Triaxial quadrupole deformation dynamics in sd-shell nuclei around $^{26}$Mg

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Large-amplitude dynamics of axial and triaxial quadrupole deformation in $^{24,26}$Mg, $^{24}$Ne, and $^{28}$Si is investigated on the basis of the quadrupole collective Hamiltonian constructed with use of the constrained Hartree-Fock-Bogoliubov plus the local quasiparticle random-phase approximation method. The calculation reproduces well properties of the ground rotational bands, and $\beta$ and $\gamma$ vibrations in $^{24}$Mg and $^{28}$Si. The $\gamma$-softness in the collective states of $^{26}$Mg and $^{24}$Ne are discussed. Contributions of the neutrons and protons to the transition properties are also analyzed in connection with the large-amplitude quadrupole dynamics.

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I. INTRODUCTION

It is known that collective deformation grows up in the middle of the sd-shell region. The appearance of the prolate ground state of $^{24}$Mg and the oblate ground state of $^{28}$Si is associated with the shell gaps $N = Z = 12$ at the prolate region and $N = Z = 14$ at the oblate region in Nilsson diagram [5], respectively. Because of the shell gaps in the deformed regions, various shapes are expected to appear in the mass number region around $^{24}$Mg and $^{28}$Si.

Moreover, triaxial deformation degree of freedom plays very important roles on the low-lying collective dynamics in this mass region [6]. In $^{24}$Mg, possibility of the triaxial deformation in the ground states has been discussed for decades [7–9]. The low-lying $K = 2$ band built on top of the $2_1^+$ state suggests that the triaxial degree of freedom is activated in the collective dynamics. In $^{28}$Si, importance of triaxiality has been suggested in connection with the large-amplitude collective dynamics of the oblate-prolate shape coexistence [10, 11].

In contrast to the well-developed deformations in $^{24}$Mg and $^{28}$Si, the deformation property of $^{26}$Mg is not yet fully clarified. Since it is a system with $N = 14$ and $Z = 12$, neutrons and protons favor different shapes separately. Indeed, so far many mean-field calculations with use of the realistic effective interactions have been performed for $^{26}$Mg within an axial symmetry restriction, and they yielded a coexistence of oblate and prolate shapes with an oblate minimum [12–14]. On the other hand, the symmetry-unrestricted mean-field calculations using a Skyrme density functional (SkM*) [15] or relativistic model [16] show extremely triaxially soft potential energy surfaces.

In the study of collective excitations in this mass region, the quasiparticle random-phase approximation (QRPA) calculations have been systematically performed by employing various effective interactions [13, 15, 18]. The QRPA is a standard tool to analyze the collective modes of excitations. However, in order to discuss the low-lying collective dynamics of nuclei which are very soft against quadrupole deformation, one should use a microscopic theory of large-amplitude collective motion instead of the small-amplitude theory such as the QRPA. The generator coordinate method (GCM) with the restriction of axial symmetry [12], and the antisymmetrized molecular dynamics + multi-configuration mixing [19] have been performed using the energy density functionals for magnesium isotopes and $^{28}$Si, respectively. However, $^{26}$Mg is soft against $\beta$ and $\gamma$ directions as shown in the potential energy surface [12, 13], and therefore the triaxial degree of freedom in addition to the axial degree of freedom should be included for the description of the low-lying collective dynamics.

Quite recently, the GCM calculations including axial and triaxial generator coordinates have been performed [16, 20, 22] for magnesium isotopes. The first applications are concentrated on the low-lying states of $^{24}$Mg, in which the small-amplitude description in the prolate mean field is rather good. In Ref. [16], the properties of the yrast states of the magnesium isotopes are discussed systematically.

The quadrupole collective Hamiltonian provides a powerful theoretical tool to investigate the large-amplitude collective motion while taking into account the $\beta$ and $\gamma$ degrees of freedom [3, 23, 24]. Recently, on the basis of the adiabatic self-consistent collective coordinate method [28, 27], a new microscopic method to construct the collective Hamiltonian has been developed, called the constrained Hartree-Fock-Bogoliubov plus local QRPA (CHFB+LQRPA) [28]. In this method, the collective potential is calculated by the CHFB equation, while the inertial functions for large-amplitude quadrupole shape vibration and the three-dimensional rotation are determined from the normal modes on the CHFB state in ($\beta, \gamma$) plane. A new point of this method is that the contributions from the time-odd mean field are taken into account in evaluating the vibrational and rotational inertial masses. So far this CHFB + LQRPA method in conjunction with the pairing-plus-quadrupole (P+Q) model [28, 30] including the quadrupole-pairing force has been successfully applied to the oblate-prolate shape coexistence in proton-rich Se and Kr isotopes [28, 31].

In this paper, we analyze the role of triaxiality in con-
connection with the large-amplitude collective motion in the
calculation of the CHFB + LQRPA method with the P+Q model. We
also discuss the roles of neutrons and protons in $N \neq Z$
uclei on the large-amplitude collective dynamics in relation to the
electric transition properties. This article is
organized as follows. In the next section, the formulation
of the CHFB+LQRPA method is briefly recapitulated.
The results of the numerical calculations are presented
in Sec. III and the role of the triaxial degree of freedom
in these nuclei is discussed in Sec. IV. Summary is given
in Sec. V.

II. FORMULATION

A. CHFB+LQRPA method

The theoretical approach, the CHFB + LQRPA method is briefly summarized in this section. See Ref. [28] for detailed description of the method.

The method enables us to derive the five-dimensional quadrupole collective Hamiltonian of the Bohr-Mottelson type [23 27]

$$\mathcal{H}_{\text{coll}} = T_{\text{vib}} + T_{\text{rot}} + V(\beta, \gamma),$$

(1)

$$T_{\text{vib}} = \frac{1}{2} D_{\beta\beta}(\beta, \gamma) \dot{\beta}^2 + D_{\beta\gamma}(\beta, \gamma) \dot{\gamma} + \frac{1}{2} D_{\gamma\gamma}(\beta, \gamma) \dot{\gamma}^2,$$

(2)

$$T_{\text{rot}} = \frac{1}{2} \sum_{k=1}^{3} J_k(\beta, \gamma) \omega_k^2,$$

(3)

where $V(\beta, \gamma)$ is the collective potential in the $(\beta, \gamma)$ plane. The quantities $T_{\text{vib}}$ and $T_{\text{rot}}$ are the vibrational and rotational kinetic energies. The inertial functions $D_{\beta\beta}, D_{\beta\gamma},$ and $D_{\gamma\gamma}$ are the vibrational masses associated with the time-derivatives of the two quadrupole deformation variables, $\beta$ and $\gamma$, and $J_k$ are the rotational moments of inertia associated with the three components of the rotational angular velocities $\omega_k$ defined with respect to the principal axes.

The collective potential and the inertial functions in the collective Hamiltonian [21] are determined microscopically in the CHFB + LQRPA method. The collective potential is determined by solving the CHFB equation

$$\langle \phi(\beta, \gamma) | \hat{H}_{\text{CHFB}}(\beta, \gamma) | \phi(\beta, \gamma) \rangle = 0,$$

(4)

where the CHFB Hamiltonian is given as

$$\hat{H}_{\text{CHFB}} = \hat{H} - \sum_{\tau=n, p} \lambda(\tau)(\beta, \gamma) \hat{N}(\tau) - \sum_{m=0, 2} \mu_m(\beta, \gamma) \hat{D}_m(\tau),$$

(5)

with the constraints on particle numbers and quadrupole deformations. Here $\hat{H}$ is the microscopic Hamiltonian, $|\phi(\beta, \gamma)\rangle$ is the CHFB state, $\lambda(\tau)(\beta, \gamma)$ and $\mu_m(\beta, \gamma)$ are the Lagrange multipliers, $\hat{N}(\tau) \equiv \hat{N}(\tau) - \hat{N}_0(\tau)$ are the particle number operators measured from $\hat{N}_0(\tau)$ which are the neutron and proton particle numbers of the nucleus. The operators $\hat{D}_m(\tau)$ are the Hermitian part of the quadrupole operators given by $\hat{D}_m(\tau) \equiv (D_{2m} + D_{2-m})/2$. The collective potential is given by

$$V(\beta, \gamma) = \langle \phi(\beta, \gamma) | \hat{H} | \phi(\beta, \gamma) \rangle.$$

(6)

On top of the CHFB state, the local normal modes are calculated by solving the LQRPA equations

$$\delta \langle \phi(\beta, \gamma) | \left[ \hat{H}_{\text{CHFB}}(\beta, \gamma), \hat{Q}_i(\beta, \gamma) \right]$$

$$- \frac{1}{i} \hat{P}_i(\beta, \gamma) | \phi(\beta, \gamma) \rangle = 0,$$

(7)

$$\delta \langle \phi(\beta, \gamma) | \left[ C_i(\beta, \gamma), \dot{Q}_i(\beta, \gamma) \right]$$

$$- \frac{1}{i} C_i(\beta, \gamma) \dot{P}_i(\beta, \gamma) | \phi(\beta, \gamma) \rangle = 0.$$  

(8)

Here $\dot{Q}_i(\beta, \gamma)$ and $\dot{P}_i(\beta, \gamma)$ are the infinitesimal generators locally defined as functions of $(\beta, \gamma)$. The quantity $C_i(\beta, \gamma) = \omega_i^2(\beta, \gamma)$ is the squared eigen frequency of the normal mode. We choose two collective modes from the LQRPA modes, following the minimal metric criterion in Ref. [28]. The vibrational masses $D_{\beta\beta}, D_{\beta\gamma},$ and $D_{\gamma\gamma}$ are determined from the transformation of the collective coordinates spanned by the two LQRPA modes into $(\beta, \gamma)$.

The rotational moments of inertia are calculated by solving the LQRPA equations for rotation on top of the CHFB state.

$$\delta \langle \phi(\beta, \gamma) | \left[ \hat{H}_{\text{CHFB}}, \hat{I}_k(\beta, \gamma) \right]$$

$$- \frac{1}{i} \hat{J}_k(\beta, \gamma) | \phi(\beta, \gamma) \rangle = 0,$$

(9)

$$\langle \phi(\beta, \gamma) | \left[ \hat{I}_k, \hat{I}_{k'} \right] | \phi(\beta, \gamma) \rangle = i \delta_{kk'},$$

(10)

where $\hat{I}_k(\beta, \gamma)$ and $\hat{J}_k$ represent the rotational angles and the angular momentum operators with respect to the three principal axes associated with the CHFB state $|\phi(\beta, \gamma)\rangle$, and $\hat{J}_k$ are the LQRPA moments of inertia.

Pauli’s prescription is used to quantize the classical collective Hamiltonian [21]. From the solution of the collective Schrödinger equation

$$\left\{ \hat{T}_{\text{vib}} + \hat{T}_{\text{rot}} + V(\beta, \gamma) \right\} \Psi_{\alpha M}(\beta, \gamma, \Omega) = E_{\alpha M} \Psi_{\alpha M}(\beta, \gamma, \Omega),$$

(11)

we obtain the collective wave function $\Psi_{\alpha M}(\beta, \gamma, \Omega)$ as functions of quadrupole deformations $(\beta, \gamma)$ and three Euler angles $\Omega$. The collective wave function is specified by the angular momentum $I$, and its projection onto the $z$-axis of the laboratory frame, $M$, and $\alpha$ distinguishes the states which have the same $I$ and $M$. 
The collective wave function is written in the following form
\[ \Psi_{aIM}(\beta, \gamma, \Omega) = \sum_{K \geq 0, \text{even}} \Phi_{aIK}(\beta, \gamma) \langle \Omega | IMK \rangle, \quad (12) \]
where \( \Phi_{aIK}(\beta, \gamma) \) is the vibrational part of the collective wave function, and the rotational part is written as
\[ \langle \Omega | IMK \rangle = \frac{1}{\sqrt{16\pi^2(1 + \delta_{k0})}} [D^I_{MK}(\Omega) + (-)^I D^I_{M-K}(\Omega)]. \quad (13) \]
Here \( D^I_{MK} \) is the Wigner's rotation matrix and \( K \) is the projection of the angular momentum onto the \( z \)-axis in the body-fixed frame.

The vibrational wave functions are normalized as
\[ \int d\beta d\gamma |\Phi_{aI}(\beta, \gamma)|^2 |G(\beta, \gamma)|^{\frac{3}{2}} = 1, \quad (14) \]
and the volume element \(|G(\beta, \gamma)|^{\frac{3}{2}} d\beta d\gamma |\) is given by
\[ |G(\beta, \gamma)|^{\frac{3}{2}} d\beta d\gamma = 2\beta^4 \sqrt{W(\beta, \gamma)R(\beta, \gamma)} \sin 3\gamma d\beta d\gamma, \quad (16) \]
where \( W(\beta, \gamma) = \{D_{\beta\beta}(\beta, \gamma)D_{\gamma\gamma}(\beta, \gamma) - [D_{\beta\gamma}(\beta, \gamma)]^2\} \beta^{-2}, \quad (17) \]
\[ R(\beta, \gamma) = D_1(\beta, \gamma)D_2(\beta, \gamma)D_3(\beta, \gamma), \quad (18) \]
and the volume element \(|G(\beta, \gamma)|^{\frac{3}{2}} d\beta d\gamma |\) is related to the moments of inertia as
\[ J_k(\beta, \gamma) = 4\beta^2 D_k(\beta, \gamma) \sin^2(\gamma - 2\pi k/3). \]
The requantization form, the symmetries, and boundary conditions of the collective Hamiltonian are described in Ref. \[23\].

The electric properties are calculated following the discussions in Refs. \[24\] [51]. The value of \( B(E2) \) and the spectroscopic quadrupole moment are given by
\[ B(E2; \alpha I \rightarrow \alpha' I') = (2I + 1)^{-1} \langle \alpha I | D^{(E2)} | \alpha' I' \rangle^2, \quad (19) \]
and
\[ Q(\alpha I) = \sqrt{\frac{16\pi}{5}} \left( \frac{I}{-I} \frac{2}{0} \frac{I}{1} \right) \langle \alpha I | D^{(E2)} | \alpha I \rangle. \quad (20) \]
The reduced matrix element in Eqs. \[19\] and \[20\] is calculated as
\[ \langle \alpha I | D^{(E2)} | \alpha' I' \rangle = \int d\beta d\gamma |G(\beta, \gamma)|^{\frac{3}{2}} \rho^{(E2)}_{\alpha I \alpha' I'}(\beta, \gamma), \quad (21) \]
where \( \rho^{(E2)}_{\alpha I \alpha' I'}(\beta, \gamma) \) is the \( E2 \) transition density. The quantities \( D_{m+}^{(E2)}(\beta, \gamma) \) are the expectation values of the \( E2 \) operator in the intrinsic frame,
\[ D_{m+}^{(E2)}(\beta, \gamma) = \langle \phi(\beta, \gamma) | \sum_{\tau = n, p} e^{(\tau)} D_{m+}^{(\tau)} | \phi(\beta, \gamma) \rangle, \quad (23) \]
where \( e^{(\tau)} \) are the neutron and proton effective charges, and \( D_{m+}^{(\tau)} \) are the neutron and proton parts of the quadrupole operators.

**B. Model Hamiltonian and parameters**

The pairing-plus-quadrupole model \[24\] [30] including the quadrupole-pairing interaction is adopted as a microscopic Hamiltonian in the present work. The single-particle model space consists of harmonic oscillator two-major shells (\( p \)-shell and \( sd \)-shell) both for neutrons and protons. The modified oscillator values are used as the spherical single-particle energies \[32\]. The values of the neutron and proton monopole pairing strengths and the quadrupole particle-hole interaction strengths are summarized in Table II. The strength of the quadrupole-pairing interaction is evaluated at the spherical CHFB state with use of the prescription proposed by Sakamoto and Kishimoto \[33\]. The interaction strengths of \( ^{24}\text{Mg} \) calculated with Skyrme SkM* density functional and the mixed surface-volume type pairing functional in Ref. \[35\]. A simple mass number dependence of the interaction strengths is used to obtain those for \( ^{26}\text{Mg}, ^{24}\text{Ne}, \) and \( ^{28}\text{Si} \). Only the quadrupole strength for \( ^{28}\text{Si} \) is increased in order to adjust the deformation of the oblate HFB minimum.

The Fock term is neglected following the conventional prescription of the P+Q model. Therefore we call the present framework Hartree-Bogoliubov (HB) instead of Hartree-Fock-Bogoliubov (HFB). Following Baranger and Kumar \[30\], the reduction factors are multiplied by the quadrupole matrix elements between the single-particle states in the upper shells, and the nuclear radial parameter \( R = 1.2A^{1/3} \) fm is used in the calculation of the harmonic oscillator length \( b \). In the calculation of \( E2 \) transition strengths and quadrupole moments, the quadrupole operator without reduction factor is used.
The two-dimensional mesh in the \( \beta \) and \( \gamma \) directions is used to express the collective Hamiltonian in the \( (\beta, \gamma) \) plane. The 60 mesh points are taken both in the range \( 0 < \beta < \beta_{\text{max}} \) and \( 0^\circ < \gamma < 60^\circ \). As for \( \beta_{\text{max}} \), 0.6 is used as common value, except \( \beta_{\text{max}} = 0.5 \) used for \( ^{26}\text{Si} \), because we could not get converged solution which satisfies the CHB equation with four constraints at large deformation \( \beta > 0.5 \) near the prolate region in \( ^{28}\text{Si} \).

### III. RESULTS

#### A. Collective potentials

The collective potentials calculated for \( N = Z \) nuclei, \( ^{24}\text{Mg} \) and \( ^{28}\text{Si} \), are shown in Figs. 1 and 2. The collective potential of \( ^{24}\text{Mg} \) shows a prolate minimum at \( \beta \sim 0.41 \), while that of \( ^{28}\text{Si} \) shows an oblate minimum at \( \beta \sim 0.26 \). As shown in Fig. 3, deformed shell gaps are found at prolately and oblately deformed regions in the present model. It corresponds to the appearance of the deformed minima in the collective potentials of \( ^{24}\text{Mg} \) and \( ^{28}\text{Si} \). In \( ^{24}\text{Mg} \), the triaxial potential valley from the prolate minimum to the oblate region exists in the small deformed region with \( \beta \sim 0.2 \). In \( ^{28}\text{Si} \), the collective potential curve around the oblate minimum is steep against the triaxial deformation. A prolate local minimum suggested by other microscopic calculations for \( ^{28}\text{Si} \) does not appear in the present calculation. Note that a potential energy surface similar to the present work is reported in the Skyrme HF + BCS calculation.

In contrast to the deep oblate and prolate minima in \( N = Z \) nuclei, the collective potentials of \( ^{26}\text{Mg} \) and \( ^{24}\text{Ne} \) presented in Figs. 3 and 4 show \( \beta \) and \( \gamma \) soft situations. The potential minima in \( ^{26}\text{Mg} \) and \( ^{24}\text{Ne} \) show small oblate deformations with \( \beta = 0.16 \) and 0.20, respectively. Around the potential minima, the collective potentials are soft against both axial and triaxial quadrupole deformations, suggesting the anharmonic situations in these nuclei.

In Fig. 6 proton and neutron monopole pairing gaps are plotted as functions of \( (\beta, \gamma) \). Basic characters of the proton pairing gaps in \( ^{26}\text{Mg} \) and \( ^{24}\text{Ne} \) are same as that in \( ^{24}\text{Mg} \), and those of neutron pairing gaps in \( ^{26}\text{Mg} \) and \( ^{24}\text{Ne} \) are same as that in \( ^{28}\text{Si} \). The proton monopole pairing gaps in \( ^{24}\text{Mg} \), \( ^{26}\text{Mg} \), and \( ^{24}\text{Ne} \) show strong \( (\beta, \gamma) \) dependence; they have the maximum at the spherical point, and they become zero around \( \beta \sim 0.4 \) in the prolate region. The neutron pairing gaps in \( ^{28}\text{Si} \), \( ^{26}\text{Mg} \), and \( ^{24}\text{Ne} \) becomes zero in the oblate region. These zero-gap regions correspond to the prolate \( Z = 12 \) and 10 shell gap and the oblate \( N = 14 \) shell gap in Fig. 3 respectively. Comparing the proton pairing gap for \( ^{26}\text{Mg} \) with that for \( ^{24}\text{Mg} \), and the neutron pairing gap for \( ^{26}\text{Mg} \) with that for \( ^{28}\text{Si} \), an interesting feature is seen in \( ^{26}\text{Mg} \); the zero-gap region extends widely in the \( (\beta, \gamma) \) plane in the case of \( ^{26}\text{Mg} \).

In the lower panel of Fig. 3 the neutron Nilsson diagram is shown as functions of \( \gamma \) at \( \beta = 0.3 \). The energies of the Nilsson orbits change gradually depending on \( \gamma \). Especially, the oblate shell gap at 14 and the prolate shell gap at 12 more smoothly vanish in the \( \gamma \) direction than in the \( \beta \) direction.
FIG. 3: Neutron Nilsson diagrams calculated within \( p \) and \( sd \)-shell model space as functions of \( \beta \) (a) and \( \gamma \) (b). Even and odd parity orbits are drawn with solid and dashed lines, respectively.

FIG. 4: (Color online) Collective potential \( V(\beta, \gamma) \) for \( ^{26}\text{Mg} \).

FIG. 5: (Color online) Collective potential \( V(\beta, \gamma) \) for \( ^{24}\text{Ne} \).

B. Properties of the LQRPA modes

As discussed in the previous subsection, one of the pairing features of the \( sd \)-shell nuclei is the collapse of the pairing gaps around the deformed shell gaps (Fig. 3). In the former applications of the LQRPA equations to Se and Kr isotopes [28, 31], where the systems are in the superconducting phase in all over the \( (\beta, \gamma) \) region considered, the properties and choices of the LQRPA modes in the normal phase has not been analyzed though they are interesting theoretical issues to be clarified. In Fig. 4 the eigen frequencies squared of the LQRPA modes \( \omega^2(\beta, \gamma) \), the vibrational part of the metric \( W(\beta, \gamma) \), and the neutron and proton monopole pairing gaps \( \Delta_0^{(n)} \) and \( \Delta_0^{(p)} \) in \( ^{24}\text{Mg} \) are plotted as functions of \( \gamma \) along the \( \beta = 0.425 \) line.

Let us label the chosen two collective modes in Fig. 7 (a) as ‘mode A’ and ‘mode B’. Mode A denotes the collective mode lower in energy at \( \gamma = 0^\circ \), and this mode jumps around \( \gamma = 12^\circ, 32^\circ, 36^\circ, \) and \( 41^\circ \). It becomes the second lowest mode at \( \gamma = 60^\circ \). Mode B denotes the collective mode higher in energy at \( \gamma = 0^\circ \), and this mode jumps around \( \gamma = 8^\circ \) and \( 25^\circ \). It becomes the lowest mode at \( \gamma = 60^\circ \).

As seen in Fig. 4 (c), in the region with \( 0^\circ < \gamma < 8^\circ \), both the neutron and proton pairing gaps vanish. In this region, lowest four eigen modes are close in energy around 1-2 MeV, and they correspond to the neutron and proton pairing vibrational modes at a normal phase (pair addition and pair annihilation). The fifth mode is chosen as a collective mode (mode A). It has the \( \gamma \)-vibrational character at the axial limit, and this mode is chosen continuously in the small \( \gamma \) where the neutron is normal. One important character in the normal system is that the collectivity of the \( \beta \) vibration is weakened [5]. In the present case, the \( \beta \) vibrational mode is found at around 7 MeV. In this energy region, the \( \beta \)-vibrational collective mode is sometimes embedded in other non-collective modes. However, the figure shows that one can always find such modes using the minimal metric criterion to select the two collective modes.

Around \( 8^\circ < \gamma < 12^\circ \), the proton becomes superconducting, and the character of the low-lying modes changes to the neutron pair addition and annihilation, proton pairing vibration, and proton pairing rotation (Nambu-Goldstone mode). Note that in Fig. 7 (a), the zero-energy pairing rotational mode is not shown.

In \( 12^\circ < \gamma < 25^\circ \), neutrons also become superconducting. As the pairing gaps significantly change as functions of deformation in this region, the low-lying vibrational modes have the pairing vibrational characters, and thus they are not chosen to evaluate the quadrupole collective
masses. In this region, the vibrational part of the metric increases.

This situation changes in $25^\circ < \gamma < 41^\circ$. The pairing vibration and quadrupole vibration mix in the lowest LQRPA mode, and the pairing-vibrational character of the lowest mode decreases. Therefore, the lowest LQRPA mode is continuously chosen as a collective mode (mode B) to oblate limit. On the other hand, the other collective mode (mode A) changes in several LQRPA modes in this region, because large quadrupole collectivity appears in these LQRPA modes. The quadrupole collectivities of these modes are similar, and the vibrational part of the metric $W$ is continuous around the jump.

Near the oblate region with $41^\circ < \gamma < 60^\circ$, the lowest two modes are chosen as the collective modes, and are decoupled with other LQRPA modes in energy. At the oblate axial limit, the lowest mode (mode B) becomes the $\gamma$ vibration, and the second lowest mode (mode A) corresponds to the $\beta$ vibration.

This analysis shows that the minimal metric criterion for selecting the two collective modes among low-lying LQRPA modes also works for the situation where the pairing gap vanishes.

C. Collective levels

In this subsection, we present the excitation spectra, electric transition properties, and collective wave functions that are obtained by solving the collective Schrödinger equation \((\text{III})\) for \(I \leq 6\) states.

I. \(^{24}\text{Mg}\)

The excitation spectra for \(^{24}\text{Mg}\) are shown in Fig. 8. The calculation yields an yrast rotational band composed of $0^+_1$, $2^+_1$, $4^+_1$, and $6^+_1$ states, and an excited side band composed of $2^+_2$, $3^+_1$, $4^+_2$, $5^+_1$, and $6^+_2$ states. These two bands are in a very good agreement with the experimental energy levels. To analyze the structure of each state, the vibrational wave functions squared are displayed in Fig. 4 where the $\beta^\gamma$ factor is multiplied, which carries the main $\beta$ dependence from the volume element $|G(\beta, \gamma)|^2d\beta d\gamma$ in Eq. (10). The members of the yrast band are localized around the prolate minimum, showing the prolate character of the yrast rotational band. The vibrational wave functions of the excited band are concentrated in the triaxial region around the prolate minimum with $\gamma \sim 20^\circ$. This can be interpreted as the $\gamma$ vibrational band of the prolate yrast state. We also analyze the $K$-component fraction for these states in Table III. The table shows that the $K$-mixing in these states are very small, and the results support the $K = 0$ ground band and $K = 2$ excited band. These features are almost unchanged even with the increase of angular momentum.

In Tables III and IV the electric properties of the low-lying states are summarized. In addition to the theoretical results calculated with the effective charges $(c^{(n)}_{\text{eff}}, c^{(p)}_{\text{eff}}) = (0.5, 1.5)$, we also list the values for pure neutron and proton contributions obtained by using $(c^{(n)}_{\text{eff}}, c^{(p)}_{\text{eff}}) = (1, 0)$ and $(0, 1)$. The calculated results reasonably reproduce the trend of the $E2$ transition strengths though they tend to underestimate the absolute values of experimental data. Using a different set of effective charges improves the systematical underestimation of the theoretical values. However, still there are some disagreement, for example, in $4^+_1 \rightarrow 2^+_1$, $3^+_1 \rightarrow 2^+_2$, and $5^+_1 \rightarrow 3^+_1$ transitions. One can see that the neutron and proton matrix elements are almost equal for all the transitions listed in the table. The prolate feature is shown also in the spectroscopic quadrupole moments. The calculated values for the yrast and the side bands are well described with the estimated values from the rotational collective model of a prolate deformation with $K = 0$ and 2, respectively. This shows the rigid deformation feature in $^{24}\text{Mg}$. 
FIG. 7: (a) The LQRPA eigen frequencies $\omega^2$, (b) the vibrational part of the metric $W$, and (c) the neutron and proton monopole pairing gaps $\Delta_0^{(\gamma)}$ in $^{24}\text{Mg}$ plotted as a function of $\gamma$ for a constant value $\beta = 0.425$. In (a), the frequencies squared of low-lying LQRPA eigen modes are denoted with the cross symbols, while the two collective modes chosen at each deformation to calculate the vibrational collective masses are denoted by the filled circles. In (b), the vibrational metrics are denoted by the filled circles, while the vibrational metrics calculated with other possible combinations of the LQRPA modes are denoted by the cross symbols.

As mentioned in Introduction, several triaxial GCM calculations are performed for the study of low-lying states in $^{24}\text{Mg}$ employing modern density functionals [20–22]. In comparison with them, the present calculation gives a remarkable agreement with the experimental data for the ground $K = 0$ band and the excited $K = 2$ band despite the schematic effective interaction and restricted model space. Especially, the agreement in the excitation energies are better than the GCM calculations, while that in the $B(E2)$ values are worse. The small $K$-mixing properties in the ground and excited bands shown in Table II are consistent with them [21, 22].

TABLE II: Probability amplitude of quantum states of $^{24}\text{Mg}$ for each $K$-component.

| $I^+\pi$ | $K = 0$ | $K = 2$ | $K = 4$ | $K = 6$ |
|----------|--------|--------|--------|--------|
|          | 1.00   | -      | -      | -      |
| $2^+_1$  | 0.99   | 0.01   | -      | -      |
| $4^+_1$  | 1.00   | 0.00   | 0.00   | -      |
| $6^+_1$  | 1.00   | 0.00   | 0.00   | 0.00   |
| $2^+_2$  | 0.03   | 0.97   | -      | -      |
| $3^+_2$  | -      | 1.00   | -      | -      |
| $4^+_2$  | 0.03   | 0.95   | 0.02   | -      |
| $5^+_2$  | -      | 1.00   | 0.00   | -      |
| $6^+_2$  | 0.02   | 0.97   | 0.01   | 0.00   |

In addition to the ground rotational band, we show two rotational bands built on the $2^+_2$ and $0^+_2$ states in Fig. 10. The values of $B(E2)$ and spectroscopic quadrupole moments are summarized in Tables V and VI, and the vibrational wave functions squared in $(\beta, \gamma)$ plane are shown in Fig. 11. The vibrational wave functions and the quadrupole moments indicate the oblate deformation of the yrast rotational band. In Table VII the $K$ mixing in each state is listed. The yrast rotational band is
momentum states, and this reduces the single-particle states plays a role in the high angular values of $Q$ from the rotational collective model using the calculated $K$ results overestimate both of the excitation energies and $E$ values of $K$ seen in Fig. 11. consistent with $K = 0$ oblate band. It is seen that the spectroscopic quadrupole moment significantly increases with the increase of the angular momentum. The values of $Q(4^+_1)$ and $Q(6^+_1)$ are larger than those estimated from the rotational collective model using the calculated $Q(2^+_1)$ value. (In the case of the $6^+_1$ state, 16.8 $e$ fm$^2$ is the estimated value for $K = 0$ rotational band.) This indicates that the $\beta$ deformation of the intrinsic state grows as the angular momentum increases in the yrast band as seen in Fig. 11.

We then compare the theoretical results for the ground band with the experimental data. The theoretical results overestimate both of the excitation energies and $B(E2)$ values for $4^+_1$ and $6^+_1$ states. The experimental values of $B(E2)$ do not follow the trend of the collective model. This indicates that the spin alignment of the single-particle states plays a role in the high angular momentum states, and this reduces the $B(E2)$ values. In the quadrupole collective Hamiltonian which we derive in the present calculation, the moments of inertia are evaluated at zero angular momentum, and the alignment of the single-particle states is not taken into account. To include such effect by using the cranked mean field approach is an interesting extension of the model, but is beyond the scope of this paper.

The vibrational wave functions of the theoretical $0^+_1$, $2^+_1$, and $4^+_1$ states have nodes in the $\beta$ direction, and this can be interpreted as the $\beta$-vibrational behavior on top of the oblate yrast states. The fact that in-band transitions in the $\beta$-vibrational band $B(E2; 2^+_1 \rightarrow 0^+_1)$ is comparable with those of the ground rotational band $B(E2; 2^+_1 \rightarrow 0^+_1)$, and the positive sign of the quadrupole moment in $2^+_1$ and $4^+_1$ supports this interpretation. However, they contain $K$ mixing and shape mixing with the

\begin{table}[h]
\centering
\caption{The values of $B(E2)$ for $^{24}$Mg listed in units of e$^2$fm$^4$. The theoretical values are calculated with the effective charges $(\epsilon_{eff}^n, \epsilon_{eff}^p) = (0.5, 1.5)$. The values calculated from pure neutron contribution $(\epsilon_{eff}^n, \epsilon_{eff}^p) = (1, 0)$ and proton contribution $(\epsilon_{eff}^p, \epsilon_{eff}^p) = (0, 1)$ are also listed. Experimental data are taken from Ref. [38].}
\begin{tabular}{cccc}
\hline
\textbf{EXP CHB+LQRPA} & \textbf{neutron} & \textbf{proton} \\
\hline
$2^+_1 \rightarrow 0^+_1$ & 88 & 63.026 & 16.189 & 15.614 \\
$4^+_1 \rightarrow 2^+_1$ & 160 & 96.171 & 24.663 & 23.838 \\
$6^+_1 \rightarrow 4^+_1$ & 155 & 108.032 & 27.684 & 26.784 \\
$3^+_1 \rightarrow 2^+_2$ & - & 103.484 & 26.686 & 25.602 \\
$4^+_2 \rightarrow 3^+_1$ & - & 80.216 & 20.674 & 19.849 \\
$5^+_1 \rightarrow 4^+_2$ & - & 57.085 & 14.713 & 14.125 \\
$6^+_2 \rightarrow 5^+_1$ & - & 47.575 & 11.220 & 11.786 \\
$4^+_2 \rightarrow 2^+_2$ & 64 & 44.673 & 11.448 & 11.076 \\
$5^+_1 \rightarrow 3^+_1$ & 149 & 65.981 & 16.944 & 16.347 \\
$6^+_2 \rightarrow 4^+_2$ & - & 83.500 & 21.415 & 20.697 \\
$4^+_1 \rightarrow 2^+_2$ & - & 0.011 & 0.003 & 0.003 \\
$2^+_2 \rightarrow 2^+_1$ & 15 & 17.197 & 4.216 & 4.327 \\
$2^+_2 \rightarrow 0^+_1$ & 8 & 4.911 & 1.179 & 1.244 \\
$3^+_1 \rightarrow 4^+_1$ & - & 5.091 & 1.241 & 1.284 \\
$3^+_1 \rightarrow 2^+_1$ & 10 & 8.180 & 1.948 & 2.078 \\
$4^+_2 \rightarrow 4^+_1$ & - & 12.100 & 2.925 & 3.059 \\
$6^+_1 \rightarrow 4^+_2$ & - & 0.018 & 0.004 & 0.005 \\
$4^+_2 \rightarrow 2^+_2$ & 5 & 3.493 & 0.849 & 0.882 \\
$5^+_1 \rightarrow 6^+_1$ & - & 4.217 & 1.018 & 1.066 \\
$5^+_1 \rightarrow 4^+_2$ & - & 7.618 & 1.825 & 1.931 \\
$6^+_2 \rightarrow 6^+_1$ & - & 9.756 & 2.348 & 2.470 \\
$6^+_2 \rightarrow 4^+_2$ & - & 3.534 & 0.867 & 0.889 \\
\hline
\end{tabular}
\end{table}

\begin{table}[h]
\centering
\caption{Spectroscopic quadrupole moments for $^{24}$Mg listed in units of $e$ fm$^2$. See also caption in Table III. Experimental data are taken from Ref. [38].}
\begin{tabular}{cccc}
\hline
\textbf{EXP CHB+LQPRA} & \textbf{neutron} & \textbf{proton} \\
\hline
$Q(2^+_1)$ & -16.6 & -15.7 & -7.97 & -7.81 \\
$Q(4^+_1)$ & -20.8 & -10.5 & -10.4 \\
$Q(6^+_1)$ & -23.7 & -12.0 & -11.8 \\
$Q(2^+_2)$ & -15.5 & 7.87 & 7.72 \\
$Q(3^+_1)$ & 0 & 0 & 0 \\
$Q(4^+_2)$ & -5.85 & -3.00 & -2.90 \\
$Q(5^+_1)$ & -13.0 & -6.58 & -6.45 \\
$Q(6^+_2)$ & -14.4 & -7.34 & -7.16 \\
\hline
\end{tabular}
\end{table}
prolate region in the vibrational wave functions.

The excited band composed of \( 2^+_2, 3^+_1, 4^+_2, 5^+_1, \) and \( 6^+_2 \) states are also found in the calculation. The main component of the vibrational wave functions is \( K = 2 \), and lies in the triaxially deformed region. However, there is no experimental information corresponding to this triaxial band. Since the prolate local minimum is not found in the collective potential, the prolate rotational band does not obtained in the energy spectrum.

\[ \begin{align*}
0^+ & \quad 2^+ & \quad 0^+_2 & \quad 0^+ \\
0 & \quad 0 & \quad 0 & \quad 0 \\
2^+_1 & \quad 3^+_1 & \quad 2^+_3 & \quad 0 \\
6^+_1 & \quad 5^+_2 & \quad 4^+_3 & \quad 6^+_2 \\
\end{align*} \]

FIG. 10: Excitation spectra calculated for \( {}^{28}\text{Si} \) by means of the CHB\text{+LQRPA} method and experimental data \([38]\).

3. \( {}^{26}\text{Mg} \)

By adding two neutrons to the prolately deformed \( {}^{24}\text{Mg} \), the character of collective dynamics in the low-lying levels drastically changes from that of \( {}^{24}\text{Mg} \). Figure 12 compares the theoretical and experimental low-lying energy levels in \( {}^{26}\text{Mg} \). In addition to the yrast band (\( 0^+_1, 2^+_2, 4^+_3 \) and \( 6^+_4 \)), we show a side band composed of \( 2^+_2, 3^+_1, 4^+_2, 5^+_1, \) and \( 6^+_2 \) states, which are connected by relatively large \( B(E2) \) values (see Table VIII).

The vibrational wave functions squared are shown in Fig. 13. A significant difference from \( {}^{24}\text{Mg} \) is seen in the deformation property of the vibrational wave functions of the ground \( 0^+_1 \) state. In contrast to the well-developed prolate structure of the \( {}^{24}\text{Mg} \) yrast band, the \( 0^+_1 \) state of \( {}^{26}\text{Mg} \) spreads over the triaxial region from oblate to prolate ones, although the shallow potential minimum is located at the oblate region. This indicates the very \( \gamma \)-soft character of the ground state. The members of the yrast band tend to localize around the prolate shape as the angular momentum increases. As for the excitation energies of the yrast band, the \( E_x(4^+_1)/E_x(2^+_1) \) ratio is 2.64 in theoretical calculation, which explains the experimental value 2.71 very well. As seen in Table IX, the spectroscopic quadrupole moments of the yrast band are consistent with the prolate deformation, but the absolute values are relatively smaller than those of \( {}^{24}\text{Mg} \). Moreover, \( Q(6^+_1) \) is almost twice of \( Q(2^+_1) \), and this is much larger than the value of \( K = 0 \) rotational collective model, indicating that the development of the prolate deformation with increase of the angular momentum. This feature is also seen in the ratio \( B(E2; 4^+_1 \rightarrow 2^+_1)/B(E2; 2^+_1 \rightarrow 0^+_1) \). The theoretical value of the ratio is 1.7, which is larger than the collective model value 1.43. However, the experimental value of the ratio 1.05 again indicates the effect of the single-particle alignment.

The quantum states in the side band distribute widely in the \( \gamma \) direction. This character is remarkably different from that of \( {}^{24}\text{Mg} \), where all the members of the \( K = 2 \) side band are localized in the triaxial region close to the prolate local minimum (Fig. 9). In particular, an oblate character develops in even angular momentum states of the side band, and the \( 4^+_2 \) and \( 6^+_2 \) states form the two-peak structure in the oblate and prolate region. This two-peak structure indicates the \( \gamma \)-soft character of the collective potential as discussed in Ref. [39].
FIG. 12: Excitation spectra calculated for $^{26}$Mg by means of the CHB+LQRPA method and experimental data \[38\].

![Excitation spectra](image)

FIG. 13: (Color online) Vibrational wave functions squared $\beta^4|\Phi_{01}(\beta, \gamma)|^2$ calculated for $^{26}$Mg.

![Vibrational wave functions](image)

TABLE V: The values of $\beta(E2)$ for $^{28}$Si listed in units of $e^2$fm$^4$. See also caption in Table IV. Experimental data are taken from Ref. \[38\].

| $\beta(E2)$ (e$^2$fm$^4$) | CHB+LQRPA neutron proton | EXP CHB+LQRPA neutron proton |
|--------------------------|--------------------------|-----------------------------|
| $0^+ \rightarrow 2^+$     | 16.7                      | 10.497                      |
| $2^+ \rightarrow 0^+$     | 9.15                      | 9.15                        |
| $2^+ \rightarrow 3^+$     | 3.12                      | 3.12                        |
| $3^+ \rightarrow 4^+$     | 1.55                      | 1.55                        |

TABLE VI: Spectroscopic quadrupole moments for $^{28}$Si listed in units of $e$ fm$^2$. See also captions in Table IV. Experimental data are taken from Ref. \[38\].

| $Q(E2)$ (e$^2$fm$^2$) | CHB+LQRPA neutron proton | EXP CHB+LQRPA neutron proton |
|-----------------------|--------------------------|-----------------------------|
| $0^+ \rightarrow 2^+$ | 12.0                      | 6.05                        |
| $2^+ \rightarrow 0^+$ | 17.8                      | 8.99                        |
| $2^+ \rightarrow 3^+$ | 22.2                      | 11.2                        |
| $3^+ \rightarrow 4^+$ | -12.6                     | -6.33                       |
| $4^+ \rightarrow 5^+$ | -3.10                     | -1.55                       |
| $5^+ \rightarrow 6^+$ | 10.3                      | 5.16                        |
| $6^+ \rightarrow 7^+$ | 1.61                      | 0.83                        |

4. $^{24}$Ne

Figures 14 and 15 display the energy spectra and vibrational wave functions squared for $^{24}$Ne, and the $\beta(E2)$ and the spectroscopic quadrupole moments are listed in Tables X and XI. The theoretical yrast band $0^+_1, 2^+_1, 4^+_1$, and $6^+_1$ has the $\gamma$-soft character similar to that of $^{26}$Mg;
the vibrational wave function of the 0\textsuperscript{+}\textsubscript{1} state has a peak at the oblate region, but spreads over the triaxial region. Moreover, the gradual shape change in the yrast states with the increase of angular momentum is also found in the calculation. The oblate peak of the vibrational wave function in the 0\textsuperscript{+}\textsubscript{1} state shifts to the prolate region in the 4\textsuperscript{+}\textsubscript{1} state.

The spectroscopic quadrupole moments of the yrast band are consistent with the prolate deformation, but the absolute values are smaller than 26\textsuperscript{Mg}.

### TABLE VIII: The values of $B(E2)$ for 26\textsuperscript{Mg} listed in units of e\textsuperscript{2}fm\textsuperscript{4}. See also caption in Table \textbf{III}. Experimental values are taken from Refs. [38, 40], and the assignment of the levels are given in Ref. [40].

| $K_2$ | EXP | CHB+LQRPA neutron | proton |
|-------|-----|--------------------|--------|
| 2\textsuperscript{+} | 2\textsuperscript{+} | 2.073 | 0.131 | 0.704 |
| 2\textsuperscript{+} | 2\textsuperscript{+} | 28.4 | 62.940 | 19.792 | 14.486 |
| 0\textsuperscript{+} | 2\textsuperscript{+} | 1.60 | 0.765 | 2.040 | 0.011 |
| 3\textsuperscript{+} | 4\textsuperscript{+} | 3.207 | 5.971 | 4.148 |
| 3\textsuperscript{+} | 4\textsuperscript{+} | 0.456 | 3.234 | 0.022 |
| 4\textsuperscript{+} | 4\textsuperscript{+} | 28.948 | 12.298 | 5.847 |
| 6\textsuperscript{+} | 4\textsuperscript{+} | 0.635 | 0.022 | 0.232 |
| 6\textsuperscript{+} | 5\textsuperscript{+} | 1.536 | 1.755 | 0.148 |
| 6\textsuperscript{+} | 5\textsuperscript{+} | 13.227 | 4.769 | 2.879 |
| 5\textsuperscript{+} | 4\textsuperscript{+} | 0.773 | 2.999 | 0.000 |
| 6\textsuperscript{+} | 6\textsuperscript{+} | 18.167 | 9.774 | 3.238 |
| 6\textsuperscript{+} | 4\textsuperscript{+} | 1.113 | 1.006 | 0.136 |

*Data taken from Ref. [40].

### TABLE IX: Spectroscopic quadrupole moment for 26\textsuperscript{Mg} listed in units of e\textsuperscript{2}fm\textsuperscript{4}. See also caption in Table \textbf{III}. Experimental data are taken from Ref. [38].

| $K_2$ | $Q_2\textsuperscript{+}$ | $Q_4\textsuperscript{+}$ | $Q_6\textsuperscript{+}$ |
|-------|-----------------|-----------------|-----------------|
| 0\textsuperscript{+} | -13.5 | -9.07 | -2.33 |
| 2\textsuperscript{+} | -14.9 | -4.44 | -8.48 |
| 4\textsuperscript{+} | -17.9 | -5.61 | -10.1 |
| 2\textsuperscript{+} | -7.17 | 2.51 | 3.95 |
| 3\textsuperscript{+} | 0 | 0 | 0 |
| 4\textsuperscript{+} | 2.74 | 2.84 | 0.88 |
| 5\textsuperscript{+} | -7.96 | -1.91 | -4.67 |
| 6\textsuperscript{+} | -3.25 | 1.60 | -2.70 |

FIG. 14: Excitation spectra calculated for 24\textsuperscript{Ne} by means of the CHB+LQRPA method and experimental data [38].

The two-peak structure in the 2\textsuperscript{+}, 4\textsuperscript{+}, and 6\textsuperscript{+} states in the side band is also seen in 24\textsuperscript{Ne} showing again the $\gamma$-soft character as well as 26\textsuperscript{Mg}.

The excitation energy of the 2\textsuperscript{+}\textsuperscript{1} state and $B(E2;2\textsuperscript{+}\textsuperscript{1} \rightarrow 0\textsuperscript{+}\textsuperscript{1})$ are reproduced by the theoretical calculation. However, the experimental energy spectrum is much more vibrational than the theoretical one.

### IV. DISCUSSION

#### A. Rotational hindrance of shape mixing in 26\textsuperscript{Mg} and 24\textsuperscript{Ne}

Here, we discuss the character of the vibrational wave functions in more detail. What is commonly seen in the calculated results for 26\textsuperscript{Mg} and 24\textsuperscript{Ne} is the localization of the vibrational wave functions squared as the increase of the angular momentum. While the ground 0\textsuperscript{+} state spreads over the $\gamma$ direction, the yrast band tends to localize in the ($\beta, \gamma$) plane around the prolate region, even though the collective potential has the shallow oblate minimum.

This rotational hindrance of shape mixing is also seen in the cases of oblate-prolate shape coexistence [39, 41]
prolate asymmetry is seen in the rotational moments of inertia, the rotational moments of inertia about the intermediate axis, the rotational kinetic energy, while it is unfavored in collective moments of inertia, this prolate region is favored in rotational kinetic energy, which increases as the deformation increases. As a result, the vibrational wave function for higher angular momentum states localizes, and the \( \beta \)-soft nature of the vibrational wave function is hindered.

The change of the yrast state structure discussed above is also seen in \(^{28}\text{Si}\). In the case of \(^{28}\text{Si}\), one can see that the deformation of the vibrational wave function grows as angular momentum increases. The minimum of the collective potential locates around \( \beta = 0.26 \), while...
the maximum of the moment of inertia locates around $\beta = 0.36$ in the oblate side. This is the reason why the deformation increases in the yrast band of $^{28}$Si. In case of $^{24}$Mg, however, the minimum of the collective potential and the maximum of the moment of inertia coincide around $\beta = 0.41$, and such a change in the structure of the yrast band does not occur.

![FIG. 16: (Color online) Rotational moments of inertia about the intermediate axis, $J_1(\beta, \gamma)$.](image)

B. Analysis with mirror nucleus method

One of the interesting issues in $N \neq Z$ nuclei in the middle of $sd$-shell region is the properties of the neutron and proton quadrupole transition matrix elements. In the mirror nucleus method \cite{43,44}, the proton matrix element $M_p$ is determined from the $E2$ transitions as $B(E2; I \rightarrow I') = M_p^2/(2I + 1)$, while the neutron one $M_n$ is determined from the same $E2$ transitions of the mirror nucleus as $B_{\text{mirror}}(E2; I \rightarrow I') = M_n^2/(2I + 1)$. Although the $M_n/M_p$ ratio should be equal to the ratio $N/Z$ in the simple collective model, it has been experimentally suggested that the $M_n/M_p$ ratio for the ground band transitions deviates from $N/Z$ in some $N \neq Z$ nuclei \cite{44,45} indicating possible difference between proton and neutron shapes or shape dynamics. Moreover, the $M_n/M_p$ ratio for the $2^+_2 \rightarrow 0^+_1$ transition in $^{26}$Mg is known to be extremely larger than the expected value $N/Z$ in mirror nucleus method and also in the analysis of $(p, p')$ scattering reactions \cite{44,47}. The value of $(M_n/M_p)/(N/Z)$ is $1.83 \pm 0.34$ \cite{38}. This clearly indicates the dominance of the neutron matrix element in this transition. In this subsection, we discuss the origin of the neutron dominance in the $2^+_2 \rightarrow 0^+_1$ transition in $^{26}$Mg in terms of the large-amplitude triaxial shape dynamics. We also discuss the yrast transition $2^+_1 \rightarrow 0^+_1$ in $^{26}$Mg and $^{24}$Ne.

1. $2^+_2 \rightarrow 0^+_1$ transition in $^{26}$Mg

In Table \textbf{XII} the experimental and theoretical values for $B(E2; 2^+_2 \rightarrow 0^+_1)$ transition are summarized. The theoretical value $(M_n/M_p)/(N/Z) = 2.15$ reproduces the neutron dominance very well. Actually, the bare proton contribution to this transition in $^{26}$Mg is more than hundred times smaller than the neutron one. As seen from Table \textbf{XII} and \textbf{XIII} such difference cannot be found in $N = Z$ nuclei. The shell model value 2.10 \cite{48} given with the effective charges $(e_n, e_p) = (0.35, 1.35)$ also explains the neutron dominance of this transition.

To analyze the mechanism of the neutron dominance in this transition, we present the $E2$ transition density $\rho_{\text{E2}}(\beta, \gamma)$ defined in Eq. \textbf{22} for $^{26}$Mg in Fig. \textbf{17}. Let us first compare $E2$ transition densities for $2^+_1 \rightarrow 0^+_1$ and $2^+_2 \rightarrow 0^+_1$. Because of the structure of the yrast vibrational wave functions, the sign of the transition density for the former in-band transition is positive all over the $(\beta, \gamma)$ deformation, while the transition density for the latter transition changes its sign in the $(\beta, \gamma)$ plane, since the vibrational wave function of an excited state has nodes in the $(\beta, \gamma)$ plane. In the case of the $2^+_2 \rightarrow 0^+_1$ transition as seen in Fig. \textbf{17}, the sign of the transition density is opposite in the prolate region and oblate region, and this results in the cancellation after the $(\beta, \gamma)$ integration. In the case of $^{26}$Mg, the proton matrix element is almost completely canceled after the integration of the transition density. Concerning the neutrons, the contribution to this transition density is relatively larger in the oblate region than in the prolate region, since the neutron favors the oblate deformation for $N = 14$ system. This situation produces the large $M_n/M_p$ ratio.

This cancellation taking place in the $(\beta, \gamma)$ plane is the result of the large-amplitude collective dynamics in the $(\beta, \gamma)$ plane, especially in the $\gamma$ direction. The importance of triaxial degree of freedom is clearly seen from Fig. \textbf{13} where the $E2$ transition density is plotted as a function of $\gamma$ for a constant value of $\beta$. It is seen that the proton transition density is almost anti-symmetric with respect to $\gamma = 30^\circ$, while the asymmetry is present for neutron transition density.

\begin{table}[h]
\centering
\begin{tabular}{|c|c|c|c|c|c|}
\hline
Nucleus & Experimental & CHB+LQRPA & Neutron & Proton & SM \\
\hline
$^{26}$Mg & 1.60 $\pm$ 0.32 & 0.765 & 2.040 & 0.011 & 3.28 \\
$^{26}$Si & 7.32 $\pm$ 2.28 & 4.82 & 2.040 & 19.7 & \textbf{3.28} \\
\hline
\end{tabular}
\caption{The values of $B(E2; 2^+_2 \rightarrow 0^+_1)$ for $^{26}$Mg and $^{26}$Si listed in units of $e^2$fm$^4$. Theoretical values for $^{26}$Mg are calculated with the effective charges $(e_n, e_p) = (0.5, 1.5)$, while those for $^{26}$Si are calculated assuming the mirror symmetry. The bare neutron and proton contributions calculated with $(e_n, e_p) = (1, 0)$ and (0,1) are also listed. Experimental data and shell model calculation are taken from Refs. \cite{38} and \cite{48}, respectively.}
\end{table}
lines show the calculations with three sets of effective charges, \((e_{\text{eff}}^{(n)}, e_{\text{eff}}^{(p)}) = (0.5, 1.5), (1, 0),\) and \((0, 1)\) are used for left, middle and right panels, respectively.

In contrast to the large \((M_n/M_p)/(N/Z)\) for the \(2\gamma^+ \rightarrow 0\gamma^+_1\) transition discussed above, the experimental value of the ratio \((M_n/M_p)/(N/Z)\) for the \(2\gamma^+ \rightarrow 0\gamma^+_1\) transition in \(26\text{Mg}\) is \(0.92 \pm 0.05\), which is close to unity. This indicates that a simple collective model picture with the usual assumption that the radius and the deformation for neutrons are consistent with those for protons is expected to hold. In the case of \(24\text{Ne}\), the experimental value of this ratio is \(0.59 \pm 0.11\). The possible suppression of the ratio from unity may suggest that the simple picture does not hold for neutrons and protons in this system, and the smaller neutron deformation than the proton deformation is expected in the ground state of \(24\text{Ne}\).}

We evaluate the \(M_n/M_p\) ratio by using the mirror nucleus method. In Table XIII \(B(E2; 2\gamma^+_1 \rightarrow 0\gamma^+_1)\) values for \(26\text{Mg}\), \(26\text{Si}\), \(24\text{Ne}\), and \(24\text{Si}\) are summarized. The relative magnitudes of \(E2\) transition probability for mirror pairs cannot be satisfactorily reproduced by the theoretical calculation both for \(A = 24\) and \(26\) systems.

The theoretical value of the ratio \((M_n/M_p)/(N/Z)\) for \(26\text{Mg}\) is 0.81. The calculated value is smaller than unity, and qualitatively reproduces the tendency of the experimental value. The shell model value \(48\) gives smaller ratio 0.69 than the present calculation. For \(24\text{Ne}\), the ratio is calculated to be 0.86. The results for \(24\text{Ne}\) fail to quantitatively describe the suppression of the ratio extracted from the central values of the experimental \(B(E2)\). For more detailed discussions, precise measurements of the \(E2\) transition strengths for \(24\text{Ne}\) and \(24\text{Si}\) are required.

### Table XIII

| System | \(M_n/M_p\) | \(N/Z\) | \(EXP\) | \(CHB+LQRPA\) | \(neutron\) | \(proton\) | \(SM\) |
|--------|-------------|--------|--------|---------------|-------------|-------------|--------|
| 26Mg   | 61.4 ± 1.8  | 52.870 | 11.131 | 13.953        | 57.1        |             |        |
| 26Si   | 70.5 ± 6.9  | 47.226 | 13.953 | 11.131        | 36.5        |             |        |
| 24Ne   | 28.0 ± 6.6  | 33.576 | 14.124 | 6.813         |             |             |        |
| 24Si   | 19.1 ± 5.9  | 48.197 | 6.813  | 14.124        |             |             |        |

\(26\text{Mg}\) and 24Ne\)

2. \(2\gamma^+_1 \rightarrow 0\gamma^+_1\) transition in 26Mg and 24Ne

In contrast to the large \((M_n/M_p)/(N/Z)\) for the \(2\gamma^+_1 \rightarrow 0\gamma^+_1\) transition discussed above, the experimental value of the ratio \((M_n/M_p)/(N/Z)\) for the \(2\gamma^+_1 \rightarrow 0\gamma^+_1\) transition in \(26\text{Mg}\) is \(0.92 \pm 0.05\), which is close to unity. This indicates that a simple collective model picture with the usual assumption that the radius and the deformation for neutrons are consistent with those for protons is expected to hold. In the case of \(24\text{Ne}\), the experimental value of this ratio is \(0.59 \pm 0.11\). The possible suppression of the ratio from unity may suggest that the simple picture does not hold for neutrons and protons in this system, and the smaller neutron deformation than the
V. SUMMARY

Large-amplitude triaxial quadrupole deformation dynamics in the low-lying states of sd-shell nuclei, $^{24}\text{Mg}$, $^{28}\text{Si}$, $^{26}\text{Mg}$, and $^{24}\text{Ne}$ are analyzed on the basis of the quadrupole collective Hamiltonian derived microscopically from the CHFB + LQRPA method.

As for the $N = Z$ systems, the calculation reproduces the prolate rotational band and the $\gamma$ vibrational band in $^{24}\text{Mg}$, and the oblate rotational band and the $\beta$ vibrational band in $^{28}\text{Si}$. As for $N \neq Z$ systems, $^{26}\text{Mg}$ and $^{24}\text{Ne}$, the collective potentials are shown to be soft against the $\beta$ and $\gamma$ deformations, and the large shape-fluctuation in the $(\beta, \gamma)$ plane is found in the vibrational wave functions of the ground states.

The yrast bands show rotational hindrance of the shape mixing, and the states localize around the prolate region as the angular momentum increases. The neutron and proton matrix elements for $2^+_1 \rightarrow 0^+_1$ transition in $^{26}\text{Mg}$ is explained in terms of the large-amplitude collective dynamics in the $\gamma$-direction. The neutron and proton matrix elements for $2^+_1 \rightarrow 0^+_1$ yrast transition are analyzed with use of the mirror nuclear method for $^{26}\text{Mg}$ and $^{24}\text{Ne}$. Also in other $N \neq Z$ nuclei, differences in the behavior of neutrons and protons in large-amplitude shape dynamics are expected to be interesting.

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