Precession mode on high-\(K\) configurations: non-collective axially-symmetric limit of wobbling motion

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Abstract
The precession mode, the rotational excitation built on the high-\(K\) isomeric state, in comparison with the recently identified wobbling mode has been studied. The random-phase-approximation (RPA) formalism, which has been developed for the nuclear wobbling motion, is invoked and the precession phonon is obtained by the non-collective axially symmetric limit of the formalism. The excitation energies and the electromagnetic properties of the precession bands in \(^{178}\)W are calculated, and it is found that the results of RPA calculations well correspond to those of the rotor model; the correspondence can be understood by an adiabatic approximation to the RPA phonon. As a by-product, it is also found that the problem of too small out-of-band \(B(E2)\) in our previous RPA wobbling calculations can be solved by a suitable choice of the triaxial deformation which corresponds to the one used in the rotor model.

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1. Introduction

In this paper, recent progress of the study on the wobbling and precession modes in nuclei is presented. These keywords, wobbling and precession, represent the motions of classical tops. They are quite interesting because they describe three-dimensional (3D) rotational motion, and so are related to a fundamental question: how does a nucleus rotate as a 3D quantum object?

It is not known how strictly these two are distinguished, but in the following, wobbling for motions of a triaxial-body and precession for those of an axially-symmetric body are used. In the lowest energy motion, the top rotates about one of the principal axes with largest moment of inertia, but when excited the angular momentum vector tilts from this axis in the body-fixed (intrinsic) frame. Then, looking at it from the laboratory frame, the body wobbles or precesses, and that is where these names come from. In classical mechanics, these two are similar; actually the difference is that the trajectory of the angular momentum vector is a pure circle in the case of precession, while it is an ellipse in the case of wobbling.

However, the atomic nucleus is a quantum system and the situation is dramatically changed: the collective rotation cannot occur about the symmetry axis. Thus, the quantum spectra corresponding to the wobbling and precession motions are completely different. In the case of wobbling, the spectra associated with an intrinsic configuration are composed of multiple rotational bands; the lowest (yrast) one represents the uniform rotation about the main rotation axis, the first (one-phonon) excited band represents a quantized motion of tilting angular momentum vector, and so on (more excited band with more tilting angle). These \((\Delta I = 2)\) rotational bands corresponds to a collective rotation about the main rotation axis with largest moment of inertia. Since the excitation of phonons, or tilting the angular momentum vector, is another type of rotation about the axis perpendicular to the main rotation axis, these excitations form \((\Delta I = 1)\) vertical sequences. In this way, the wobbling...
motions show a complicated band structure, horizontal rotational sequences and vertical phonon-like excitations.

On the other hand, the angular momentum along the symmetry axis (the main rotation axis) is generated by quasi-particle alignments in the high-K isomeric configurations, and no collective rotation exists about it in the case of precession. There are no horizontal sequences leaving only one vertical band for each intrinsic configuration, which is nothing but a collective rotation about the perpendicular axis to the high-K angular momentum.

Recently, the wobbling phonon spectra have been identified among the triaxial superdeformed (TSD) bands in Lu nuclei [1, 2], up to two-phonon excitations. By using the microscopic framework, the random-phase approximation (RPA), we have studied the wobbling phonon in the HF-Lu region [3, 4]. The precession bands are rotational bands excited on the prolate high-K isomers and have been known for many years, see e.g. [5]. We have recently investigated the properties of the precession modes applying the same RPA formalism in comparison with the wobbling modes [6, 7]. The content of this paper is largely based on the results of [7], and a further development on the out-of-band $B(E2)$ transition probability of wobbling phonon excitations (see the section 3).

2. Precession mode as a phonon

The study of wobbling and precession modes has a long history (see references quoted in [7]). Our recent works on wobbling motions rely on the microscopic framework developed in 1979 by Marshalek [8]. At almost the same time, a very important work for the precession mode has been done here in Lund in 1981 by Andersson et al [9] (cf also [10, 11]). In this work, the microscopic RPA formalism was used, which is known to be suitable to describe vibrational excitations. The reason why the RPA is employed for such specific rotational motions as the wobbling and precession modes can be easily understood in the case of a precession band, i.e. the high-K rotational band.

The spectra of a high-K rotor is well known:

$$E_{\text{high-K}}(I) = \frac{1}{2J_\perp} [I(I + 1) - K^2], \quad (I \geq K). \quad (1)$$

Putting $n = I - K$ and assuming $K$ is large, it is easy to see that this leads to an approximately harmonic spectrum,

$$E_{\text{high-K}}(n) = \omega_{\text{prec}} \left( n + \frac{1}{2} + \frac{n(n + 1)}{K} \right), \quad (2)$$

where $\omega_{\text{prec}} \equiv K / J_\perp$ is the precession phonon energy. The remaining anharmonic term is of order $(1/K)$ and can be neglected at the high-K limit. This harmonic picture is also valid for the $E2$ transitions,

$$B(E2) = \left( \frac{5}{16\pi} \right) Q_0^2 \langle I_f K 20 | I_i K \rangle^2. \quad (3)$$

By taking the high-K asymptotic limit in the Clebsch–Gordan coefficient,

$$\begin{align*}
B(E2; n + 1 \rightarrow n) &\propto \left( n + \frac{1}{2} \right), \\
B(E2; n + 2 \rightarrow n) &\propto \frac{3}{2} \left( \frac{n(n + 1)}{K^2} \right),
\end{align*}$$

in which the one-phonon transition is of order $(1/K)$ and the direct transition from the two-phonon state is of order $(1/K^2)$, so that the direct two-phonon transitions are prohibited in the high-K limit.

Now it is clear that the phonon picture is valid and the RPA formalism can be used to describe the precession mode microscopically, which has been done by Andersson et al [9]. The key of their work is to adopt the so-called symmetry-restoring separable type interaction: the residual interaction used in the RPA is constructed in such a way that the rotational symmetry broken by the deformed mean-field Hamiltonian is recovered. This uniquely determines the complete form of the residual interaction, and there is no adjustable parameter in the RPA calculation. The dispersion equation is simply given as $\omega S(\omega) = 0$, where the $\omega = 0$ solution is the rotational symmetry-restoring mode, i.e. the Nambu–Goldstone (NG) mode, $J_\perp = J_\perp + 1 J_\parallel$. Here and in the following we assume the $x$-axis as the high-K alignment axis. The equation for non-NG modes can be formally cast into the same form as $\omega_{\text{prec}} \equiv K / J_\perp$.

$$\omega_{\text{prec}} = \frac{K}{J_\perp(\omega_{\text{prec}})}, \quad (4)$$

but with microscopically defined moment of inertia,

$$J_\perp(\omega_{\text{prec}}) = \frac{1}{2} \sum_{\mu<\nu} \left[ \frac{|J_\mu(\nu)|^2}{E_{\mu\nu} - \omega_{\text{prec}}} + \frac{|J_\nu(\mu)|^2}{E_{\mu\nu} + \omega_{\text{prec}}} \right].$$

However, the inertia $J_\perp(\omega_{\text{prec}})$ is energy dependent and actually equation (4) is a nonlinear equation to obtain the RPA eigenenergy. Electromagnetic transitions can be calculated as the squared amplitudes of the RPA phonon $X_{\text{prec}}$ with respect to appropriate transition operators $Q_{\mu\nu}$:

$$B(\lambda, \Delta I) \approx \left( |K||Q_{\mu\nu=\Delta I}, X_{\text{prec}}^\dagger|K \right)^2 \quad (K \gg 1). \quad (5)$$

Here, it should be stressed that this RPA formalism of Andersson et al is obtained by taking the non-collective axially symmetric limit ($\gamma = 60$ or $-120^\circ$ in the Lund convention) of the RPA wobbling formalism of Marshalek [7]. In this way, the precession and wobbling can be considered as similar kinds of collective excitation modes.

The results of RPA calculations for precession modes in $^{178}$W [5] are presented in figure 1. In this nucleus, there are many high-K isomers observed, on which the rotational bands exist: 11 isomers investigated are ranging from four-quasi-particle states $(2\nu = 2\pi)$ to ten-quasi-particle states $(6\nu = 4\pi)$. We have used the Nilsson potential as a mean field with appropriate deformation parameters and pairing gap parameters. Except for four cases, $K^\pi = 18^\pi, 25^\pi, 28^\pi, 29^\pi$, the one-phonon excitation energies are well reproduced. As for these four isomers, the precession energies are too small; namely, the calculated moments of inertia are too large. We found that the overestimation of the calculated moments
of inertia is due to the proton contributions in these four configurations, which include the $\pi [541]_{1/2}^{1/2}$ Nilsson state originating from the $\pi h_{9/2}$ high-$j$ decoupled orbit. Apparently the contribution from this orbit is overestimated in the present calculations, see [7] for more detailed discussions.

As for the electromagnetic transitions, one can directly compare the calculated and measured transition probabilities, e.g. equation (3) for $B(E2)$. Thus, by using the asymptotic forms of the Clebsch–Gordan coefficients, $B(E2)$ and $B(M1)$ are parameterized in the high-$K$ limit as,

$$B(E2 : K + 1 \rightarrow K)_{\text{rot}} \approx \frac{15}{16\pi} \frac{1}{K} Q_0^2,$$
$$B(M1 : K + 1 \rightarrow K)_{\text{rot}} \approx \frac{3}{4\pi} (g_K - g_R)^2 K.\quad (7)$$

On the other hand, they are calculated by equation (5) in the RPA formalism. By equating these two expressions, we define $Q$ moment or $g_K - g_R$ factor calculated within the RPA. Namely, the results of RPA calculations have been parameterized in the same way as the rotor model, and they are compared with experimentally extracted ones in figure 2 for $B(M1)$. In this figure, the results of the simple mean-field approximation are included, namely using $g_K$ calculated as $\langle \mu_3 \rangle/\langle J_i \rangle$ ($\mu_3$ is the $M1$ operator) with each high-$K$ state, and a common value of $g_K$ calculated in the same way but with the ground state. As it is clearly seen in figure 2, the RPA calculation gives a much better description of $B(M1)$ values. As for the $E2$ transitions, there is no experimental data available, but by comparing the calculated RPA $Q$ moments with the mean-field estimate of $Q$ moments it is found that they coincide well except for four configurations, which include the $\pi h_{9/2}$ decoupled orbit (the figure is not shown, see [7]).

The results for the excitation energies and transition probabilities indicate that the high-$K$ rotor model picture is realized if the model parameters, moment of inertia, $Q$ moment and $g$-factors, are calculated appropriately by means of the RPA. This fact can be naturally understood by using an adiabatic approximation studied in [10, 11]; i.e. in this approximation, $X_{\text{prec}}^{(E)} \approx (J_i + K\Theta_{\gamma})_{\text{RPA}}/\sqrt{2K}$, while the NG mode is $X_{\text{NG}} = J_i/\sqrt{2K}$. By using this form it is easy to confirm, for example, $(Q_0)_{\text{mean-field}} \approx (Q_0)_{\text{RPA}}$; the situation is more complicated for the $g$-factors, see [7] for more detailed discussions.

3. Out-of-band transition of a wobbling phonon

Now let us come to the case of the wobbling mode. It has been shown that the RPA calculation well corresponds to the rotor model picture in the case of precession, especially for $B(E2)$. It can be shown that a similar consideration in terms of the adiabatic approximation can be also applied for the wobbling mode. However, the $B(E2)$ value of our RPA wobbling calculations in Lu nuclei was too small, by about $1/2 - 1/3$, compared with the experimental data, whose magnitudes are well reproduced by the rotor model. This has been a serious problem, see figure 3 and [3, 4]. Considering the results of the precession modes, however, it is difficult to understand the discrepancy between the results of the RPA calculation and the rotor model. Therefore, we looked for the reason why the RPA calculation gave such smaller values.

Two reasons have been found: one is related to the model space of the RPA calculations, and the other is to the definition of the triaxiality parameter $\gamma$ in the Nilsson potential. As for the model space, we used the 5-major shells but it was not enough; this effect, however, is only about 20% and not the major effect. More important is the value of the $\gamma$ parameter. It is believed that the $\gamma$ value is about $20^\circ$ in the TSD bands in the Lu region. We have used $20^\circ$ for the $\gamma$ in the Nilsson potential, $\gamma (\text{Nilso}) = 20^\circ$. It is found, however, that the $\gamma$-value defined by the density distribution, $\gamma (\text{dens})$, is only about $10^\circ$. Here, $\gamma (\text{dens})$ is defined by the expectation values of the quadrupole operators (the Lund convention for sign of $\gamma$),
The discrepancy between $\gamma$(Nils) and $\gamma$(dens) is just a matter of definition, see e.g. appendix of [14]. For the pure harmonic oscillator potential, the definition of $\gamma$(Nils) gives

$$
\gamma(\text{Nils}) = \tan^{-1} \frac{\sqrt{3}(\omega_x - \omega_y)}{2\omega_z - \omega_x - \omega_y}.
$$

(9)

The self-consistency of the potential (Mottelson condition), $\langle \sum_{k=1}^{3}(\gamma^2_{ik})_n \rangle \approx 1/\omega^2_k$ ($k = x, y, z$), is approximately satisfied in the Nilsson potential, and then by equation (8),

$$
\gamma(\text{dens}) \approx \gamma(\text{self}) = \tan^{-1} \frac{\sqrt{3}(1/\omega^2_x - 1/\omega^2_y)}{2/\omega^2_z - 1/\omega^2_x - 1/\omega^2_y},
$$

(10)

where $\gamma(\text{self})$ is the triaxiality based on the shape (ellipsoid) of an equi-potential surface for the anisotropic harmonic oscillator. In this way, the relation between $\gamma$(Nils) and $\gamma$(dens) in the upper panel of figure 3 can be naturally understood.

4. Summary

The RPA precession formalism by Andersson et al can be obtained by a non-collective axially-symmetric limit of the RPA wobbling formalism by Marshalek. By using this formalism, it is shown that the RPA calculation gives fairly good agreements for both the excitation energies and $B(M1)$ transitions for the precession phonons on high-$K$ isomers in $^{178}$W. This result can be interpreted by an adiabatic approximation, and shows a good correspondence between the RPA calculation and the rotor model; especially $B(E2)$ or $Q$ moments, and $B(M1)$ or $g$-factors [7].

There is an important feedback to the calculation of recently observed nuclear wobbling motions. The problem of small $B(E2)$ ratios in our previous RPA calculations can be solved if one uses a proper value of the triaxiality parameter $\gamma(\text{dens}) \approx 20^\circ$ in the density distribution. This does not completely solve the problem, however, because the Nilsson–Strutinsky calculation gives minima at $\gamma(\text{Nils}) \approx 30^\circ$ in the Nilsson potential, which corresponds to $\gamma(\text{dens}) \approx 10^\circ$. Thus, it raises a more fundamental question: why the Nilsson–Strutinsky calculation does not provide enough triaxiality, which is required for explaining the measured $B(E2)$ ratio of the wobbling phonon band. Finally, it is shown that the RPA can describe the 3D rotational motion (in the small amplitude approximation) at least at the same level as the rotor model. In other words, the rotor model may be justified by the microscopic RPA calculations.

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