<thead>
<tr>
<th></th>
<th>$^{102}$Zr</th>
<th>$^{110}$Zr</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>HFBTHO</td>
<td>HFBFFT</td>
</tr>
<tr>
<td>$E_{\text{tot}}$</td>
<td>(-859.65)</td>
<td>(-859.69)</td>
</tr>
<tr>
<td>$E_{\text{kin},n}$</td>
<td>(1202.02)</td>
<td>(1200.96)</td>
</tr>
<tr>
<td>$E_{\text{kin},p}$</td>
<td>(651.25)</td>
<td>(651.22)</td>
</tr>
<tr>
<td>$E_{\text{pair},n}$</td>
<td>(-3.39)</td>
<td>(-2.50)</td>
</tr>
<tr>
<td>$E_{\text{pair},p}$</td>
<td>(-1.97)</td>
<td>(-1.42)</td>
</tr>
<tr>
<td>$\tilde{E}_{\text{kin},n}$</td>
<td>(1198.63)</td>
<td>(1199.53)</td>
</tr>
<tr>
<td>$E_{\text{kin},p}$</td>
<td>(649.28)</td>
<td>(649.79)</td>
</tr>
<tr>
<td>$\Delta_n$</td>
<td>(0.69)</td>
<td>(0.69)</td>
</tr>
<tr>
<td>$\Delta_p$</td>
<td>(0.56)</td>
<td>(0.56)</td>
</tr>
<tr>
<td>$r_{\text{rms}}$</td>
<td>(4.58)</td>
<td>(4.58)</td>
</tr>
<tr>
<td>$Q_{20,n}$</td>
<td>(639)</td>
<td>(639)</td>
</tr>
<tr>
<td>$Q_{20,p}$</td>
<td>(411)</td>
<td>(411)</td>
</tr>
<tr>
<td>$V_n$</td>
<td>(-284.57)</td>
<td>(-367.00)</td>
</tr>
<tr>
<td>$V_p$</td>
<td>(-284.57)</td>
<td>(-372.00)</td>
</tr>
</tbody>
</table>
Table 6.6: Similar to Table 6.5, but for the ground state and fission isomer of $^{240}$Pu.

<table>
<thead>
<tr>
<th>$^{240}$Pu</th>
<th>ground state</th>
<th>fission isomer</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>HFBTHO</td>
<td>HFBFFT</td>
</tr>
<tr>
<td>$E_{\text{tot}}$</td>
<td>$-1802.11$</td>
<td>$-1802.43$</td>
</tr>
<tr>
<td>$E_{\text{kin,n}}$</td>
<td>$2938.92$</td>
<td>$2939.94$</td>
</tr>
<tr>
<td>$E_{\text{kin,p}}$</td>
<td>$1520.95$</td>
<td>$1521.46$</td>
</tr>
<tr>
<td>$E_{\text{pair,n}}$</td>
<td>$-3.11$</td>
<td>$-2.30$</td>
</tr>
<tr>
<td>$E_{\text{pair,p}}$</td>
<td>$-1.54$</td>
<td>$-1.22$</td>
</tr>
<tr>
<td>$\tilde{E}_{\text{kin,n}}$</td>
<td>$2935.81$</td>
<td>$2937.64$</td>
</tr>
<tr>
<td>$\tilde{E}_{\text{kin,p}}$</td>
<td>$1519.40$</td>
<td>$1520.25$</td>
</tr>
<tr>
<td>$\Delta_n$</td>
<td>$0.44$</td>
<td>$0.44$</td>
</tr>
<tr>
<td>$\Delta_p$</td>
<td>$0.33$</td>
<td>$0.33$</td>
</tr>
<tr>
<td>$r_{\text{rms}}$</td>
<td>$5.93$</td>
<td>$5.93$</td>
</tr>
<tr>
<td>$Q_{20,n}$</td>
<td>$1784$</td>
<td>$1782$</td>
</tr>
<tr>
<td>$Q_{20,p}$</td>
<td>$1166$</td>
<td>$1165$</td>
</tr>
<tr>
<td>$V_n$</td>
<td>$-284.57$</td>
<td>$-360.00$</td>
</tr>
<tr>
<td>$V_p$</td>
<td>$-284.57$</td>
<td>$-355.00$</td>
</tr>
</tbody>
</table>

As a 3D coordinate-space solver, HFBFFT performs better than HO-basis-based solvers for the study of deformed and weakly bound systems. There are also many features we plan to add to HFBFFT to make it more versatile, such as the deformation constraint, the blocking procedure for odd-$A$ and odd-odd systems, and the pairing regularization that removes the dependence of pairing strengths on the cutoff. We are also going to further optimize the performance of HFBFFT for modern supercomputers using GPU architectures.
Chapter 7

Conclusions

As shown in this dissertation, the nuclear-DFT framework has broad applications to various nuclear physics problems. Research works discussed in this dissertation develop the framework, and employ it to study nuclear ground-state properties, collective rotation and beta decays. We also demonstrate that the combination of the nuclear DFT and tools developed in other fields (e.g., the localization function developed in the electronic DFT, statistics, and high-performance computing) can be helpful.

Chapter 3 shows the NLF patterns in two rotating systems, the CHO model and SD\textsuperscript{152}Dy. These two examples demonstrate the “constructive-interference” mechanism that explains the usefulness of the NLF in the visualization of the internal structure; they also reveal the close connection between NLF patterns and high-lying occupied s.p. orbits. Chapter 4 discusses the origin of reflection-asymmetric ground-state shapes. The roles that different multipole components of the energy play in the onset of pear-like deformations are examined for Ra and Yb isotopic chains, and high-order components are small but important due to the strong cancellation between monopole and octupole parts. Besides, the coupling between close-lying orbits with $\Delta \ell = \Delta j = 3$ across the Fermi energy is shown to be responsible for the reflection asymmetry. Chapter 5 presents the procedure and results of the model calibration for beta-decay calculations. The time-odd Skyrme couplings, isoscalar pairing strength, and effective axial-vector coupling are fitted within the $\chi^2$-optimization.
framework; their uncertainties and correlations are also obtained in the fit, with the number of effective parameters determined by the PCA. The $\chi^2$ optimization paves the way for the Bayesian model calibration, and can be utilized to provide high-quality beta-decay inputs with quantified uncertainties for $r$-process simulations. Finally, chapter 6 finally discusses the numerical implementation and benchmark of the new HFB solver HFBFFT. This new solver uses the canonical-HFB formalism in the 3D coordinate-space representation, so it is an efficient and reliable tool for the studies of weakly bound and large-deformed nuclei. HFBFFT is highly performant and well parallelized, and its correctness is ensured by careful benchmarks against other HFB solvers. We expect to add more features to HFBFFT and further optimize it for future applications.

Generally speaking, there are three main ingredients necessary in the nuclear-DFT research, namely, (i) universal and well calibrated EDF parametrizations, (ii) versatile and highly performant solvers, and (iii) useful approaches to extract physics information. Works presented in this dissertation has significantly contributed to all these aspects.
APPENDICES
Appendix A

List of my contributions

   - Performed cranked HF calculations for rotating $^{152}$Dy.
   - Carried out the analysis and prepared figures regarding the results of rotating $^{152}$Dy and the cranked harmonic-oscillator model.
   - Wrote the first draft for the paper.

   - Prepared Figs. 11 and 12 to facilitate discussions on s.p. levels.
   - Checked equations in Sec. III and proofread the draft.

   - Performed a large portion of work regarding the code development.
   - Wrote the last two paragraphs in Sec. 2.2 and the whole Sec. 3.6.
   - Checked all the equations and proofread the draft.

- Improved the performance of PNFAM by replacing loops with BLAS routines.


- Added new codes to PyNFAM to connect the physics model with the $\chi^2$-optimization routine.
- Produced physics-model outputs for the GP emulator building.

6. Presentations

- “Model calibration for beta-decay calculations”, TRIUMF theory seminar, online seminar hosted by TRIUMF, Dec 2021.
- “Skyrme EDF parameter calibration for beta-decay calculations”, 2021 NUCLEI collaboration meeting, online, Jun 2021.
- “Nucleon localization function in rotating nuclei”, 2021 APS April meeting, online, Apr 2021.
- “Nucleon localization function in rotating systems”, 2020 NUCLEI collaboration meeting, online, Jun 2020.


T. Shafer, J. Engel, C. Fröhlich, G. C. McLaughlin, M. Mumpower, and R. Surman. \(\beta\) decay of deformed \(r\)-process nuclei near \(A = 80\) and \(A = 160\), including odd-\(A\) and odd-odd nuclei, with the skyrme finite-amplitude method. \emph{Phys. Rev. C}, 94(5):055802, November 2016.


[201] Evaluated nuclear structure data file (ENSDF).


