

Particle-number conserving analysis for the 2-quasiparticle and high- K multi-quasiparticle states in doubly-odd $^{174,176}\text{Lu}^*$

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Abstract: Two-quasiparticle bands and low-lying excited high- K four-, six-, and eight-quasiparticle bands in the doubly-odd $^{174,176}\text{Lu}$ are analyzed by using the cranked shell model (CSM) with the pairing correlations treated by a particle-number conserving (PNC) method, in which the blocking effects are taken into account exactly. The proton and neutron Nilsson level schemes for $^{174,176}\text{Lu}$ are taken from the adjacent odd- A Lu and Hf isotopes, which are adopted to reproduce the experimental bandhead energies of the one-quasiproton and one-quasineutron bands of these odd- A Lu and Hf nuclei, respectively. Once the quasiparticle configurations are determined, the experimental bandhead energies and the moments of inertia of these two- and multi-quasiparticle bands are well reproduced by PNC-CSM calculations. The Coriolis mixing of the low- K ($K=|\Omega_1-\Omega_2|$) two-quasiparticle band of the Gallagher-Moszkowski doublet with one nucleon in the $\Omega=1/2$ orbital is analyzed.

Key words: doubly-odd nucleus, moment of inertia, particle-number conserving method

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1 Introduction

The level structure of doubly-odd deformed nuclei is among the most complex topics in nuclear physics because of the complexity of level structure associated with contributions from both valence protons and neutrons. However, they often provide a wealth of nuclear structure phenomena. The rotational bands of deformed doubly-odd nuclei exhibit several unusual features and anomalies such as Gallagher-Moszkowski (GM) splittings [1], Newby shifts [2], chiral structure [3], signature inversion [4, 5], etc., which have been investigated from both experimental and theoretical sides [6–18]. A striking feature of axially symmetric deformed rare-earth nuclei with $A\sim 180$ is the observation of a large number of long-lived isomers [19]. In particular, many very long-lived isomers are observed in the doubly-odd nuclei. The most famous isomer $K^\pi = 9^-$ ($9/2^-[514]\otimes 9/2^+[624]$) in ^{180}Ta has a half-life longer than 1.2×10^{15} years [20], while the ground state $K^\pi = 1^+$ ($7/2^+[404]\otimes 9/2^+[624]$) has a half-life of about a few hours. These high- K bands arise from the coupling of a few orbitals near both proton and neutron Fermi surfaces, with large angular momentum projections Ω_i on the nuclear symmetry axes

($K = \sum_i \Omega_i$); e.g., the proton orbitals $\pi 7/2^+[404](g_{7/2})$, $\pi 9/2^-[514](h_{11/2})$, $\pi 5/2^+[402](d_{5/2})$, and the neutron orbitals $\nu 7/2^+[633](i_{13/2})$, $\nu 7/2^-[514](h_{9/2})$, $\nu 9/2^+[624](i_{13/2})$, etc.

Most of the earlier studies in this region have focused on the neutron deficient nuclei. With the development of experimental techniques, more and more measurements have been taken for the neutron rich nuclei. As for the neutron rich doubly-odd Lu nuclei, many experiments have been performed not only for the 2-quasiparticle (qp) structures, but also for the high- K multi-qp isomers [21–29].

It is well known that pairing correlations are very important in the low angular momentum region, where they are manifested by reducing the moments of inertia (MOIs) of the rigid-body estimation [30]. Due to the blocking effects, the MOIs of 1-qp bands in odd- A nuclei are usually larger than those of the ground state bands (gsbs) in adjacent even-even nuclei [31]. The blocking effects on MOI's of multi-qp bands are even more important [32]. In Ref. [33], the systematically observed low-lying high- K multi-qp bands in even-even and odd- A Hf and Lu isotopes ($170\leq A\leq 178$) [34] are analyzed using the cranked shell model (CSM) with pairing correlations

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treated by a particle-number conserving (PNC) method [35, 36]. Once an appropriate Nilsson level scheme near the Fermi surface for a given nucleus is adopted to reproduce the experimental bandhead energies of 1-qp bands in an odd- A nucleus, the experimental MOIs of these 1-qp bands and various kinds of low-lying high- K multi-qp bands can be reproduced well by the PNC-CSM calculations. In contrary to the conventional Bardeen-Cooper-Schrieffer (BCS) or Hartree-Fock-Bogolyubov (HFB) approaches, in the PNC method, the Hamiltonian is solved directly in a truncated Fock-space [37]. So the particle-number is conserved and the Pauli blocking effects are taken into account exactly. Note that the PNC scheme has been used in both relativistic and nonrelativistic mean field models [38, 39], in which the single-particle states are calculated from the self-consistent mean field potentials instead of the Nilsson potential. The PNC-CSM has also been used to investigate the heaviest nuclei with proton number $Z \approx 100$ [40–42].

In this paper, using the Nilsson level schemes of the odd- A Lu and Hf isotopes adopted in Ref. [33], we present the results of the doubly-odd $^{174,176}\text{Lu}$ from PNC-CSM calculations. The paper is organized as follows. A brief introduction of the PNC treatment of pairing correlations within the CSM is presented in Section 2. The PNC-CSM calculations of the excitation energies and MOIs for states in $^{174,176}\text{Lu}$ are presented in Section 3. A brief summary is given in Section 4.

2 A brief introduction to PNC-CSM

The CSM Hamiltonian of an axially symmetric nucleus in the rotating frame is [36, 43],

$$H_{\text{CSM}} = H_0 + H_P = H_{\text{Nil}} - \omega J_x + H_P, \quad (1)$$

where H_{Nil} is the Nilsson Hamiltonian [44], $-\omega J_x$ is the Coriolis interaction with the cranking frequency ω about the x axis (perpendicular to the nuclear symmetry z axis). H_P is the monopole pairing interaction

$$H_P = -G \sum_{\xi\eta} a_{\xi}^{\dagger} a_{\bar{\xi}}^{\dagger} a_{\bar{\eta}} a_{\eta}, \quad (2)$$

where $\bar{\xi}$ ($\bar{\eta}$) labels the time-reversed state of a Nilsson state ξ (η), and G is the effective strengths of the monopole pairing interaction, which is determined by the experimental odd-even differences in nuclear binding energies.

Instead of the usual single-particle level truncation used in common shell model calculations, a cranked many-particle configuration (CMPC) truncation (Fock space truncation) is adopted which is crucial to make the PNC calculations for low-lying excited states both workable and sufficiently accurate [37, 45]. An eigenstate of

H_{CSM} can be written as

$$|\psi\rangle = \sum_i C_i |i\rangle, \quad (C_i \text{ real}), \quad (3)$$

where $|i\rangle$ is an eigenstate of the one-body operator H_0 , i.e., a CMPC. By diagonalizing H_{CSM} in a sufficiently large CMPC space, sufficiently accurate solutions for low-lying excited eigenstates of H_{CSM} can be obtained.

The angular momentum alignment of state $|\psi\rangle$ is,

$$\langle\psi|J_x|\psi\rangle = \sum_i C_i^2 \langle i|J_x|i\rangle + 2 \sum_{i<j} C_i C_j \langle i|J_x|j\rangle, \quad (4)$$

and the kinematic MOI of state $|\psi\rangle$ is

$$J^{(1)} = \frac{1}{\omega} \langle\psi|J_x|\psi\rangle. \quad (5)$$

The occupation probability n_{μ} of the cranked orbital $|\mu\rangle$ is $n_{\mu} = \sum_i |C_i|^2 P_{i\mu}$, where $P_{i\mu} = 1$ if $|\mu\rangle$ is occupied in $|i\rangle$, and $P_{i\mu} = 0$ otherwise.

In the following PNC-CSM calculations, the Hamiltonian (1) is diagonalized in a sufficiently large CMPC space to obtain the low-lying excited eigenstates $|\psi\rangle$'s. The dimension of the CMPC space is about 700 for protons and 800 for neutrons. As we are only interested in the yrast and low-lying excited states, the number of the important CMPCs involved (weight $> 1\%$) is not very large (usually < 20) and almost all the CMPCs with weight $> 0.1\%$ are counted in, so the solutions to the low-lying excited states are quite accurate.

We note that because $R_x(\pi) = e^{-i\pi J_x}$, $[J_x, J_z] \neq 0$, the signature scheme invalidates the quantum number K . It has been pointed out that [36, 46], although $[J_x, J_z] \neq 0$, we have $[R_x(\pi), J_z^2] = 0$. Thus we can construct simultaneous eigenstates of $[R_x(\pi), J_z^2]$. Each CMPC $|i\rangle$ in Eq. (3) is chosen as a simultaneous eigenstate of (H_0, J_z^2) . However, K is still commonly used as a convenient quantum number to label rotational bands of deformed nuclei. For the gsb of an even-even nucleus, parity $\pi = +$, $\alpha = 0$, $K = 0$, $I = 0, 2, 4, \dots$. For an odd- A nucleus, if the Nilsson orbital 1 is blocked by an unpaired nucleon, we have two sequences of rotational levels with $\pi = \pi_1$, $K = \Omega_1$, $\alpha = \pm 1/2$, $I \geq K$. For a 2-qp band in a doubly-odd nucleus, when the proton and neutron Nilsson orbitals 1 and 2 are blocked by two unpaired nucleons, we have four sequences of rotational levels, $\pi = \pi_1 \pi_2$, $K = |\Omega_1 \pm \Omega_2|$, $\alpha = 0, 1$, $I \geq K$. The situation is similar for pair-broken (multi-qp) bands, etc.

3 Results and discussions

3.1 Parameters

Comparison of the PNC-CSM calculations on excitation energies and MOIs for the low-lying states in $^{174,176}\text{Lu}$ with the experimental values is shown in this section. The deformation parameters $\varepsilon_2 = 0.262, \varepsilon_4 =$

0.0402 ($\varepsilon_2 = 0.260, \varepsilon_4 = 0.0512$) for ^{174}Lu (^{176}Lu) are taken from Lund systematics [47], i.e., an average of the neighboring even-even Yb and Hf isotopes. The proton and neutron Nilsson level schemes of ^{174}Lu (^{176}Lu) are taken from those of ^{173}Lu and ^{175}Hf (^{177}Lu and ^{177}Hf) in Ref. [33], which are adopted to reproduce the bandhead energies of low-lying 1-quasiproton and 1-quasineutron bands in ^{173}Lu and ^{175}Hf (^{177}Lu and ^{177}Hf).

The effective pairing strengths can be determined by the odd-even differences in nuclear binding energies, and are connected with the dimension of the truncated CMPC space. The CMPC space for $^{174,176}\text{Lu}$ is constructed in the proton $N=4, 5$ shells and the neutron $N=5, 6$ shells. The dimensions of the CMPC space are about 700 for protons and 800 for neutrons in our calculation. The truncated CMPC's energies are about $0.8\hbar\omega_0$ for protons and $0.7\hbar\omega_0$ for neutrons, where $\hbar\omega_0 = 41A^{-1/3}$ MeV. The effective pairing interaction strengths for ^{174}Lu (^{176}Lu) are $G_p=0.33$ MeV (0.30 MeV) for protons and $G_n=0.30$ MeV (0.28 MeV) for neutrons, respectively. The stability of the PNC-CSM calculation results against the change of the dimension of the CMPC space has been investigated in Refs. [36, 42, 45, 48]. In the present calculations, almost all the CMPCs with weight $> 0.1\%$ are taken into account, so the solutions to the low-lying excited states are accurate enough. A larger CMPC space with renormalized pairing strengths gives essentially the same results.

When an unpaired proton and an unpaired neutron in a deformed doubly-odd nucleus are coupled, the projections of their total angular momentum on the nuclear symmetry axis, Ω_p and Ω_n , can produce two states with $K_> = \Omega_p + \Omega_n$ and $K_< = |\Omega_p - \Omega_n|$. They follow the GM coupling rules [1]:

$$K_> = |\Omega_p + \Omega_n|, \text{ if } \Omega_p = A_p \pm \frac{1}{2} \text{ and } \Omega_n = A_n \pm \frac{1}{2},$$

$$K_< = |\Omega_p - \Omega_n|, \text{ if } \Omega_p = A_p \pm \frac{1}{2} \text{ and } \Omega_n = A_n \mp \frac{1}{2}.$$

Note that the neutron-proton residual interaction is not included in our calculation, so we cannot get the GM doublets splittings.

3.2 ^{174}Lu

The cranked Nilsson levels near the Fermi surface of ^{174}Lu are given in Fig. 1. The positive (negative) parity levels are denoted by the blue (red) lines. The signature $\alpha = +1/2$ ($\alpha = -1/2$) levels are denoted by the solid (dotted) lines. Fig. 1 shows that there exists a proton sub-shell at $Z=70$ and a neutron sub-shell at $N=104$.

Using this Nilsson level scheme, first we calculate the bandhead energies of 2- and multi-qp states in ^{174}Lu with pairing interaction included. The results are shown in Table 1. Most of the data are well reproduced by the PNC-CSM calculations even without the neutron-proton residual interaction, especially those low-lying high- K

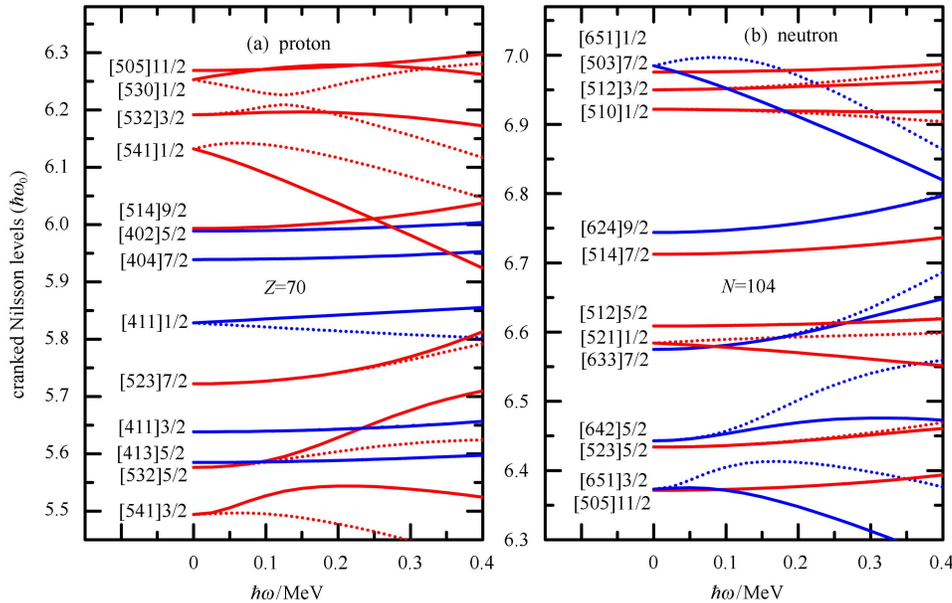


Fig. 1. (color online) The cranked Nilsson levels near the Fermi surface of ^{174}Lu for (a) protons and (b) neutrons. The deformation parameters ($\varepsilon_2 = 0.262, \varepsilon_4 = 0.0402$) are taken from Lund systematics [47], i.e., an average of the neighboring even-even Yb and Hf isotopes. The proton (neutron) Nilsson level scheme of ^{174}Lu is taken from that of ^{173}Lu (^{175}Hf) in Ref. [33], which is adopted to reproduce the bandhead energies of low-lying 1-quasiproton (1-quasineutron) bands of ^{173}Lu (^{175}Hf). The positive (negative) parity levels are denoted by the blue (red) lines. The signature $\alpha = +1/2$ ($\alpha = -1/2$) levels are denoted by the solid (dotted) lines.

multi-qp states. The only obvious exception is $K^\pi=2^+$ ($\pi 1/2[541] \otimes \nu 5/2^- [512]$). In our calculation the energy of $\pi 1/2[541]$ is too high. It is very hard to reproduce the correct location of this orbital in the Nilsson level scheme. For ^{174}Lu , the last occupied proton Nilsson orbital is $\pi 7/2^+[404]$ and the last occupied neutron Nilsson orbital is $\nu 5/2^- [512]$, which is consistent with the data. The experimental values of these 2-qp bands indicate that the orbital $\nu 7/2^+[633]$ is closer to the neutron Fermi surface than $\nu 1/2^- [521]$, which is reversed in our calculation. We should note that the level sequence in odd- A Hf isotopes in the $170 < A < 180$ mass region [49] is consistent with what we used. This level sequence inversion still needs to be investigated. For the 4-, 6-, and 8-qp states, the possible configurations and the corresponding bandhead energies are also shown. The results are quite similar to those multi-qp blocking calculations in Ref. [28]. For the state at 4710 keV, the calculated energies of the two configuration assignments $K^\pi=22^+$ and $K^\pi=22^-$ are both close to the data. So these two configurations are both possible for the observed state with excitation energy 4710 keV, which is consistent with the conclusion in Ref. [28]. Later on, the possible configuration assignments for $K^+=14^-, 15^+, \text{ and } 16^+$ will be analyzed according to the MOIs. For the calculations of the excitation energies and the MOIs, the configuration

assignments can be reasonably made.

The experimental kinematic MOIs for each band are extracted by

$$\frac{J^{(1)}(I)}{\hbar^2} = \frac{2I+1}{E_\gamma(I+1 \rightarrow I-1)}, \quad (6)$$

separately for each signature sequence within a rotational band. The relation between the rotational frequency ω and the angular momentum I is

$$\hbar\omega(I) = \frac{E_\gamma(I+1 \rightarrow I-1)}{I_x(I+1) - I_x(I-1)}, \quad (7)$$

where $I_x(I) = \sqrt{(I+1/2)^2 - K^2}$, K is the projection of nuclear total angular momentum along the symmetry z axis of an axially symmetric nuclei.

Figure 2 shows the experimental and calculated MOIs of the 2-qp bands in ^{174}Lu . As we mentioned in Section 2, for a 2-qp band in a doubly-odd nucleus there are four sequences of rotational levels. The experimental MOIs of the high- K ($K_> = \Omega_1 + \Omega_2$) and low- K ($K_< = |\Omega_1 - \Omega_2|$) bands are denoted by up and down triangles (both full and empty), respectively. The full and empty triangles denote the signature $\alpha=0$ and 1 bands, respectively. The calculated MOIs by the PNC-CSM are denoted by the dotted lines ($\alpha=1, \alpha_\pi = +1/2, \alpha_\nu = +1/2$), the dashed lines ($\alpha=1, \alpha_\pi = -1/2, \alpha_\nu = -1/2$), the short dashed lines ($\alpha=0, \alpha_\pi = -1/2, \alpha_\nu = +1/2$), the solid lines ($\alpha=0, \alpha_\pi = +1/2, \alpha_\nu = -1/2$). The blue dash dotted lines are the calculated results which are 50% admixture of the high- K and low- K states. The pairing interaction strengths $G_p=0.33$ MeV and $G_n=0.30$ MeV are determined by the experimental odd-even differences in nuclear binding energies.

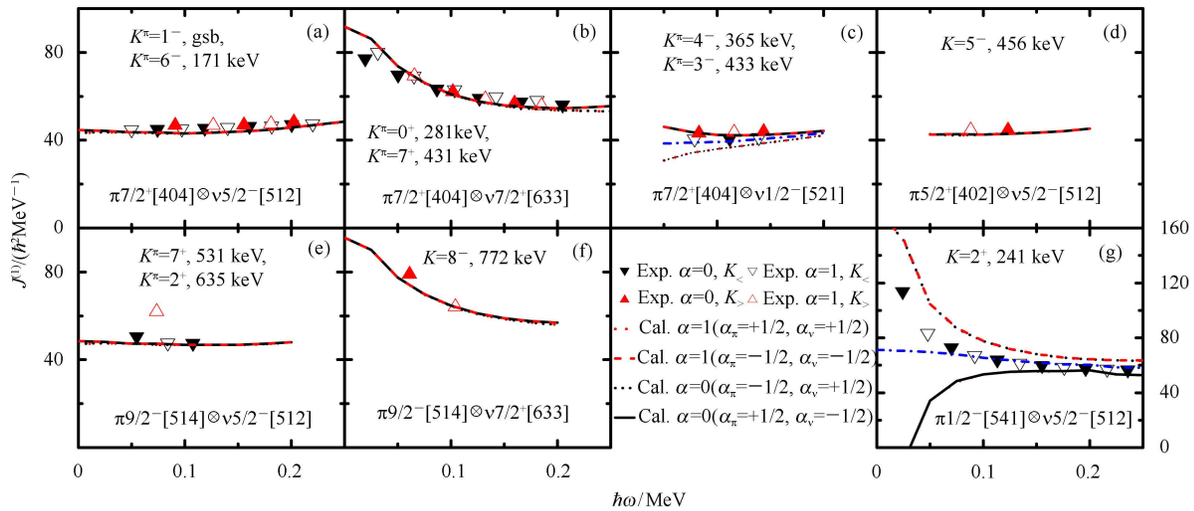


Fig. 2. (color online) The experimental and calculated MOIs $J^{(1)}$ of the 2-qp bands in ^{174}Lu . The experimental values are taken from Refs. [22, 23, 27, 50]. The experimental MOIs of the high- K ($K_> = \Omega_1 + \Omega_2$) and low- K ($K_< = |\Omega_1 - \Omega_2|$) bands are denoted by up and down triangles (both full and empty), respectively. The full and empty triangles denote the signature $\alpha=0$ and 1 bands, respectively. The calculated MOIs by the PNC-CSM are denoted by the dotted lines ($\alpha=1, \alpha_\pi = +1/2, \alpha_\nu = +1/2$), the dashed lines ($\alpha=1, \alpha_\pi = -1/2, \alpha_\nu = -1/2$), the short dashed lines ($\alpha=0, \alpha_\pi = -1/2, \alpha_\nu = +1/2$), the solid lines ($\alpha=0, \alpha_\pi = +1/2, \alpha_\nu = -1/2$). The blue dash dotted lines are the calculated results which are 50% admixture of the high- K and low- K states. The pairing interaction strengths $G_p=0.33$ MeV and $G_n=0.30$ MeV are determined by the experimental odd-even differences in nuclear binding energies.

lines ($\alpha=0, \alpha_\pi=-1/2, \alpha_\nu=+1/2$), the solid lines ($\alpha=0, \alpha_\pi=+1/2, \alpha_\nu=-1/2$). It can be seen that nearly all the experimental MOIs are reproduced by the PNC-CSM calculation quite well. An interesting thing is that there is almost no signature splitting in the $K_>$ or $K_<$ bands. The splitting happens between the GM doublet. The data of $K^\pi=7^+$ ($\pi 9/2^- [514] \otimes \nu 5/2^- [512]$) seems much larger than its GM partner band $K^\pi=2^+$. One possible reason is that the spin-parity assignment and the configuration assignment for this band may be unreasonable because there is almost no signature splitting in both $\pi 9/2^- [514]$ and $\nu 5/2^- [512]$.

It is well known that when one of the nucleons is in an $\Omega=1/2$ orbital, the GM doublet has $\Delta K=1$, and accordingly the two bands are expected to be Coriolis admixed. This effect can be very significant in the $K_<$ band, which has been identified in the doubly-odd rare-earth nuclei [13, 21]. In Fig. 2 the blue dash dotted lines are the calculated results which are 50% admixing of the high- K and low- K states of a GM doublet. As pointed out in Ref. [21], the mixing in $K^\pi=2^+$ ($\pi 1/2^- [541] \otimes \nu 5/2^- [512]$) band is very strong, which is almost 50% mixing with its high- K GM partner band. Our calculation shows that 50% mixing can reproduce the data quite well in the higher ω region, while in the very low ω region ($\hbar\omega < 0.05$ MeV), the mixing is less prominent, which is about 30%. Another Coriolis mixing band is the $K^\pi=3^-$ ($7/2^+ [404] \otimes \nu 1/2^- [521]$) band. This band also can be reproduced quite satisfactorily using 50% mixing which shows that in this GM doublet the mixing is as strong as that in $K^\pi=2^+$ ($\pi 1/2^- [541] \otimes \nu 5/2^- [512]$).

Figure 3 shows the experimental and calculated kinematic MOIs of the multi-qp bands in ^{174}Lu . All of these multi-qp isomers are high- K states. The experimental MOIs are denoted by the full up triangles (signature $\alpha=0$) and the open up triangles (signature $\alpha=1$), respectively. The calculated MOIs by the PNC-CSM are denoted by the solid lines (signature $\alpha=0$) and the dotted lines (signature $\alpha=1$), respectively. It can be seen that there is no signature splitting in these multi-qp isomers.

There are several possible configuration assignments for these multi-qp states. For $K^\pi=14^-$ band at 2063 keV, there are two possible configuration assignments. In Ref. [28], the configuration is assigned as $\pi 9/2^- [514] \otimes \nu^3 7/2^+ [633] 5/2^- [512] 7/2^- [514]$ based on the g_K value, the excitation energy and the alignment. In Fig. 3(b), MOIs of these two configurations are calculated to compare with the data. An alternative configuration $\pi 7/2^+ [404] \otimes \nu^3 7/2^+ [633] 9/2^+ [624] 5/2^- [512]$ for $K^\pi=14^-$ band is denoted as Config1 (red line). Both the calculated MOIs of these two configurations are very close to the data. The excitation energies of these two

configurations from the PNC-CSM calculations in Table 1 are also very close to each other, so it is very hard to say which configuration is possible from our calculation. For another two states, $K^\pi=15^+$ at 2732 keV and $K^\pi=16^+$ at 2876 keV, the configurations are chosen as $\pi 9/2^- [514] \otimes \nu^3 7/2^+ [633] 9/2^+ [624] 5/2^- [512]$ and $\pi^3 7/2^+ [404] 7/2^- [523] 9/2^- [514] \otimes \nu 9/2^+ [624]$, respectively. Fig. 3(c) and (d) shows that the experimental MOIs for $K^\pi=15^+$ and $K^\pi=16^+$ bands can be reproduced quite well using these two configuration assignments, respectively. Other configuration assignments are also possible for these two states. Config2 denotes an alternative configuration $\pi^3 7/2^+ [404] 7/2^- [523] 9/2^- [514] \otimes \nu 7/2^+ [633]$ for $K^\pi=15^+$ band. Config3 denotes an alternative configuration $\pi^3 7/2^+ [404] 7/2^- [523] 9/2^- [514] \otimes \nu 9/2^+ [624]$ for $K^\pi=16^+$ band. Obviously, the calculated MOIs of these two possible configurations (see Fig. 3(c)

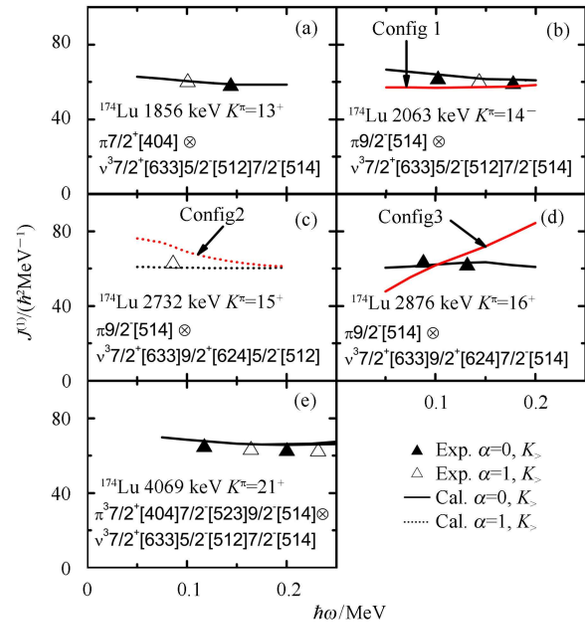


Fig. 3. (color online) The experimental and calculated MOIs $J^{(1)}$ of the multi-qp bands in ^{174}Lu . The experimental values are taken from Ref. [28]. The experimental MOIs are denoted by full up triangles (signature $\alpha=0$) and open up triangles (signature $\alpha=1$), respectively. The calculated MOIs by the PNC-CSM are denoted by the solid lines (signature $\alpha=0$) and the dotted lines (signature $\alpha=1$), respectively. Config1 denotes an alternative configuration $\pi 7/2^+ [404] \otimes \nu^3 7/2^+ [633] 9/2^+ [624] 5/2^- [512]$ for $K^\pi=14^-$ band. Config2 denotes an alternative configuration $\pi^3 7/2^+ [404] 7/2^- [523] 9/2^- [514] \otimes \nu 7/2^+ [633]$ for $K^\pi=15^+$ band. Config3 denotes an alternative configuration $\pi^3 7/2^+ [404] 7/2^- [523] 9/2^- [514] \otimes \nu 9/2^+ [624]$ for $K^\pi=16^+$ band.

Table 1. The experimental and calculated 2- and multi-qp states in ^{174}Lu . The experimental values are taken from Refs. [22, 23, 27, 28, 50].

K^π	configuration ^a	$E_{\text{exp}}/\text{keV}$	$E_{\text{cal}}/\text{keV}$
$1^-, 6^-$	$\pi 7/2^+ \otimes \nu 5/2^-$	0, 171	0
2^+	$\pi 1/2^- \otimes \nu 5/2^-$	241	1330
$0^+, 7^+$	$\pi 7/2^+ \otimes \nu 7/2^+$	281, 431	258
$4^-, 3^-$	$\pi 7/2^+ \otimes \nu 1/2^-$	365, 433	195
5^-	$\pi 5/2^+ \otimes \nu 5/2^-$	456	370
$7^+, 2^+$	$\pi 9/2^- \otimes \nu 5/2^-$	531, 635	399
8^-	$\pi 9/2^- \otimes \nu 7/2^+$	772	657
13^+	$\pi 7/2^+ \otimes \nu^3 7/2^+, 5/2^-, 7/2^-$	1856	1638
14^-	$\pi 9/2^- \otimes \nu^3 7/2^+, 5/2^-, 7/2^-$	2063	2037
14^-^b	$\pi 7/2^+ \otimes \nu^3 7/2^+, 9/2^+, 5/2^-$	2063	1873
15^+	$\pi 9/2^- \otimes \nu^3 7/2^+, 9/2^+, 5/2^-$	2732	2274
15^+^c	$\pi^3 7/2^+, 7/2^-, 9/2^- \otimes \nu 7/2^+$	2732	2490
16^+	$\pi 9/2^- \otimes \nu^3 7/2^+, 9/2^+, 7/2^-$	2876	2901
16^+^d	$\pi^3 7/2^+, 7/2^-, 9/2^- \otimes \nu 9/2^+$	2876	2885
19^+	$\pi 9/2^- \otimes \nu^5 7/2^+, 9/2^+, 1/2^-, 5/2^-, 7/2^-$	3741	3629
21^+	$\pi^3 7/2^+, 7/2^-, 9/2^- \otimes \nu^3 7/2^+, 5/2^-, 7/2^-$	4069	4127
22^-	$\pi^3 7/2^+, 7/2^-, 9/2^- \otimes \nu^3 7/2^+, 9/2^+, 5/2^-$	4710	4362
22^+^e	$\pi^3 7/2^+, 7/2^-, 9/2^- \otimes \nu^3 9/2^+, 5/2^-, 7/2^-$	4710	4950
23^-	$\pi^3 7/2^+, 7/2^-, 9/2^- \otimes \nu^3 7/2^+, 9/2^+, 7/2^-$	5062	4991
26^-	$\pi^3 7/2^+, 7/2^-, 9/2^- \otimes \nu^5 7/2^+, 9/2^+, 1/2^-, 5/2^-, 7/2^-$	5850	6119

^a Protons (π): $5/2^+ : 5/2^+ [402]; 7/2^+ : 7/2^+ [404]; 1/2^- : 1/2^- [541]; 7/2^- : 7/2^- [523]; 9/2^- : 9/2^- [514]$.

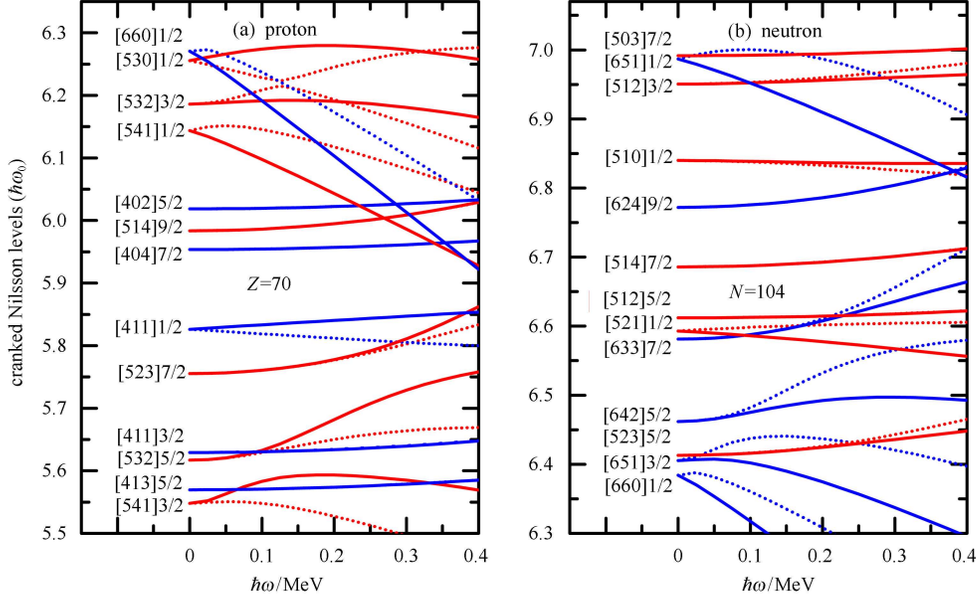
Neutrons (ν): $1/2^- : 1/2^- [521]; 5/2^- : 5/2^- [512]; 7/2^- : 7/2^- [514]; 7/2^+ : 7/2^+ [633]; 9/2^+ : 9/2^+ [624]$.

^b An alternative configuration for $K^\pi=14^-$, which is denoted as Config1 in Fig. 3(b).

^c An alternative configuration for $K^\pi=15^+$, which is denoted as Config2 in Fig. 3(c).

^d An alternative configuration for $K^\pi=16^+$, which is denoted as Config3 in Fig. 3(d).

^e An alternative spin-parity assignment for the state at 4710 keV.


 Fig. 4. (color online) The cranked Nilsson levels near the Fermi surface of ^{176}Lu for (a) protons and (b) neutrons.

The deformation parameters ($\varepsilon_2=0.260, \varepsilon_4=0.0512$) are taken from Lund systematics [47], i.e., an average of the neighboring even-even Yb and Hf isotopes. The proton (neutron) Nilsson level scheme of ^{176}Lu is taken from that of ^{177}Lu (^{177}Hf) in Ref. [33], which is adopted to reproduce the bandhead energies of low-lying 1-quasiproton (1-quasineutron) bands of ^{177}Lu (^{177}Hf). The positive (negative) parity levels are denoted by the blue (red) lines. The signature $\alpha=+1/2$ ($\alpha=-1/2$) levels are denoted by the solid (dotted) lines.

and (d)) are not consistent with the data, which is consistent with Ref. [28]. The configuration of the 6-qp bands $K^\pi=21^+$ at 4069 keV is assigned as $\pi 3^7/2^+[404] 7/2^- [523] 9/2^- [514] \otimes 7/2^+[633] 5/2^- [512] 7/2^- [514]$. In Fig. 3 the data can be well reproduced by this configuration, which also confirms the configuration assignment for this state.

3.3 ^{176}Lu

The cranked Nilsson levels near the Fermi surface of ^{176}Lu are given in Fig. 4. The positive (negative) parity levels are denoted by the blue (red) lines. The signature $\alpha=+1/2$ ($\alpha=-1/2$) levels are denoted by the solid (dotted) lines. It should be noted that the proton level sequence for ^{176}Lu is slightly different from that of ^{174}Lu . The level sequence of $\pi 9/2^- [514]$ and $\pi 5/2^+[402]$ is conversed in these two nuclei. The bandhead energies of the 2-qp states shown in Table 2 confirm this. Most of the data are well reproduced by the PNC-CSM calculations.

The experimental and calculated MOIs of the 2-qp bands in ^{176}Lu are shown in Fig. 5. Most of the data can be reproduced quite well by the PNC-CSM calcu-

lations. Note that in ^{176}Lu there are four GM doublets which have $\Delta K=1$. For $\pi 1/2^- [541] \otimes \nu 7/2^- [514]$ and $\pi 1/2^+[411] \otimes \nu 7/2^- [514]$, the 50% mixing of the low- K states with the high- K states can reproduce the data very well. The results show that in these two states the

Table 2. The experimental and calculated 2-qp states in ^{176}Lu . The experimental values are taken from Ref. [51].

K^π	configuration	$E_{\text{exp}}/\text{keV}$	$E_{\text{cal}}/\text{keV}$
$7^-, 0^-$	$\pi 7/2^+[404] \otimes \nu 7/2^- [514]$	0, 123	0
$1^+, 8^+$	$\pi 9/2^- [514] \otimes \nu 7/2^- [514]$	194, 488	208
$1^+, 8^+$	$\pi 7/2^+[404] \otimes \nu 9/2^+[624]$	339, 425	409
$1^-, 6^-$	$\pi 5/2^+[402] \otimes \nu 7/2^- [514]$	386, 564	432
$4^+, 3^+$	$\pi 1/2^- [541] \otimes \nu 7/2^- [514]$	635, 734	1293
$1^-, 6^-$	$\pi 7/2^+[404] \otimes \nu 5/2^- [512]$	638, 766	518
$3^-, 4^-$	$\pi 7/2^+[404] \otimes \nu 1/2^- [510]$	658, 788	1014
$4^-, 3^-$	$\pi 1/2^+[411] \otimes \nu 7/2^- [514]$	723, 843	528
$7^+, 2^+$	$\pi 5/2^+[402] \otimes \nu 9/2^+[624]$	734, 866	841
0^-	$\pi 9/2^- [514] \otimes \nu 9/2^+[624]$	780	617
$4^-, 3^-$	$\pi 7/2^+[404] \otimes \nu 1/2^- [521]$	908, 958	658
$0^+, 7^+$	$\pi 7/2^- [523] \otimes \nu 7/2^- [514]$	1057, 1274	1047

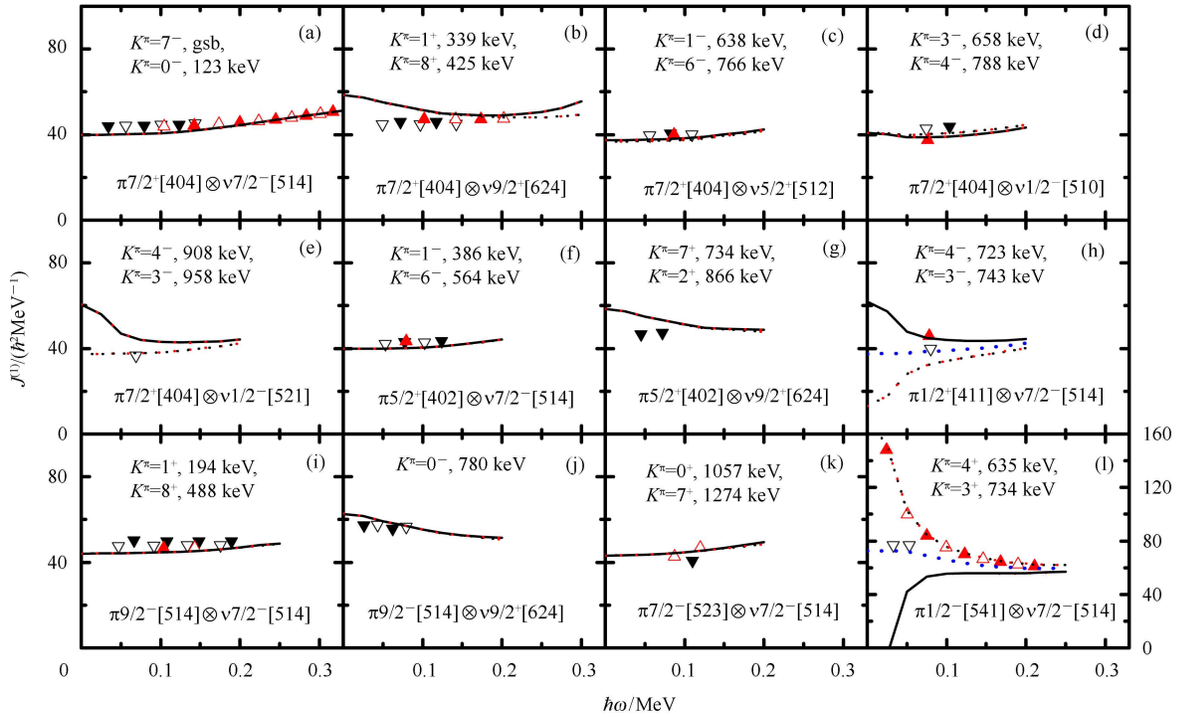


Fig. 5. (color online) The experimental and calculated MOIs $J^{(1)}$ of the 2-qp bands in ^{176}Lu . The experimental values are taken from Refs. [29, 51]. The experimental MOIs of the high- K ($K_> = \Omega_1 + \Omega_2$) and low- K ($K_< = |\Omega_1 - \Omega_2|$) bands are denoted by up and down triangles (both full and empty), respectively. The full and empty triangles denote the signature $\alpha=0$ and 1 bands, respectively. The calculated MOIs by the PNC-CSM are denoted by the dotted lines ($\alpha=1, \alpha_\pi=+1/2, \alpha_\nu=+1/2$), the dashed lines ($\alpha=1, \alpha_\pi=-1/2, \alpha_\nu=-1/2$), the short dashed lines ($\alpha=0, \alpha_\pi=-1/2, \alpha_\nu=+1/2$), the solid lines ($\alpha=0, \alpha_\pi=+1/2, \alpha_\nu=-1/2$). The blue dash-dotted lines are the calculated results which are 50% admixture of the high- K and low- K states. The pairing interaction strengths $G_p=0.30$ MeV and $G_n=0.28$ MeV are determined by the experimental odd-even differences in nuclear binding energies.

Coriolis mixing is also very strong. But there are exceptions. Because the signature splitting of the neutron orbital $\nu 1/2^-$ [510] in the PNC-CSM calculation is very small, so the Coriolis mixing in the $\pi 7/2^+[404] \otimes \nu 1/2^-$ [510] GM doublet cannot be determined by the MOIs. It is very interesting that the experimental MOIs of $K^\pi = 3^-$ ($\pi 7/2^+[404] \otimes \nu 1/2^-$ [521]) bands can be well reproduced by the PNC-CSM calculations without Coriolis mixing. We note the energy splitting of this GM doublet (50 keV) is much smaller than others (usually about 100 keV). So the absence of the Coriolis mixing of this state is due to the weaker proton-neutron residual interaction.

4 Summary

In summary, 2-qp bands and low-lying excited high- K 4-, 6-, and 8-qp bands in the doubly-odd $^{174,176}\text{Lu}$ are analyzed by using the cranked shell model with the pair-

ing correlations treated by a particle-number conserving method, in which the blocking effects are taken into account exactly. The proton and neutron Nilsson level schemes for $^{174,176}\text{Lu}$ are taken from the adjacent odd- A Lu and Hf isotopes, which are adopted to reproduce the experimental bandhead energies of the one-quasiproton and one-quasineutron bands of these odd- A Lu and Hf nuclei, respectively. The quasiparticle configurations are determined and the experimental bandhead energies and the MOIs of these 2- and multi-qp bands are well reproduced by PNC-CSM calculations. The Coriolis mixing of the low- K ($K = |\Omega_1 - \Omega_2|$) 2-qp band of the GM doublet involving $\Omega = 1/2$ is analyzed. The possible configuration assignments of the multi-qp isomers in ^{174}Lu are also analyzed both from the bandhead energies and the MOIs.

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