Theoretical study of triaxial shapes of neutron-rich Mo and Ru nuclei

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Background: Whether atomic nuclei can possess triaxial shapes at their ground states is still a subject of ongoing debate. According to theory, good prospects for low-spin triaxiality are in the neutron-rich Mo-Ru region. Recently, transition quadrupole moments in rotational bands of even-mass neutron-rich isotopes of molybdenum and ruthenium nuclei have been measured. The new data have provided a challenge for theoretical descriptions invoking stable triaxial deformations.

Purpose: To understand experimental data on rotational bands in the neutron-rich Mo-Ru region, we carried out theoretical analysis of moments of inertia, shapes, and transition quadrupole moments of neutron-rich even-even nuclei around ¹¹⁰Ru using self-consistent mean-field and shell model techniques.

Methods: To describe yrast structures in Mo and Ru isotopes, we use nuclear density functional theory (DFT) with the optimized energy density functional UNEDF0. We also apply triaxial projected shell model (TPSM) to describe yrast and positive-parity, near-yrast band structures.

Results: Our self-consistent DFT calculations predict triaxial ground-state deformations in ^{106,108}Mo and ^{108,110,112}Ru and reproduce the observed low-frequency behavior of moments of inertia. As the rotational frequency increases, a negative- γ structure, associated with the aligned $\nu(h_{11/2})^2$ pair, becomes energetically favored. The computed transition quadrupole moments vary with angular momentum, which reflects deformation changes with rotation; those variations are consistent with experiment. The TPSM calculations explain the observed band structures assuming stable triaxial shapes.

Conclusions: The structure of neutron-rich even-even nuclei around ¹¹⁰Ru is consistent with triaxial shape deformations. Our DFT and TPSM frameworks provide a consistent and complementary description of experimental data.

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I. INTRODUCTION

Neutron-rich molybdenum and ruthenium isotopes are known to exhibit shape changes and shape-coexistence phenomena [1–6]. With increasing neutron number, triaxial deformation is expected to appear in their ground states due to the occupation of $1\nu h_{11/2}$ and $1\pi g_{9/2}$ intruder orbitals [7].

Experimentally, the clearest signature of triaxial shapes comes from the γ -ray spectroscopy of rotating nuclei [8,9]. Following Ref. [10], which reported evidence for rotationallike behavior in the very neutron-rich even-even Zr-Pd region, numerous experiments were devoted to investigations of shape transitions and rotational properties in this region. The first systematic high-spin study of collective band structures was undertaken in Ref. [11], which reported deformed configurations in ^{103,104,107}Zr and ^{107,108}Mo and ¹⁰⁸Mo.

The early work [2] on ¹⁰⁴Ru was indicative of a transition from spherical to triaxial shapes. The collective nature of neutron-rich nuclei and triaxiality in ¹¹⁰Ru and ¹¹²Ru was confirmed in Ref. [12] by reporting a steady decrease of the γ -band bandhead energy. More evidence of collective triaxial behavior of Ru isotopes came from the spectroscopy of fission fragments [13–18]. The decrease of transition quadrupole moments at high spin showed that the triaxial deformation in neutron-rich Mo isotopes could be spindependent [19]. Another piece of experimental information on triaxial deformations came from the measurement of the quasi- γ band in ¹¹⁰Mo [20,21]. A more detailed information on quadrupole collectivity was obtained by Coulomb excitation studies, which succeeded in determining unique sets of *E*2 and *M*1 matrix elements in ¹⁰⁴Ru [3] and ¹¹⁰Mo [3] and extracting triaxial deformations using the collective quadrupole invariant approach. Rotational bands in ¹⁰⁶Mo and ^{108,110,112}Ru were investigated in Refs. [4,22], which reported chiral doublets associated with triaxial nuclear rotation. In recent papers [6,23], transition quadrupole moments of rotational bands in neutron-rich, even mass ^{102–108}Mo and ^{108–112}Ru nuclei were measured for the spin range of 8–16 \hbar , suggesting γ -softness effects or even triaxiality in these nuclei.

Theoretically, triaxial ground states in this region have been investigated with different models. In Ref. [7], based on macroscopic-microscopic approach, triaxial ground-state (g.s.) minima were found in the neutron-rich Mo isotopes with N = 62-66 and also in Ru isotopes. In a systematic survey of Refs. [24,25], the largest shell effects due to triaxial deformations were found around ¹⁰⁸Ru. The interacting boson model analysis of Ref. [26] did not find any candidates for stable triaxiality in this region. In Ref. [27], potential energy surface (PES) calculations for Ru isotopes were carried out with Hartree-Fock (HF) and interacting boson models, and shallow triaxial minima were found for N = 64-70 (see also Ref. [28]). In the self-consistent Hartree-Fock Bogoliubov study of Ref. [29] with Gogny D1S interaction, triaxial deformations were predicted for even-even isotopes $^{104-110}$ Mo and 104,106 Ru.

Collective rotation has been shown to enhance triaxial minima in even-even Mo and Ru isotopes [7,30,31]. For that reason, those nuclei are candidates for the presence of novel collective modes associated with triaxial rotation, such as wobbling motion and chiral bands [8,9]. The angular momentum alignment pattern in the lowest bands of Mo and Ru isotopes was also explored within the projected shell model approach [32,33]. The axial study [32] provided a reasonable description of yrast spectra and electromagnetic properties of ^{100–112}Ru. This work was extended in Ref. [33], which also contains TPSM analysis of ¹¹⁰Mo and ¹¹⁴Ru. In the case of ¹¹⁴Ru, stable γ deformation turned out to be crucial for reproducing the data.

In an attempt to explain the recent data on transition quadrupole moments in Mo and Ru nuclei, cranked relativistic mean-field calculations [23] predicted axial prolate and oblate ground states in those nuclei. However, the angular momentum dependence of resulting transition quadrupole moments was not consistent with observations. As concluded in Ref. [23]: "Attempts to describe the observations in mean-field based models, specifically cranked relativistic Hartree-Bogoliubov theory, illustrate the challenge theory faces and the difficulty to infer information on γ -softness and triaxiality from the data." To shed some light on this puzzle, and to further explore the importance of triaxial deformation in this mass region, we apply the cranked self-consistent Hartree-Fock-Bogoliubov (CHFB) method and TPSM to the rotational properties of neutron-rich, even-even Mo and Ru isotopes.

This article is organized as follows. Section II gives a brief introduction to CHFB and TPSM models used in this work. In Sec. III, the results of calculations for Mo and Ru isotopes are presented and compared with experiment. Therein, we discuss potential energy and Routhian surfaces, quasiparticle Routhian spectra, and equilibrium deformations. To test the stability of CHFB minima with respect to angular momentum orientation, we carry our tilted-axis cranking calculations employing the Kerman-Onishi conditions. Finally, the conclusions of this work are given in Sec. IV.

II. THE MODEL

A. Cranked Skyrme-Hartree-Fock-Bogoliubov model

Our CHFB calculations were performed with the DFT solver HFODD (version 2.49t) [34]. Parity and *y*-signature $[\hat{R}_y = \exp(-i\pi \hat{J}_y)]$ symmetries are conserved; the corresponding quantum numbers are denoted as π and *r*. The quasiparticle HFB wave functions were expanded in 800 spherical harmonic oscillator basis states with the oscillator frequency of $\hbar\omega = 49.2 \text{ MeV}/A^{1/3}$. We have tested that such a basis provides a very reasonable precision for the observables studied.

In the particle-hole channel, we employ the global Skyrme energy density functional UNEDF0 optimized in Ref. [35]. In the pairing channel, we take the zero-range density-dependent pairing force [36] with the Lipkin-Nogami correction for particle-number fluctuations. The original pairing strengths are taken as $(V_0^{\nu}, V_0^{\pi}) = (-170.374, -199.202)$ MeV fm³, with a cutoff energy in the quasiparticle spectrum of $E_{\text{cut}} = 60$ MeV. In the present calculation, the strengths of the pairing force for neutrons and protons have been increased by 5% to reproduce the kinematic moment of inertia of the g.s. band (g.b.) of ¹⁰⁶Mo. As discussed below, the calculated potential energy surfaces are not sensitive to such a small variation of pairing strengths.

In the multidimensional potential energy surface calculations, the constraints are imposed on expectation values of multipole moments. We use the augmented Lagrangian method [37] to perform the constrained iterations. The total Routhian calculations were computed within the principal-axis cranking approach [9]. However, to study the stability of the resulting triaxial minima with respect to the orientation of the angular momentum vector, we applied the Kerman-Onishi conditions implemented as in Refs. [38,39]. Since the Lipkin-Nogami method is not strictly variational, the g.b. minimum at nonzero angular momentum was obtained by minimizing the constrained total Routhian surface. This increases precision of calculations, especially when the minima are soft [40].

B. Triaxial projected shell model

Recently, multi-quasiparticle TPSM approach has been developed and it has been shown to provide a consistent description of yrast, γ (K = 2) and $\gamma\gamma$ (K = 4) bands in transitional nuclei [41,42]. In this method, the three-dimensional projection technique is employed to project out the good angular-momentum states from product states built upon quasiparticle (q.p.) configurations of triaxially deformed Nilsson+BCS model. The shell model Hamiltonian is then diagonalized in this angular-momentum projected basis. The TPSM space includes multi-quasiparticle states; hence, is capable of describing near-yrast band structures at high-spins [33,43–46].

The TPSM basis employed in this study consists of 0-q.p. vacuum, two-proton, two-neutron, and the four-q.p. configurations [45]. The q.p. basis chosen is adequate to describe high-spin states up to angular momentum $I \sim 20$. In the present analysis we shall, therefore, restrict our discussion to this spin regime. As in the earlier TPSM calculations, we use the pairing plus quadrupole-quadrupole Hamiltonian [33,47]:

$$\hat{H} = \hat{H}_0 - \frac{1}{2}\chi \sum_{\mu} \hat{Q}^{\dagger}_{\mu} \hat{Q}_{\mu} - G_M \hat{P}^{\dagger} \hat{P} - G_Q \sum_{\mu} \hat{P}^{\dagger}_{\mu} \hat{P}_{\mu}, \quad (1)$$

where \hat{H}_0 is the single-particle spherical Nilsson Hamiltonian, χ is the strength of the quadrupole-quadrupole force related in a self-consistent way to deformation of the q.p. basis, and G_M and G_Q are the strengths of the monopole and quadrupole pairing terms, respectively. The configuration space employed corresponds to three principal oscillator shells \mathcal{N}_{osc} : $\nu[3,4,5]$ and $\pi[2,3,4]$. The pairing strengths have been parametrized as in Refs. [32,48] in terms of two constants G_1 and G_2 . In this work, we choose $G_1 = 16.22$ MeV and $G_2 = 22.68$ MeV; with these pairing strengths we approximately reproduce the experimental odd-even mass differences in this region. The



FIG. 1. (Color online) PES in (Q_{20}, Q_{22}) plane in CHFB+UNEDF0 for ¹⁰⁶Mo and ¹⁰⁸Mo. Left: standard pairing strengths. Right: pairing strengths increased by 5%, see text. The difference between contour lines is 0.5 MeV.

quadrupole pairing strength G_Q is assumed to be proportional to G_M , and the proportionality constant was set to 0.18 [32,48]. The single-particle basis is that of the deformed Nilsson Hamiltonian parametrized in terms of axial (ε) and triaxial (ε') deformations related to the standard Bohr triaxiality parameter γ by $\gamma = \arctan(\varepsilon'/\varepsilon)$.

III. RESULTS AND DISCUSSIONS

A. CHFB results

1. Ground-state potential energy surfaces

The g.s. UNEDF0 PESs for ^{106,108}Mo and ^{108,110,112}Ru are shown in Figs. 1 and 2, respectively. All these nuclei are predicted to be triaxial in their ground states. It is seen that the PESs are practically not affected by a 5% change in the pairing strengths. In particular, the triaxial minima appearing at $(Q_{20}, Q_{22}) \approx (8.0-9.5, 2.0-3.0)$ b are not affected by pairing. The corresponding g.s. quadrupole deformations (β_2, γ) are displayed in Table I. For ^{106,108}Mo, we predict the triaxial g.s. minima at $(\beta_2, \gamma) \approx (0.19, 17^\circ)$. Similar results were obtained in the macroscopic-microscopic calculations of Refs. [7,25,49] and HFB+D1S calculations [29]. For ^{108,110,112}Ru, we also predict triaxial g.s. minima; this is consistent with Refs. [7,31] and HF+SIII calculations of Ref. [12]. Triaxial g.s. shapes for

TABLE I. Bohr quadrupole deformation parameters β and γ calculated in CHFB+UNEDF0 for ^{106,108}Mo and ^{108,110,112}Ru.

| | ¹⁰⁶ Mo | ¹⁰⁸ Mo | ¹⁰⁸ Ru | ¹¹⁰ Ru | ¹¹² Ru |
|-----------|-------------------|-------------------|-------------------|-------------------|-------------------|
| β_2 | 0.19 | 0.18 | 0.16 | 0.16 | 0.15 |
| γ | 16° | 18° | 24° | 25° | 24° |



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FIG. 2. (Color online) Similar to Fig. 1, but for ¹⁰⁸Ru, ¹¹⁰Ru, and ¹¹²Ru.

^{108,110}Ru were also obtained in the survey [25] but ¹¹²Ru was calculated to be axial.

2. High spin behavior

The angular momentum alignment pattern of Mo and Ru nuclei is governed by the $vh_{11/2}$ and $\pi g_{9/2}$ high-j q.p. excitations, which give rise to strong shape polarization effects [7]. Figures 3 and 4 show self-consistent CHFB+UNEDF0 1-q.p. Routhian diagrams versus rotational frequency for ¹⁰⁶Mo and ¹¹²Ru, respectively. In both cases, the alignment of $\nu(h_{11/2})^2$ and $\pi(g_{9/2})^2$ pairs occurs at similar rotational frequencies of $\hbar\omega \approx 0.3$ MeV. At higher rotational frequencies, a transition is expected from the g.b. configuration to aligned $\nu(h_{11/2})^2$ and $\pi(g_{9/2})^2$ 2-q.p. bands, and then to a 4-q.p. $\nu(h_{11/2})^2 \pi(g_{9/2})^2$ band. These two consecutive crossings are difficult to follow in CHFB calculations, as this would require a diabatic-configuration extension [50-52] of the current framework. Such an extension is highly nontrivial in CHFB as the self-consistent mean fields associated with aligned configurations are expected to differ significantly from those of the g.b. [7]. Moreover, pairing correlations in the aligned bands are quenched and this causes numerical instabilities around the band crossing. Therefore, to provide interpretation of the transition quadrupole moments at higher angular momenta, we carry out cranked Skyrme-Hartree-Fock (CHF) calculations without pairing at $\hbar \omega > 0.3$ MeV. In this case, diabatic configurations can be defined by the



FIG. 3. (Color online) One-quasiparticle Routhian diagram for ¹⁰⁶Mo obtained with CHFB+UNEDF0. The parity π and signature r of individual levels are indicated in the following way: $\pi = +, r = +i$ – solid line; $\pi = +, r = -i$ – dotted line; $\pi = -, r = +i$ – dot-dashed line; $\pi = -, r = -i$ – dashed line. The thin line indicates the Fermi energy.

number of single-particle Routhians occupied in the four parity-signature blocks [53]. Specifically, each neutron and proton configurations is defined by four occupation numbers $[N_{++}, N_{+-}, N_{-+}, N_{--}]$ representing the number of particles



FIG. 4. (Color online) Similar to Fig. 3, but for ¹¹²Ru.



FIG. 5. (Color online) Angular momentum alignment for ¹⁰⁶Mo and ^{108,112}Ru. CHFB ($\hbar \omega < 0.3$ MeV) and CHF ($\hbar \omega > 0.3$ MeV) calculations are compared to experiment [14–16].

 $N_{\pi,r}$ occupying single-particle states of given π and r. The lowest total Routhian with $\pi = +$ and r = 1 is associated with the yrast configuration.

The angular momentum alignment (total angular momentum as a function of rotational frequency) is shown in Fig. 5 for ¹⁰⁶Mo, ¹⁰⁸Ru, and ¹¹²Ru. Below the predicted band crossing at $\hbar\omega \approx 0.3$ MeV, our calculations reproduce experiment well. (Note that our pairing strengths were adjusted to match the kinematic moment of inertia of ¹⁰⁶Mo.) The first band crossing, associated with the alignment of the $h_{11/2}$ neutron pair, is seen in ^{108,112}Ru data slightly below $\hbar\omega = 0.4$ MeV, and it is significantly delayed in ¹⁰⁶Mo. The predicted aligned configuration above the band crossing has a fairly different shape as compared to that of the g.b., and it is difficult to follow the g.b. at $\hbar\omega > 0.3$ MeV.

To investigate the evolution of nuclear shapes with rotation, we compute the equilibrium β_2 and γ deformations for low-lying $\pi = +, r = 1$ bands in ^{106,108}Mo (Fig. 6) and ^{108,110,112}Ru (Fig. 7). In all cases considered, the triaxial paired g.b. undergoes small centrifugal stretching in the direction of β_2 . For instance, in the case of ¹⁰⁸Ru, β_2 increases from the value of 0.15 at $\hbar\omega = 0$ to 0.17 at $\hbar\omega = 0.3$ MeV.

At higher spins $(10 \le I \le 36)$, when pairing is neglected in our calculations, it is useful to label many-body configurations $[N_{++}, N_{+-}, N_{-+}, N_{--}]$ by the number of occupied intruder levels, i.e., $\mathcal{N}_{osc} = 4$ protons (primarily $g_{9/2}$) and $\mathcal{N}_{osc} = 5$ neutrons (primarily $h_{11/2}$). For instance, the aligned configuration $\pi(9,9,12,12) \otimes \nu(17,17,15,15)$ in ¹⁰⁶Mo [shown by circles in Fig. 6(a)] can be denoted as $\pi 4^4 \nu 5^4$, and the same label applies to the $\pi(9,9,12,12) \otimes \nu(18,18,15,15)$ in ¹⁰⁸Mo [shown by up-triangles in Fig. 6(b)].

The quadrupole deformations β_2 of aligned bands are predicted to be in the range of $0.12 \leq \beta_2 \leq 0.16$, which represents a reduction as compared to the shapes of paired ground-state bands. The aligned bands remain triaxial with



FIG. 6. (Color online) Summary of equilibrium deformations of the lowest $\pi = +, r = 1$ bands in ^{106,108}Mo calculated with CHFB+UNEDF0 (ground band) and CHF+UNEDF0 (aligned bands). The rotational frequency is varied from zero to $\hbar\omega = 0.6$ MeV. The corresponding range of angular momentum is indicated. The aligned band are classified according to the number of occupied high- \mathcal{N} intruder levels ($\mathcal{N} = 5$ and 4 for neutrons and protons, respectively).

 γ values around -30° up to $\hbar\omega = 0.6$ MeV. This finding is consistent with the results of early Ref. [7] employing cranked macroscopic-microscopic approach. At the highest rotational frequencies considered, our calculations predict the appearance of aligned triaxial configurations with $\gamma > 0$, which eventually terminate at oblate shapes ($\gamma \approx 60^{\circ}$), see, e.g., Fig. 6(b).

To study the stability of different triaxial minima at high spins, we analyzed related diabatic total Routhians in the (Q_{20}, Q_{22}) plane. In Fig. 8 we show the total Routhian surfaces at $\hbar\omega = 0.5$ MeV for the selected low-lying aligned configurations in ¹⁰⁶Mo discussed in Fig. 6(a). For all those configurations, the collective triaxial minimum with γ between -30° and -15° appears lowest in energy. For the configuration $\pi 4^4 \nu 5^4$ shown in Fig. 8(a), we also predict a noncollective oblate state with I = 34 that represents a termination point of $\gamma > 0$ band.



FIG. 7. (Color online) Similar to Fig. 6, but for ^{108,110,112}Ru.



FIG. 8. (Color online) Diabatic total Routhian surfaces for ¹⁰⁶Mo at $\hbar \omega = 0.5$ MeV calculated in CHF+UNEDF0 for the configurations: (a) $\pi(9,9,12,12) \otimes \nu(17,17,15,15) [\pi 4^4 \nu 5^4$ in Fig. 6(a)]; (b) $\pi(10,10,11,11) \otimes \nu(17,17,15,15) (\pi 4^6 \nu 5^4)$; and (c) $\pi(10,10,11,11) \otimes \nu(16,16,16,16) (\pi 4^6 \nu 5^6)$.

To eliminate spurious minima that are unstable with respect to the angular momentum orientation, we also investigated the dependence of the Routhians on the angular momentum tilting angle θ with respect to the axis of rotation (y axis). To this end, we used the tilted-axis-cranking formalism of Refs. [38,39]. The calculations were performed for the aligned bands in ¹⁰⁶Mo At $\hbar \omega < 0.5$ MeV, the total Routhians of triaxial ($\gamma <$ 0) configurations $\pi 4^4 \nu 5^4$, $\pi 4^5 \nu 5^5$, and $\pi 4^6 \nu 5^4$ of Fig. 8(a) show a minimum at $\theta = 0^\circ$. At $\hbar \omega \approx 0.5$ MeV, the Routhians become very soft in θ , indicating a large-amplitude collective motion in this direction. This instability is not present for a $(\pi = -, r = 1)$ configuration $\pi(9, 9, 12, 12) \otimes \nu(18, 17, 15, 14)$ $(\pi 4^4 \nu 5^5)$, which shows a pronounced minimum at $\theta = 90^\circ$ associated with $\gamma > 0$. This result is consistent with the deformation-driving effect of aligned $h_{11/2}$ neutrons orbitals discussed in Ref. [7].

The transition quadrupole moments along the yrast band in ^{106,108}Mo and ^{108,110,112}Ru are shown in Fig. 9 as a function of rotational frequency. At low rotational frequencies $\hbar\omega < 0.3$ MeV, there is a gradual increase of Q_t with ω due to the centrifugal stretching effect seen in Figs. 6 and 7. As discussed earlier, at higher frequencies cranking calculations are performed without pairing. While this approximation seriously affects the predicted angular momentum alignment shown in Fig. 5, the equilibrium shapes obtained in the CHF



FIG. 9. (Color online) Transition quadrupole moments for ^{106,108}Mo and ^{108,110,112}Ru calculated in CHFB (open circles) and CHF (dots) compared to experiment. The Q_t value at I = 2 is taken from Ref. [54] and the high-spin values from Ref. [23].

are reasonable approximations to those obtained in the full CHFB framework [55,56], and reproduce experimental Q_t values for aligned configurations [57,58]. As seen in Fig. 9, the predicted transition quadrupole moments in aligned bands are slightly reduced with respect to the low-spin region due to the deformation reduction associated with the aligned $\nu h_{11/2}$ and $\pi g_{9/2}$ pairs. This reduction is generally consistent with experiment, except perhaps for ¹¹⁰Ru, where theory overestimates the measured Q_t values above $\hbar \omega = 0.3$ MeV.

B. TPSM results

The TPSM calculations proceed in several stages. In the first stage, the deformed basis is constructed based on the eigenstates of the triaxially deformed Nilsson potential. The β_2

TABLE II. Triaxial quadrupole deformation parameters γ employed in the TPSM calculation for ^{106,108}Mo and ^{108,110,112}Ru isotopes.

| Nucleus | ¹⁰⁶ Mo | ¹⁰⁸ Mo | ¹⁰⁸ Ru | ¹¹⁰ Ru | ¹¹² Ru |
|---------|-------------------|-------------------|-------------------|-------------------|-------------------|
| γ | 20° | 25° | 29° | 28° | 25° |



FIG. 10. (Color online) Comparison between experimental [59] and calculated band structure of 112 Ru.

deformation has been chosen such that the lowest quadrupole transition from $2^+ \rightarrow 0^+$ is reproduced. The non-axial deformation parameter γ is chosen from the minimum of the g.s. PES obtained in TPSM calculations. For ¹⁰⁸Mo and ^{108,110}Ru, where the PES minima are γ -soft, triaxial deformation was



FIG. 11. (Color online) Comparison of experimental and calculated band structures in 106 Mo at four values of triaxial deformation γ .

adjusted to the experimental bandhead energy of the γ band as it is known to be quite sensitive to the γ deformation. The adopted values of γ are listed in Table II. It is seen that strongly triaxial shapes are expected in all cases, and this confirms the CHFB+UNEDF0 results.

In the next step, the good angular-momentum basis is obtained from the triaxial Nilsson+BCS wave functions by using



FIG. 12. (Color online) Transition quadrupole moments for ^{106,108}Mo and ^{108,110,112}Ru calculated in TPSM (solid line) compared to experiment. The Q_t value at I = 2 is taken from Ref. [54] and the high-spin values from Ref. [23]. For ¹⁰⁶Mo, we also show TPSM results at $\gamma = 5^{\circ}, 15^{\circ}$, and 25° (dash-dotted lines).

the three-dimensional angular-momentum projection operator. Finally, the shell model Hamiltonian (1) is diagonalized in this good-angular-momentum basis.

1. Band structures

The TPSM band structures and the associated electromagnetic transition rates obtained in TPSM are quite rich and will be discussed in a separate paper [60]. Figure 10 shows the results for ¹¹²Ru as an representative example, as in this nucleus all bands are known up to high spins. It is seen that TPSM reproduces the experimental band energies quite well. The calculations slightly overestimate the bandhead energy of the $\gamma \gamma$ band; a similar result was also obtained for, e.g., ¹⁰⁸Mo and ¹⁰⁸Ru. The first q.p. $(h_{11/2})^2$ neutron alignment is predicted around I = 8, and the transition to a 4-q.p. $nu(h_{11/2})^2 \pi (g_{9/2})^2$ band is expected to occur around I = 16.

To illustrate the importance of the γ degree of freedom in the description of the band structures in the Mo and Ru isotopes, we have carried out TPSM calculations for ¹⁰⁶Mo for a range of γ values. The obtained band structures at $\gamma = 5^{\circ}, 15^{\circ}, 20^{\circ}$, and 25° are shown in Fig. 11. At $\gamma = 5^{\circ}$, the calculated γ and $\gamma\gamma$ bands are shifted with respect to experiment by more than 1 MeV. At $\gamma = 15^{\circ}, \gamma$ and $\gamma\gamma$ bands are shifted down in energy, and it is at $\gamma = 20^{\circ}$ that all the three bands are reproduced fairly accurately. At still higher value of $\gamma = 25^{\circ}$, the deviation from experiment grows again. For ¹⁰⁶Mo, therefore, $\gamma = 20^{\circ}$ is the optimum triaxial deformation in TPSM. Interestingly, the same value of γ is predicted by CHFB+UNEDF0, see Table I.

2. Transition quadrupole moments

Using the TPSM wave functions and standard *E*2 effective charges ($e_n = 0.5e$ and $e_p = 1.5e$) we have evaluated the transition quadrupole moments along the yrast line of the studied Mo and Ru isotopes, see Fig. 12. The overall behavior of the measured Q_t values is reproduced quite well by the TPSM approach. The drop in Q_t observed for all the studied isotopes around I = 8 is due to the quasiparticle alignment of the $h_{11/2}$ neutron pair, and in some isotopes there is a further drop around I = 16 due to a consecutive alignment involving the $g_{9/2}$ proton pair. Figure 12(a) also shows the TPSM predictions for Q_t in ¹⁰⁶Mo at $\gamma = 5^\circ$, 15°, and 25°. Again, it is seen that the best reproduction of experimental data is obtained at the optimum value of $\gamma = 20^\circ$.

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IV. CONCLUSIONS

Stimulated by the recent experimental data on the transition quadrupole moments in the rotational bands of neutron-rich Mo and Ru nuclei [23], we studied the shapes of the band structures in ^{106,108}Mo, and ^{108,110,112}Ru. We employed two complementary theoretical models: self-consistent CHFB+UNEDF0 approach and TPSM.

The triaxial PESs obtained in CHFB+UNEDF0 show stable g.s. triaxial minima in all cases. At higher angular momenta, the consecutive band crossings along the yrast line are expected, associated with the alignment of $\nu(h_{11/2})^2$ and $\pi(g_{9/2})^2$ pairs. The quadrupole deformations β_2 in the aligned bands are predicted to be reduced, but the shapes remain strongly triaxial. This result confirms predictions of an earlier works [7,31] based on the cranked macroscopic-microscopic method with the Woods-Saxon average potential. The decrease of the corresponding transition quadrupole moments above $\hbar\omega \sim 0.3$ MeV reflects the change in β_2 and γ due to the q.p. alignment, and this reduction is consistent with experiment.

The results obtained with TPSM paint the same picture as CHFB calculations, and strongly favor the triaxial interpretation. Indeed, both the energies of yrast, γ , and $\gamma\gamma$ bands, and transition quadrupole moments are well described assuming stable triaxial shapes. Similar as in our CHFB calculations, transition quadrupole moments obtained in TPSM exhibit a reduction at neutron and proton band crossings.

In summary, according to our analysis, high-spin behavior of ^{106,108}Mo and ^{108,110,112}Ru is consistent with triaxial rotation. The predicted triaxial g.s. minima are fairly shallow, and this perhaps is why in some calculations, e.g., the cranked relativistic Hartree-Bogoliubov model of Ref. [23], axial configurations may be slightly favored.

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